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(54) **NON-HUMAN ANIMALS EXPRESSING HUMANIZED CD3 COMPLEX**

NICHT-MENSCHLICHE TIERE MIT EXPRESSION HUMANISIERTER CD3-KOMPLEXE
ANIMAUX NON HUMAINS EXPRIMANT UN COMPLEXE CD3 HUMANISÉ

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Description**FIELD OF INVENTION**

5 [0001] A genetically modified mouse is provided that comprises in its genome a nucleic acid sequence encoding a humanized CD3 ϵ , a humanized CD3 δ , and humanized CD3 γ . Thus, genetically modified mice that express humanized CD3 complex are provided. Also provided herein is a model for preclinical testing of CD3-based therapeutics, e.g., CD3-based antibodies, e.g., CD3-based bispecific antibodies.

10 BACKGROUND OF THE INVENTION

15 [0002] In addition to the T cell receptor subunits, e.g., highly variable TCR α and TCR β , the T cell receptor complex on the surface of a T cell comprises invariant CD3 ϵ , CD3 δ , and CD3 γ chains, which form heterodimers consisting of CD3 $\epsilon\delta$ and CD3 $\epsilon\gamma$. Also associated with the TCR/CD3 complex is the ζ chain, which is present as a disulfide-linked homodimer.

20 [0003] CD3 chains play a crucial role in T cell receptor assembly, transport to the cell surface, endocytosis of surface receptors, T cell development, and T cell signaling. For example, it has been demonstrated through studies of deficiencies of various CD3 subunits, that CD3 chains are important for double negative (CD4-CD8- or DN) to double positive (CD4+CD8+ or DP) to single positive (CD4+ or CD8+ or SP) T cell transition. In addition, each of CD3 ϵ , CD3 δ , and CD3 γ chains contains one immunoreceptor tyrosine-based activation motif (ITAM) while the ζ chain dimer contains 6 total ITAMs. These motifs serve as signaling modules, and are phosphorylated by associated kinases upon TCR engagement.

25 [0004] Antibodies against CD3 have been shown to cluster CD3 on T cells, thereby causing T cell activation in a manner similar to the engagement of the TCR by peptide-loaded MHC molecules. Thus, anti-CD3 antibodies have been proposed as therapeutic candidates aimed at activation of T cells. In addition, bispecific antibodies that are capable of binding CD3 and a target antigen have been proposed for therapeutic uses involving targeting T cell immune responses to tissues and cells expressing the target antigen.

30 [0005] A convenient animal model for preclinical testing of mono- and bispecific CD3-based therapeutic antibodies is particularly desired.

35 [0006] WO2003/006639 discloses a transgenic animal comprising a human polypeptide involved in the recognition and/or antigenic activation by T cells. Kuhn et al., Sci Transl Med. 3(68):68ra10 (2011), discloses transgenic mice expressing the human ϵ chain of the CD3 complex.

SUMMARY OF THE INVENTION

40 35 [0007] Provided herein is a genetically modified mouse according to claim 1. The endogenous mouse CD3 locus is genetically modified to encode an extracellular domain of human CD3 ϵ , an extracellular domain of human CD3 δ , and an extracellular domain of human CD3 γ . The endogenous mouse CD3 locus is genetically modified so as not to express functional extracellular domain(s) of the corresponding mouse protein(s). In one embodiment, the endogenous mouse CD3 locus further encodes transmembrane and cytoplasmic domains of corresponding endogenous mouse CD3 protein(s), wherein the animal expresses a chimeric CD3 protein on the surface of its T cells comprising the extracellular domain of the human CD3 protein and the transmembrane and cytoplasmic domains of the endogenous mouse CD3 protein. In one embodiment, a nucleic acid sequence(s) encoding the extracellular domain of the human CD3 in the mouse is operably linked to nucleic acid sequence(s) encoding transmembrane and cytoplasmic domains of the corresponding endogenous mouse CD3 protein(s). In a particular embodiment, the non-human animal comprises (a) at an endogenous CD3 ϵ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 ϵ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 ϵ , (b) at an endogenous CD3 δ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 δ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 δ , and (c) at an endogenous CD3 γ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 γ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 γ , wherein the mouse expresses chimeric CD3 ϵ , CD3 δ , and CD3 γ proteins on the surface of its T cells. In some embodiments, the extracellular domain of the human CD3 protein in the mouse comprises the sequence selected from the group consisting of SEQ ID NO:33, SEQ ID NO:34, and SEQ ID NO:35. In some embodiments, the mouse comprises extracellular domains of human CD3 proteins which comprise the sequences of SEQ ID NO:33, SEQ ID NO:34 and SEQ ID NO:35.

45 50 55 [0008] In some embodiments, the genetically modified non-human animal described herein comprises a nucleic acid sequence encoding an extracellular domain of human CD3 protein operably linked to a CD3 promoter. Thus, in some embodiments, the mouse described herein comprises a nucleic acid sequence encoding the extracellular domain of a

human CD3 ε , operably linked to a CD3 promoter, an extracellular domain of human CD3 δ , operably linked to a CD3 promoter, and an extracellular domain of human CD3 γ operably linked to a CD3 promoter. In one embodiment, the CD3 promoter is a non-human animal CD3 promoter. In one embodiment, the CD3 promoter is a human CD3 promoter. In one embodiment, the CD3 promoter is an endogenous mouse CD3 promoter.

5 [0009] Thus, in one embodiment, provided herein is a genetically modified mouse, wherein the mouse comprises (a) at an endogenous mouse CD3 ε locus a nucleic acid sequence encoding an extracellular domain of a human CD3 ε operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 ε , (b) at an endogenous mouse CD3 δ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 δ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 δ , and (c) at an endogenous mouse CD3 γ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 γ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 γ , and the mouse expresses humanized CD3 ε , CD3 δ , and CD3 γ proteins on the surface of its T cells. In one embodiment, the amino acid sequence of the humanized CD3 ε protein in said mouse is set forth in SEQ ID NO:24, the amino acid sequence of the humanized CD3 δ protein is set forth in SEQ ID NO:25, and the amino acid sequence of the humanized CD3 γ protein is set forth in SEQ ID NO:26. In one embodiment, the genetically modified mouse provided herein comprises a nucleic acid sequence encoding an extracellular domain of human CD3 operably linked to a mouse CD3 promoter. In one embodiment, the promoter is an endogenous mouse CD3 promoter. In another embodiment, the genetically modified mouse provided herein comprises a nucleic acid sequence encoding an extracellular domain of human CD3 operably linked to a human CD3 promoter. In one embodiment, the mouse displays similar CD4+ to CD8+ cell ratio in the thymus as compared to a mouse that is not genetically modified to express humanized CD3 proteins. In one embodiment, the mouse CD4+ to CD8+ T cell ratio in the thymus that is within 30%, within 25%, within 20%, within 15%, within 12%, within 10%, within 5%, or within 2% of the CD4+ to CD8+ cell ratio of a mouse that is not genetically modified to express humanized CD3 proteins. In one embodiment, the mouse displays similar T and B cell percentages in spleen, lymph nodes, and peripheral blood as a mouse that is not genetically modified to express humanized CD3 proteins. In one embodiment, the mouse displays similar numbers of circulating white blood cells, lymphocytes, monocytes, neutrophils, eosinophils, and basophils as a mouse that is not genetically modified to express humanized CD3 proteins.

10 [0010] Disclosed herein is a genetically modified mouse comprising at an endogenous mouse CD3 locus a nucleic acid sequence encoding an extracellular domain of human CD3 protein, wherein the human CD3 protein is selected from the group consisting of CD3 ε , CD3 δ , CD3 γ , CD3 ζ , and a combination thereof. The mouse of the invention comprises extracellular domains of human CD3 ε , CD3 δ , and CD3 γ . In one embodiment of the mouse, the extracellular domain of human CD3 ε is set forth in SEQ ID NO:33, the extracellular domain of human CD3 δ is set forth in SEQ ID NO:34, and the extracellular domain of human CD3 γ is set forth in SEQ ID NO:35. The mouse expresses a humanized CD3 ε , a humanized CD3 δ , and a humanized CD3 γ . In one embodiment of the mouse, the humanized CD3 ε is set forth in SEQ ID NO:24, the humanized CD3 δ is set forth in SEQ ID NO:25, and the humanized CD3 γ is set forth in SEQ ID NO:26. The mouse further comprises mouse CD3 ε , CD3 δ , and CD3 γ transmembrane and cytoplasmic domains. In one embodiment, the mouse further comprises endogenous mouse CD3 ε , CD3 δ , and CD3 γ transmembrane and cytoplasmic domains.

15 [0011] In another aspect, provided herein is a method of making a genetically modified mouse expressing a humanized CD3 protein, according to claim 7. In one embodiment of the method, the animal does not comprise a functional extracellular domain(s) of the corresponding mouse protein(s). In the method, the animal comprises at the endogenous CD3 locus a nucleic acid sequence encoding an extracellular domain of human CD3 ε , an extracellular domain of human CD3 δ , and an extracellular domain of human CD3 γ . In one embodiment of the method, the extracellular domain of human CD3 ε is set forth in SEQ ID NO:33, the extracellular domain of human CD3 δ is set forth in SEQ ID NO:34, and the extracellular domain of human CD3 γ is set forth in SEQ ID NO:35. In one embodiment of the method, the animal does not comprise functional extracellular domain(s) of the corresponding mouse protein(s). In one particular embodiment, the method comprises replacing at the endogenous CD3 locus an extracellular domain of a non-human CD3 protein(s) with a corresponding extracellular domain of a human CD3 protein(s). In one embodiment of the method, the animal further comprises a nucleic acid sequence(s) encoding transmembrane and cytoplasmic domains of corresponding endogenous mouse CD3 protein(s). In one embodiment of the method, replacement is at the endogenous mouse CD3 locus. In one embodiment the mouse expresses a humanized CD3 protein selected from the group consisting of a humanized CD3 ε set forth in SEQ ID NO: 24, a humanized CD3 δ set forth in SEQ ID NO:25, a humanized CD3 γ set forth in SEQ ID NO:26, and a combination thereof. In one embodiment of the method, the replacement is made in a single ES cell, and the single ES cell is introduced into the mouse embryo to make a mouse.

20 [0012] In yet another aspect, provided herein is a mouse model, e.g., a mouse model for testing a CD3-based bispecific antigen-binding protein, wherein the antigen-binding protein is capable of binding both CD3 and an antigen of interest, the mouse model comprising a mouse genetically modified to encode an extracellular domain of human CD3 protein, wherein the human CD3 protein is CD3 ε , CD3 δ , and CD3 γ , and comprising cell expressing or comprising the non-mouse

antigen of interest. In one embodiment of the mouse model, the nucleic acid sequence(s) of the humanized CD3 protein(s) is located at the endogenous CD3 locus. In one embodiment of the mouse model, the antigen-binding protein has been introduced into said mouse. The mouse expresses human CD3 ϵ , CD3 δ , and CD3 γ extracellular domains. The mouse model, the mouse further expresses mouse CD3 ϵ , CD3 δ , and CD3 γ transmembrane and cytoplasmic domains.

5 [0013] In one embodiment of the mouse model, the mouse comprises a xenograft of a tumor expressing the antigen of interest. In one embodiment of the mouse model, the cell expressing or comprising the antigen of interest is a tumor cell. In one embodiment of the mouse model, the bispecific antigen-binding protein selected binds to both the humanized CD3 protein and the antigen of interest. In one embodiment of the mouse model, the antigen of interest is a human antigen. In one embodiment of the mouse model, the antigen binding protein is capable of binding a monkey CD3 protein.

10 In one embodiment of the mouse model, the antigen of interest is a tumor associated antigen. In such an embodiment, the tumor associated antigen may be selected from the group consisting of ALK, BAGE proteins, BIRC5 (survivin), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, EGFR variant III, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, GAGE proteins, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5,

15 EBV-derived LMP2, L1CAM, MAGE proteins, MLANA, MSLN, MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), RAGE proteins, RET, RGS5, ROR1, SART1, SART3, SLAMF7, SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMPRSS2, Thompson-nouvelle antigen, TNFRSF17, TYR, UPK3A, VTCN1, WT1.

20 [0014] In another embodiment, the antigen of interest is an infectious disease associated antigen. In such an embodiment, the mouse may be infected with an infectious agent. In one such embodiment, the infectious disease associated antigen may be a viral antigen and the viral antigen is selected from the group consisting of HIV, hepatitis A, hepatitis B, hepatitis C, herpes virus (e.g., HSV-1, HSV-2, CMV, HAV-6, VZV, Epstein Barr virus), adenovirus, influenza virus, flavivirus, echovirus, rhinovirus, coxsackie virus, coronavirus, respiratory syncytial virus, mumps virus, rotavirus, measles virus, rubella virus, parvovirus, vaccinia virus, HTLV, dengue virus, papillomavirus, molluscum virus, poliovirus, rabies virus, JC virus, ebola virus, and arboviral encephalitis virus antigen. In another such embodiment, the infectious disease associated antigen may be a bacterial antigen and the bacterial antigen is selected from the group consisting of chlamydia, rickettsia, mycobacteria, staphylococci, streptococci, pneumonococci, meningococci, gonococci, klebsiella, proteus, serratia, pseudomonas, legionella, diphtheria, salmonella, bacilli, cholera, tetanus, botulism, anthrax, plague, leptospira, and Lyme disease bacterial antigen.

30 [0015] In one embodiment of the provided mouse model, the CD3-based antigen-binding protein is an antibody. In one embodiment, the CD3-based antigen-binding protein is a human or humanized antigen-binding protein. Such mouse model may allow testing for efficacy and/or toxicity of the antigen-binding protein in the mouse.

35 [0016] Also provided herein is a method of screening a drug candidate that target an antigen of interest comprising (a) introducing the antigen of interest into a genetically modified mouse comprising an endogenous mouse CD3 locus genetically modified to encode an extracellular domain of a human CD3 protein, wherein the human CD3 protein is CD3 ϵ , CD3 δ , and CD3 γ , (b) contacting the mouse with a drug candidate of interest, wherein the drug candidate is directed against the human CD3 and the antigen of interest, and (c) determining if the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest. In the method, the genetically modified mouse comprises at the endogenous mouse CD3 locus a nucleic acid sequence encoding an extracellular domain of human CD3 ϵ , an extracellular domain of human CD3 δ , and an extracellular domain of human CD3 γ . In one embodiment of the method, the mouse does not comprise a functional extracellular domain of the corresponding mouse protein(s). In the method, the mouse comprises a nucleic acid sequence(s) encoding transmembrane and cytoplasmic domains of corresponding endogenous mouse CD3 protein(s). In the method, the nucleic acid sequence(s) encoding the extracellular domain of the human CD3 is operably linked to the nucleic acid sequence(s) encoding transmembrane and cytoplasmic domains of the corresponding endogenous mouse CD3 protein(s). In one embodiment of the method, the extracellular domain of a human CD3 ϵ is set forth in SEQ ID NO:33, the extracellular domain of a human CD3 δ is set forth in SEQ ID NO:34, and the extracellular domain of a human CD3 γ is set forth in SEQ ID NO:35. Thus, in one particular embodiment of the method, the mouse expresses a humanized CD3 ϵ protein comprising an amino acid sequence set forth in SEQ ID NO:24, a humanized CD3 δ protein comprising an amino acid sequence set forth in SEQ ID NO:25, and a humanized CD3 γ protein comprising an amino acid sequence set forth in SEQ ID NO:26.

45 [0017] In a particular embodiment of the method of screening drug candidates described herein, the step of introducing the antigen of interest into the mouse described herein comprises expressing in the mouse the antigen of interest. In one embodiment, the step of expressing in the mouse the antigen of interest comprises genetically modifying the mouse to express the antigen of interest. In one embodiment, the step of introducing the antigen of interest comprises infecting the mouse with the antigen of interest. In one embodiment of the method, the step of introducing comprises introducing into said mouse a cell expressing the antigen of interest. In various embodiments of the method, the cell can be a tumor cell, a bacterial cell, or a cell infected with a virus. Thus, in some embodiments of the method, the mouse comprises

and infection which is either a viral or bacterial infection. Thus, the antigen of interest can be an infectious disease associated antigen. In one embodiment, the antigen of interest is a viral antigen, and the viral antigen is selected from the group consisting of HIV, hepatitis A, hepatitis B, hepatitis C, herpes virus (e.g., HSV-1, HSV-2, CMV, HAV-6, VZV, Epstein Barr virus), adenovirus, influenza virus, flavivirus, echovirus, rhinovirus, coxsackie virus, coronavirus, respiratory syncytial virus, mumps virus, rotavirus, measles virus, rubella virus, parvovirus, vaccinia virus, HTLV, dengue virus, papillomavirus, molluscum virus, poliovirus, rabies virus, JC virus, ebola virus, and arboviral encephalitis virus antigen. In another embodiment, the antigen of interest is an infectious disease associated antigen, which is a bacterial antigen selected from the group consisting of chlamydia, rickettsia, mycobacteria, staphylococci, streptococci, pneumonococci, meningococci, gonococci, klebsiella, proteus, serratia, pseudomonas, legionella, diphtheria, salmonella, bacilli, cholera, tetanus, botulism, anthrax, plague, leptospira, and Lyme disease bacterial antigen.

[0018] In another embodiment of the method of screening drug candidates, the antigen of interest is a tumor associated antigen. In one embodiment of the method, the tumor associated antigen is selected from the group consisting of ALK, BAGE proteins, BIRC5 (survivin), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, EGFR variant III, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, GAGE proteins, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5, EBV-derived LMP2, L1CAM, MAGE proteins, MLANA, MSLN, MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), RAGE proteins, RET, RGS5, ROR1, SART1, SART3, SLAMF7, SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMPRSS2, Thompson-nouvelle antigen, TNFRSF17, TYR, UPK3A, VTCN1, WT1.

[0019] In some embodiments of the method of screening drug candidates, the mouse is an immunocompetent mouse. In some embodiments of the method described herein, the antigen of interest is a human antigen of interest.

[0020] In some embodiments of the method, the drug candidate is an antibody. In some embodiments, the drug candidate is an antigen-binding protein. In some embodiments, the drug candidate is a bispecific antibody or a bispecific antigen binding protein. In some embodiments, the bispecific antigen binding protein is capable of binding both human CD3 protein and the antigen of interest. In one embodiment, the drug candidate is capable of recognizing a monkey CD3 protein.

[0021] In some embodiments of the method of screening drug candidates, the drug candidate is capable of reducing, eliminating, or preventing tumor growth as compared to an agent that does not target the antigen of interest. In some embodiments of such method the step of determining if the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest comprises a tumor volume assay or a T cell mediated tumor cell killing assay.

[0022] In other embodiments, the drug candidate is capable of reducing, eliminating, or preventing bacterial or viral infection as compared to an agent that does not target the antigen of interest. In some such embodiments, the step of determining if the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest comprises the measurement of viral or bacterial titers.

[0023] Disclosed herein is a mouse model, for testing safety, efficacy, and pharmacokinetics of combination drug therapies wherein the combination therapy includes a drug, e.g., an antigen-binding protein, that binds a human CD3 molecule. Such combination therapies are aimed at targeting specific tumors, infections, or other diseases described herein which can benefit from the recruitment and/or activation of T cells.

BRIEF DESCRIPTION OF THE DRAWINGS

[0024]

FIG. 1 depicts the structure of T cell receptor complex. The complex comprises two CD3 ϵ subunits, one CD3 δ subunit, one CD3 γ subunit, and two CD3 ζ subunits, complexed with the TCR $\alpha\beta$ heterodimer on a T cell surface. Asterisks indicate the locations of the ITAM motifs.

FIGS. 2A and B are the schematic representation (not to scale) of the humanized CD3 $\gamma\delta\epsilon$ large targeting vector.

FIG. 2A depicts the large targeting vector before the selection cassette (Neo) deletion, with human CD3E, CD3D, and CD3G sequence knock-in locations indicated. A, B, C, D, E, F, and G indicate location of the junction nucleic acid sequences represented in Table 1. **FIG. 2B** depicts the large targeting vector after deletion of the selection cassette (Neo); similarly to FIG. 2A, locations of the human CD3E, CD3D, and CD3G are indicated. A-B, C, D, E, F, and G are locations of the junction nucleic acid sequences represented in Tables 1 and 3.

FIG. 3 depicts the amino acid sequences of the humanized CD3 proteins in the humanized CD3 $\gamma\delta\epsilon$ mice. The CD3 protein sequences of human origin are underlined.

FIG. 4 depicts alignments between mouse and human CD3e, CD3d, and CD3g sequences. The 5' and 3' ends of the human sequences that were introduced into mouse CD3 loci are marked with * and **, respectively.

FIG. 5A, top row, is a FACs analysis plot demonstrating normal distribution of CD4+ and CD8+ thymocytes in wild

type (WT), heterozygous humanized CD3 $\gamma\delta\epsilon$ (HET), or homozygous humanized CD3 $\gamma\delta\epsilon$ (HO) mice. **FIG. 5B**, top row, is data depicting percentages as well as numbers of B and T cells in peripheral blood of indicated animals. **FIG. 5B** bottom row is data depicting percentages of T and B cells in the spleen of indicated animals. **FIG. 5C** shows V β repertoire polyclonality in CD4+ and CD8+ T cells obtained from the spleens of the humanized CD3 $\gamma\delta\epsilon$ mice.

FIGS. 6A-B are a demonstration of viral LCMV titers in the spleens of either wild type control or humanized CD3 $\gamma\delta\epsilon$ mice in mice infected with LCMV Clone 13 (**FIG. 6A**), or LCMV clone 13 following prior LCMV Armstrong clone infection (**FIG. 6B**).

FIG. 7 is data from the FACS analysis of splenocytes from wild type (WT), heterozygous humanized CD3 $\gamma\delta\epsilon$ (hCD3 $\gamma\delta\epsilon$ Het), or homozygous humanized CD3 $\gamma\delta\epsilon$ (hCD3 $\gamma\delta\epsilon$ Ho) mice sorted with two anti-human CD3 antibodies that also cross-react with monkey CD3 (ah/mfCD3-2 and ah/mfCD3-1), two anti-human CD3 antibodies that are human CD3 specific (ahCD3-1 and ahCD3-2), control anti-mouse CD3 (amCD3-2C11), unrelated control human IgG (control hIgG) and secondary only antibody control (2nd only). MFI values are listed in the tables below each graph.

FIG. 8 demonstrates response to anti-CD3 antibodies in humanized CD3 $\gamma\delta\epsilon$ mice. **FIG. 8A** demonstrates transient T and B cell depletion in blood of mice treated with anti-CD3 antibodies; either T cell depletion on day 1 for each antibody indicated (left figure), or T and B cell depletion and recovery over 14 days for each antibody tested (middle and right figures). **FIG. 8B** depicts an increase in concentration of cytokines released (IFN γ , KC, TNF α , IL-6, and IL-10) 2 hours after treatment with indicated antibodies.

FIG. 9 demonstrates splenocytes proliferation (measured as fold activation over cells only) upon treatment with increasing amounts of indicated antibodies in wild type (WT) and humanized CD3 $\gamma\delta\epsilon$ homozygous (hCD3 $\gamma\delta\epsilon$ Ho) mice.

FIG. 10 is a table summarizing various properties of the humanized CD3 mouse model.

FIG. 11A demonstrates the effect of anti-CD3 antibody (Ab-1; bispecific antibody recognizing CD3 and CD20, tested at two different concentrations) on tumor volume of B16F10.9/CD20 tumors when treatment is initiated at the same time as tumor implantation (prophylactic model). **FIG. 11B** demonstrates the effect of anti-CD3 antibody (Ab-1; bispecific antibody recognizing CD3 and CD20, tested at two different concentrations) on tumor volume of already established B16F10.9/CD20 tumors (therapeutic model).

DETAILED DESCRIPTION

30 Definitions

[0025] The present invention provides genetically modified mice according to claim 1. Disclosed herein are genetically modified mice that comprise in their genome, e.g., in their germline, genetically modified CD3 loci encoding humanized CD3 proteins, e.g., chimeric human/mouse CD3 proteins. Also disclosed are embryos, cells, and tissues comprising the same, methods of making the same, as well as methods of using the same. Unless defined otherwise, all terms and phrases used herein include the meanings that the terms and phrases have attained in the art, unless the contrary is clearly indicated or clearly apparent from the context in which the term or phrase is used.

[0026] "CD3," as used herein, includes an antigen which is expressed on T cells as part of the multimolecular T cell receptor (TCR) complex; the multimolecular TCR complex formed from association of homodimers and/or heterodimers comprising one or more of the following receptor chains: CD3-epsilon (ϵ), CD3-delta (δ), CD3-zeta (ζ), and CD3-gamma (γ) (See **FIG. 1**). Sequences and GenBank Accession Numbers of human and mouse CD3-delta, CD3-zeta, and CD3-gamma are presented in Table 4 below. Throughout the application, ϵ or epsilon can also be written as E, δ or delta can also be written as D, ζ or zeta can also be written as Z, and γ or gamma can also be written as G.

[0027] As used herein, "an antibody that binds CD3" or an "anti-CD3 antibody" includes antibodies and antigen-binding fragments thereof that specifically recognize a single CD3 subunit (e.g., epsilon, delta, gamma or zeta), as well as antibodies and antigen-binding fragments thereof that specifically recognize a dimeric complex of two CD3 subunits (e.g., gamma/epsilon, delta/epsilon, and zeta/zeta CD3 dimers). The antibodies and antigen-binding fragments of the present invention may bind soluble CD3 and/or cell surface expressed CD3. Soluble CD3 includes natural CD3 proteins as well as recombinant CD3 protein variants such as, e.g., monomeric and dimeric CD3 constructs, that lack a transmembrane domain or are otherwise unassociated with a cell membrane.

[0028] The term "conservative," when used to describe a conservative amino acid substitution, includes substitution of an amino acid residue by another amino acid residue having a side chain R group with similar chemical properties (e.g., charge or hydrophobicity). Conservative amino acid substitutions may be achieved by modifying a nucleotide sequence so as to introduce a nucleotide change that will encode the conservative substitution. In general, a conservative amino acid substitution will not substantially change the functional properties of interest of a protein, for example, the ability of CD3 proteins to play a role in T cell receptor assembly and signaling. Examples of groups of amino acids that have side chains with similar chemical properties include aliphatic side chains such as glycine, alanine, valine, leucine, and isoleucine; aliphatic-hydroxyl side chains such as serine and threonine; amide-containing side chains such as

asparagine and glutamine; aromatic side chains such as phenylalanine, tyrosine, and tryptophan; basic side chains such as lysine, arginine, and histidine; acidic side chains such as aspartic acid and glutamic acid; and, sulfur-containing side chains such as cysteine and methionine. Conservative amino acids substitution groups include, for example, valine/leucine/soleucine, phenylalanine/tyrosine, lysine/arginine, alanine/valine, glutamate/aspartate, and asparagine/glutamine.

5 In some embodiments, a conservative amino acid substitution can be a substitution of any native residue in a protein with alanine, as used in, for example, alanine scanning mutagenesis. In some embodiments, a conservative substitution is made that has a positive value in the PAM250 log-likelihood matrix disclosed in Gonnet et al. ((1992) Exhaustive Matching of the Entire Protein Sequence Database, *Science* 256:1443-45).

10 [0029] In some embodiments, the substitution is a moderately conservative substitution wherein the substitution has a nonnegative value in the PAM250 log-likelihood matrix.

[0030] Thus, encompassed by the invention is a genetically modified mouse expressing a humanized CD3 protein(s) comprising conservative amino acid substitutions in the amino acid sequence described herein.

15 [0031] One skilled in the art would understand that in addition to the nucleic acid residues encoding humanized CD3 proteins described herein, due to the degeneracy of the genetic code, other nucleic acids may encode the polypeptides of the invention. Therefore, in addition to a genetically modified mouse that comprises in its genome nucleotide sequences encoding humanized CD3 proteins described herein, a mouse that comprises in its genome nucleotide sequences that differ from those described herein due to the degeneracy of the genetic code are also provided.

20 [0032] The term "identity" when used in connection with sequence includes identity as determined by a number of different algorithms known in the art that can be used to measure nucleotide and/or amino acid sequence identity. In some embodiments described herein, identities are determined using a ClustalW v. 1.83 (slow) alignment employing an open gap penalty of 10.0, an extend gap penalty of 0.1, and using a Gonnet similarity matrix (MacVector™ 10.0.2, MacVector Inc., 2008). The length of the sequences compared with respect to identity of sequences will depend upon the particular sequences. In various embodiments, identity is determined by comparing the sequence of a mature protein from its N-terminal to its C-terminal. In various embodiments, when comparing a humanized sequence to a human sequence, the human portion of the humanized sequence (but not the non-human portion) is used in making a comparison for the purpose of ascertaining a level of identity between a human sequence and a humanized sequence.

25 [0033] The term "operably linked" includes a juxtaposition wherein the components so described are in a relationship permitting them to function in their intended manner. As such, a nucleic acid sequence encoding a protein may be operably linked to regulatory sequences (e.g., promoter, enhancer, silencer sequence, etc.) so as to retain proper transcriptional regulation. In addition, various portions of the humanized protein of the invention may be operably linked to retain proper folding, processing, targeting, expression, and other functional properties of the protein in the cell. Unless stated otherwise, various domains of the humanized protein of the invention are operably linked to each other. Operable linkage of a human extracellular domain of a CD3 protein and nonhuman transmembrane and cytoplasmic domains can be achieved by expressing these components as a contiguous fusion protein from a nucleic acid coding sequence.

30 [0034] The term "replacement" in reference to gene replacement includes placing exogenous genetic material at an endogenous genetic locus, thereby replacing all or a portion of the endogenous gene with an orthologous or homologous nucleic acid sequence. In one instance, an endogenous non-human gene or fragment thereof is replaced with a corresponding human gene or fragment thereof. For example, DNA encoding the extracellular domain of a mouse or other non-human CD3 protein can be replaced with DNA encoding the extracellular domain of the corresponding human protein A corresponding human gene or fragment thereof is a human gene or fragment that is an ortholog of, a homolog of, or is substantially identical or the same in structure and/or function, as the endogenous non-human gene or fragment thereof that is replaced. As demonstrated in the Examples below, nucleotide sequences encoding endogenous non-human CD3 extracellular domains were replaced by nucleotide sequences corresponding to human CD3 extracellular domains.

35 [0035] "Functional" as used herein, e.g., in reference to a functional protein, includes a protein that retains at least one biological activity normally associated with the native protein. For example, in some embodiments of the invention, a replacement at an endogenous locus (e.g., replacement at endogenous non-human CD3 loci) results in a locus that fails to express a functional endogenous protein.

40 [0036] The term "locus" as in CD3 locus includes the genomic DNA comprising a CD3 coding region. The different CD3 genes CD3 ϵ , CD3 δ , CD3 γ map proximate to one another the same chromosome. Thus depending on the context, reference to an endogenous CD3 locus may refer to a locus including some or all of these coding regions or an individual coding region. For example, if only one of the human CD3s, such as CD3 ϵ is introduced into a non-human animal, then the nucleic acid encoding that CD3 preferably modifies the locus of the corresponding non-human CD3. If several human CD3's are introduced into a non-human animal such as CD3 ϵ , CD3 δ , CD3 γ , then the modified endogenous locus includes the coding regions of each of CD3 ϵ , CD3 δ , and CD3 γ . A CD3 locus can also refer to the locus of CD3 ζ , which occupies a different chromosome than CD3 ϵ , CD3 δ , and CD3 γ . If human CD3 ζ is introduced together with any of human CD3 ϵ , CD3 δ , or CD3 γ , then two or more CD3 loci can be modified on different chromosomes. Other sequences may be included in the CD3 locus that have been introduced for the purposes of genetic manipulation, e.g., selection cassettes, restriction

sites, etc.

[0037] The term "germline" in reference to an immunoglobulin nucleic acid sequence includes a nucleic acid sequence that can be passed to progeny.

[0038] The phrase "immunoglobulin molecule" includes two immunoglobulin heavy chains and two immunoglobulin light chains. The heavy chains may be identical or different, and the light chains may be identical or different.

[0039] The term "antibody", as used herein, includes immunoglobulin molecules comprising four polypeptide chains, two heavy (H) chains and two light (L) chains inter-connected by disulfide bonds. Each heavy chain comprises a heavy chain variable domain and a heavy chain constant region (C_H). The heavy chain constant region comprises three domains, C_H1 , C_H2 and C_H3 . Each light chain comprises a light chain variable domain and a light chain constant region (C_L). The heavy chain and light chain variable domains can be further subdivided into regions of hypervariability, termed complementarity determining regions (CDR), interspersed with regions that are more conserved, termed framework regions (FR). Each heavy and light chain variable domain comprises three CDRs and four FRs, arranged from amino-terminus to carboxy-terminus in the following order: FR1, CDR1, FR2, CDR2, FR3, CDR3, FR4 (heavy chain CDRs may be abbreviated as HCDR1, HCDR2 and HCDR3; light chain CDRs may be abbreviated as LCDR1, LCDR2 and LCDR3).

[0040] The term "high affinity" antibody refers to an antibody that has a K_D with respect to its target epitope about of 10^{-9} M or lower (e.g., about 1×10^{-9} M, 1×10^{-10} M, 1×10^{-11} M, or about 1×10^{-12} M).

[0041] The phrase "bispecific antibody" includes an antibody capable of selectively binding two epitopes. Bispecific antibodies generally comprise two arms, each binding a different epitope (e.g., two heavy chains with different specificities)-either on two different molecules (e.g., different epitopes on two different immunogens) or on the same molecule (e.g., different epitopes on the same immunogen). If a bispecific antibody is capable of selectively binding two different epitopes (a first epitope and a second epitope), the affinity of the first antibody arm for the first epitope will generally be at least one to two or three or four or more orders of magnitude lower than the affinity of the first antibody arm for the second epitope, and vice versa. Epitopes specifically bound by the bispecific antibody can be on the same or a different target (e.g., on the same or a different protein). Exemplary bispecific antibodies include those with a first antibody arm specific for a tumor antigen and a second antibody arm specific for a cytotoxic marker, e.g., an Fc receptor (e.g., $Fc\gamma RI$, $Fc\gamma RII$, $Fc\gamma RIII$, etc.) or a T cell marker (e.g., $CD3$, $CD28$, etc.). In one embodiment of the present invention, one arm of the bispecific antibody is specific for $CD3$. Further, a bispecific antibody with a first arm specific for a tumor antigen and a second arm specific for a toxin can be paired so as to deliver a toxin (e.g., saporin, vinca alkaloid, etc.) to a tumor cell. Other exemplary bispecific antibodies include those with a first arm specific for an activating receptor (e.g., B cell receptor, $Fc\gamma RI$, $Fc\gamma RIIA$, $Fc\gamma RIIIA$, $Fc\gamma RI$, T cell receptor, etc.) and a second arm specific for an inhibitory receptor (e.g., $Fc\gamma RIIB$, $CD5$, $CD22$, $CD72$, $CD300a$, etc.). Such bispecific antibodies can be constructed for therapeutic conditions associated with cell activation (e.g., allergy and asthma). Bispecific antibodies can be made, for example, by combining heavy chains that recognize different epitopes of the same immunogen. For example, nucleic acid sequences encoding heavy chain variable sequences that recognize different epitopes of the same immunogen can be fused to nucleic acid sequences encoding the same or different heavy chain constant regions, and such sequences can be expressed in a cell that expresses an immunoglobulin light chain. A typical bispecific antibody has two heavy chains each having three heavy chain CDRs, followed by (N-terminal to C-terminal) a C_H1 domain, a hinge, a C_H2 domain, and a C_H3 domain, and an immunoglobulin light chain that either does not confer epitope-binding specificity but that can associate with each heavy chain, or that can associate with each heavy chain and that can bind one or more of the epitopes bound by the heavy chain epitope-binding regions, or that can associate with each heavy chain and enable binding of one or both of the heavy chains to one or both epitopes. Similarly, the phrase "multispecific antibody" includes an antibody capable of selectively binding multiple epitopes (e.g., two, three, four epitopes).

[0042] The phrase "complementarity determining region," or the term "CDR," includes an amino acid sequence encoded by a nucleic acid sequence of an organism's immunoglobulin genes that normally (i.e., in a wild-type animal) appears between two framework regions in a variable region of a light or a heavy chain of an immunoglobulin molecule. A CDR can be encoded by, for example, a germline sequence or a rearranged or unrearranged sequence, and, for example, by a naive or a mature B cell. A CDR can be somatically mutated (e.g., vary from a sequence encoded in an animal's germline), humanized, and/or modified with amino acid substitutions, additions, or deletions. In some circumstances (e.g., for a CDR3), CDRs can be encoded by two or more sequences (e.g., germline sequences) that are not contiguous (e.g., in an unrearranged nucleic acid sequence) but are contiguous in a B cell nucleic acid sequence, e.g., as the result of splicing or connecting the sequences (e.g., V-D-J recombination to form a heavy chain CDR3).

[0043] The phrase "functional fragment" includes fragments of antigen-binding proteins such as antibodies that can be expressed, secreted, and specifically bind to an epitope with a K_D in the micromolar, nanomolar, or picomolar range. Specific recognition includes having a K_D that is at least in the micromolar range, the nanomolar range, or the picomolar range.

[0044] The phrase "heavy chain," or "immunoglobulin heavy chain" includes an immunoglobulin heavy chain sequence, including immunoglobulin heavy chain constant region sequence, from any organism. Heavy chain variable domains include three heavy chain CDRs and four FR regions, unless otherwise specified. Fragments of heavy chains include

CDRs, CDRs and FRs, and combinations thereof. A typical heavy chain has, following the variable domain (from N-terminal to C-terminal), a C_H1 domain, a hinge, a C_H2 domain, and a C_H3 domain. A functional fragment of a heavy chain includes a fragment that is capable of specifically recognizing an epitope (e.g., recognizing the epitope with a K_D in the micromolar, nanomolar, or picomolar range), that is capable of expressing and secreting from a cell, and that comprises at least one CDR. A heavy chain variable domain is encoded by a variable region gene sequence, which generally comprises V_H, D_H, and J_H segments derived from a repertoire of V_H, D_H, and J_H segments present in the germline. Sequences, locations and nomenclature for V, D, and J heavy chain segments for various organisms can be found on the website for the International Immunogenetics Information System (IMGT database).

[0045] The phrase "light chain" includes an immunoglobulin light chain sequence from any organism, and unless otherwise specified includes human kappa and lambda light chains and a VpreB, as well as surrogate light chains. Light chain variable domains typically include three light chain CDRs and four framework (FR) regions, unless otherwise specified. Generally, a full-length light chain includes, from amino terminus to carboxyl terminus, a variable domain that includes FR1-CDR1-FR2-CDR2-FR3-CDR3-FR4, and a light chain constant region. A light chain variable domain is encoded by a light chain variable region gene sequence, which generally comprises V_L and J_L segments, derived from a repertoire of V and J segments present in the germline. Sequences, locations and nomenclature for V and J light chain segments for various organisms can be found on the website for the International Immunogenetics Information System (IMGT database). Light chains include those, e.g., that do not selectively bind any epitopes recognized by antigen-binding protein (e.g., antibody) in which they appear. Light chains also include those that bind and recognize, or assist the heavy chain with binding and recognizing, one or more epitopes selectively bound by the antigen-binding protein (e.g., an antibody) in which they appear.

[0046] The term "antigen-binding protein" as used herein includes antibodies and various naturally produced and engineered molecules capable of binding the antigen of interest. Such include, e.g., domain-specific antibodies, single domain antibodies (e.g., derived from camelids and fish, etc.), domain-deleted antibodies, chimeric antibodies, CDR-grafted antibodies, diabodies, triabodies, tetrabodies, minibodies, nanabodies (e.g., monovalent nanobodies, bivalent nanobodies, etc.), small modular immunopharmaceuticals (SMIPs), shark variable IgNAR domains, etc. Antigen-binding protein may also include antigen-binding fragments such as, e.g., (i) Fab fragments; (ii) F(ab')2 fragments; (iii) Fd fragments; (iv) Fv fragments; (v) single-chain Fv (scFv) molecules; (vi) dAb fragments; and (vii) minimal recognition units consisting of the amino acid residues that mimic the hypervariable region of an antibody (e.g., an isolated complementarity determining region (CDR) such as a CDR3 peptide), etc.

[0047] The term "cell" includes any cell that is suitable for expressing a recombinant nucleic acid sequence. Cells include those of prokaryotes and eukaryotes (single-cell or multiple-cell), bacterial cells (e.g., strains of *E. coli*, *Bacillus* spp., *Streptomyces* spp., etc.), mycobacteria cells, fungal cells, yeast cells (e.g., *S. cerevisiae*, *S. pombe*, *P. pastoris*, *P. methanolica*, etc.), plant cells, insect cells (e.g., SF-9, SF-21, baculovirus-infected insect cells, *Trichoplusia ni*, etc.), non-human animal cells, human cells, or cell fusions such as, for example, hybridomas or quadromas. In some embodiments, the cell is a human, monkey, ape, hamster, rat, or mouse cell. In some embodiments, the cell is eukaryotic and is selected from the following cells: CHO (e.g., CHO K1, DXB-11 CHO, Veggie-CHO), COS (e.g., COS-7), retinal cell, Vero, CV1, kidney (e.g., HEK293, 293 EBNA, MSR 293, MDCK, HaK, BHK), HeLa, HepG2, WI38, MRC 5, Colo205, HB 8065, HL-60, (e.g., BHK21), Jurkat, Daudi, A431 (epidermal), CV-1, U937, 3T3, L cell, C127 cell, SP2/0, NS-0, MMT 060562, Sertoli cell, BRL 3A cell, HT1080 cell, myeloma cell, tumor cell, and a cell line derived from an aforementioned cell. In some embodiments, the cell comprises one or more viral genes, e.g. a retinal cell that expresses a viral gene (e.g., a PER.C6™ cell). In some embodiments, the cell is an ES cell.

[0048] A humanized CD3 protein means a CD3 protein in which, in one embodiment, an extracellular domain is of human sequence. The transmembrane and cytoplasmic domains can also be human but are preferably non-human endogenous sequences. A CD3 protein including sequences from different species, particularly a human extracellular domain, and non-human transmembrane and cytoplasmic domains, can also be referred to as a chimeric CD3 protein.

Genetically Modified Humanized CD3 Animals

[0049] Disclosed herein are genetically modified non-human animals (e.g., rodents, e.g., mice or rats) that comprise in their genome (e.g., in their germline genome) a nucleic acid sequence encoding a humanized CD3 protein (e.g., a humanized CD3 ϵ , CD3 γ , CD3 δ , or combination thereof). The present invention provides genetically modified mice that comprise in their genome nucleotide sequences encoding humanized CD3 δ , humanized CD3 γ , and humanized CD3 ϵ proteins. Thus, in some embodiments of the invention, the mouse expresses a humanized CD3 $\gamma\delta\epsilon$ complex on the surface of its T cells such that the humanized CD3 $\gamma\delta\epsilon$ forms a complex with the T cell receptor expressed on the same T cell.

[0050] CD3 molecule is commonly a target of agents that are aimed at modulating T cell immunity, and several anti-CD3 antibodies have been developed for that purpose (e.g., muromonab-CD3 or OKT3). Anti-CD3 antibodies such as OKT3 are used as immunosuppressive agents (e.g., in transplant rejection) but are also studied for their therapeutic potential in autoimmune diseases (e.g., Crohn's disease, type I diabetes, ulcerative colitis, etc.).

[0051] Additionally, CD3 molecules are also being studied as targets for bispecific agents, e.g., bispecific antibodies, because of the ability of anti-CD3 bispecific antibodies to recruit T cells to a target cell, e.g., a cell that expresses a particular antigen of interest. Exemplary anti-CD3 bispecific antibodies are described in U.S. Patent Application Publication No. 2014/0088295, and US. Patent Application Publication No. 2015/0266966.

5 **[0052]** During preclinical drug development stage, candidate agents are typically studied based on their efficacy, toxicity, and other pharmacokinetic and pharmacodynamics properties. Candidate agents, such as antibodies, typically target a human antigen -- as the end goal of investigation is to develop a human therapy. Many preclinical studies are conducted in large animals such as primates as their physiology and drug metabolism are most similar to humans. Several antibodies developed to CD3 (e.g., OKT3) are known not to cross-react to non-human CD3, particularly to 10 primate CD3. To conduct effective preclinical investigations relating to efficacy, toxicity, and other parameters of a drug candidate, first, the drug candidate must be determined to recognize primate CD3 molecule.

15 **[0053]** However, a separate factor complicating development of anti-CD3 therapy is that large primates such as chimpanzees are endangered and in many countries studies in chimpanzees are prohibited; while studies in other primates, e.g., cynomolgus monkeys (*Macaca fascicularis*), may raise ethical concerns. For example, for all of the above reasons, to date there is no effective primate model of human tumors. Thus, any preliminary data on a specific therapeutic candidate that can be obtained in a smaller animal model, such as a rodent, e.g., a mouse, can be helpful in determining further progress of preclinical investigations in large primates.

20 **[0054]** Preclinical studies in small animal models, such as mice, have traditionally been conducted using drug surrogates. For example, when a clinical candidate targeting a specific human antigen is in development, some preclinical data is generated in a mouse using a molecule, e.g., an antigen binding protein or an antibody, which specifically targets a mouse homolog of the antigen of interest. Information about efficacy, various dosing regimens, toxicity and side effects, and other aspects of drug administration is gathered from such drug surrogate studies. However, such findings are limited because it is not the actual drug that is in development or its human target that is being studied.

25 **[0055]** Thus, the most useful small animal model to conduct preliminary preclinical studies is a non-human animal, e.g., a rodent, that expresses a human or humanized CD3 protein, and allows the testing of anti-CD3 drug candidates that also cross-react with cynomolgus monkey CD3, allowing for subsequent primate preclinical studies. The present invention provides such an intricate mouse model.

30 **[0056]** Thus, provided herein is a genetically modified non-human animal comprising in its genome a nucleic acid sequence(s) encoding an extracellular domain of a human CD3 protein. In some examples, the CD3 protein is selected from the group consisting of CD3 γ , CD3 δ , CD3 ϵ , CD3 ζ , and a combination thereof. In some examples, the CD3 protein is selected from the group consisting of CD3 γ , CD3 δ , CD3 ϵ , and a combination thereof. In some embodiments, the CD3 protein comprises CD3 γ , CD3 δ , and CD3 ϵ polypeptide chains. Thus, the genetically modified mouse comprises in its genome a nucleic acid sequence(s) encoding an extracellular domain of a human CD3 γ , an extracellular domain of a human CD3 δ , and an extracellular domain of a human CD3 ϵ . In some such embodiments, the extracellular domains of human CD3 γ , CD3 δ , and CD3 ϵ may be encoded by a single nucleic acid. In some embodiments, the extracellular domains of human CD3 γ , CD3 δ , and CD3 ϵ are encoded by separate nucleic acids.

35 **[0057]** In some embodiments, the mouse described herein retains endogenous nonhuman CD3 promoter(s) and/or regulatory elements (e.g., endogenous nonhuman CD3 γ , CD3 δ , and/or CD3 ϵ promoters and/or regulatory elements). In other embodiments, the mouse comprises human CD3 promoter(s) and regulatory elements.

40 **[0058]** Although it has been postulated that the majority of antibodies generated against CD3 recognize CD3 ϵ epitopes (see, Tunnicliffe et al. (1989) International Immunology, 1(5):546-50), there are a number of agents that may either recognize other CD3 subunits (e.g., CD3 γ or CD3 δ) or require assembly of the CD3 complex for binding. Thus, the genetically modified mouse that comprises in its genome a nucleic acid sequence(s) encoding an extracellular domain of a human CD3 γ , an extracellular domain of a human CD3 δ , and an extracellular domain of a human CD3 ϵ , provides an advantage since it can accommodate an agent that would bind any of the CD3 subunits or the CD3 complex.

45 **[0059]** Exemplary CD3 proteins are presented in the alignment in FIG. 4. A mouse CD3 ϵ protein sequence can be found in GenBank Accession Number NP_031674 and SEQ ID NO:27, while a human CD3 ϵ protein sequence can be found in GenBank Accession Number NP_000724 and SEQ ID NO:28. A mouse CD3 δ protein sequence can be found in GenBank Accession Number NP_038515 and SEQ ID NO:29, while a human CD δ protein sequence can be found in GenBank Accession Number NP_000723 and SEQ ID NO:30. A mouse CD3 γ protein sequence can be found in GenBank Accession Number NP_033980 and SEQ ID NO:31, while a human CD3 γ protein sequence can be found in GenBank Accession Number NP_000064 and SEQ ID NO:32.

50 **[0060]** In the invention, the nucleic acid sequence(s) encoding an extracellular domain of a human CD3, e.g., an extracellular domain of a human CD3 γ , human CD3 δ , and human CD3 ϵ , are located at an endogenous mouse CD3 locus. In other words, such a nucleic acid modifies the endogenous CD3 locus to encode the human CD3 polypeptide. In some embodiments of the invention, the mouse does not comprise a functional extracellular domain of the corresponding mouse CD3 protein because of genetic modification of the endogenous locus so that the functional extracellular domain is not expressed. In some embodiments of the invention, the nucleic acid sequence(s) encoding an extracellular

domain of a human CD3 replaces corresponding nucleic acid sequence(s) encoding endogenous mouse CD3. Thus, in some embodiments, the nucleic acid sequence encoding the extracellular domain of a human CD3 γ replaces the nucleic acid sequence encoding the extracellular domain of endogenous mouse CD3 γ , the nucleic acid sequence encoding the extracellular domain of a human CD3 δ replaces the nucleic acid sequence encoding the extracellular domain of endogenous mouse CD3 δ , and the nucleic acid sequence encoding the extracellular domain of a human CD3 ϵ replaces the nucleic acid sequence encoding the extracellular domain of endogenous mouse CD3 ϵ . In some embodiments, the replacement does not comprise the replacement of a nucleic acid sequence encoding endogenous signal sequence. In another embodiment, the replacement comprises the replacement of the nucleic acid sequence encoding endogenous signal sequence with the nucleic acid sequence encoding a human signal sequence.

[0061] In some aspects of the invention, the extracellular domain comprises the region of the protein(s) that is not a transmembrane or a cytoplasmic domain, e.g., the region of the protein that appears on the surface of the cell and that, in part, when assembled in a complex interacts with the extracellular domains of other components of TCR signaling complex, e.g., TCR alpha and beta extracellular domains. In various embodiments described herein, extracellular domain refers to the domain of the protein expressed on the cell surface and, unless indicated otherwise, does not include the signal sequence which is typically proteolytically cleaved prior to cell surface expression. In some embodiments of the invention, the extracellular domain of CD3 ϵ comprises amino acids 17-130 of the amino acid sequence set forth in SEQ ID NO:24 (set forth separately as SEQ ID NO:33). In some such embodiments, the animal comprises the nucleic acid sequence encoding an endogenous CD3 ϵ signal sequence, e.g., signal sequence at amino acids 1-16 of SEQ ID NO:24. In other embodiments of the invention, the animal comprises the nucleic acid sequence encoding a human CD3 ϵ signal sequence. In some embodiments of the invention, the extracellular domain of CD3 δ comprises amino acids 19-105 of the amino acid sequence set forth in SEQ ID NO:25 (set forth separately as SEQ ID NO:34). In some such embodiments, the mouse comprises the nucleic acid sequence encoding an endogenous CD3 δ signal sequence, e.g., signal sequence at amino acids 1-18 of SEQ ID NO:25. In other embodiments of the invention, the mouse comprises the nucleic acid sequence encoding a human CD3 δ signal sequence. In some embodiments, the extracellular domain of CD3 γ comprises amino acids 20-116 of the amino acid sequence set forth in SEQ ID NO:26 (set forth separately as SEQ ID NO:35). In some such embodiments, the mouse comprises the nucleic acid sequence encoding endogenous CD3 γ signal sequence, e.g., signal sequence at amino acids 1-19 of SEQ ID NO:26. In other embodiments of the invention, the mouse comprises the nucleic acid sequence encoding a human CD3 γ signal sequence.

[0062] In some aspects of the invention, the mouse comprises a nucleic acid sequence encoding transmembrane and cytoplasmic domains of endogenous CD3 protein, e.g., corresponding endogenous CD3 protein. Thus, in one embodiment, the mouse comprises a nucleic acid sequence encoding the extracellular domain of the human CD3 protein operably linked to the nucleic acid sequence encoding transmembrane and cytoplasmic domains of the corresponding endogenous mouse CD3 protein so that a chimeric protein comprising the extracellular domain of the human CD3 protein and the transmembrane and cytoplasmic domains of the corresponding endogenous mouse CD3 protein is expressed. Thus, in one aspect, the animal comprises at an endogenous CD3 locus a nucleic acid sequence(s) encoding an extracellular domain of a human CD3 protein operably linked to a nucleic acid sequence(s) encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3. In one embodiment, the mouse comprises at an endogenous CD3 ϵ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 ϵ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 ϵ , at an endogenous CD3 δ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 δ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 δ , and at an endogenous CD3 γ locus a nucleic acid sequence encoding an extracellular domain of a human CD3 γ operably linked to a nucleic acid sequence encoding transmembrane and cytoplasmic domains of an endogenous mouse CD3 γ . Use of chimeric CD3 proteins with a human extracellular domain and endogenous transmembrane and cytoplasmic domains allows for interaction of drugs with specificity for human CD3 but may also allow to recapitulate the interaction with the endogenous T-cell receptor and its signal transduction components compared with a fully human CD3 protein.

[0063] The mouse expresses extracellular domains of human CD3 protein. In some aspects, the mouse expresses an extracellular domain of human CD3 ϵ set forth in SEQ ID NO:33. In some aspects, the mouse expresses an extracellular domain of human CD3 δ set forth in SEQ ID NO:34. In some aspects, the mouse expresses an extracellular domain of human CD3 γ set forth in SEQ ID NO:35.

[0064] The non-human animal is a mouse.

[0065] In one embodiment, the mouse is of a C57BL strain selected from C57BL/A, C57BL/An, C57BL/GrFa, C57BL/Ka-LwN, C57BL/6, C57BL/6J, C57BL/6ByJ, C57BL/6N, C57BL/10, C57BL/10ScSn, C57BL/10Cr, and C57BL/Ola. In another embodiment, the mouse is a 129 strain selected from the group consisting of a strain that is 129P1, 129P2, 129P3, 129X1, 129S1 (e.g., 129S1/SV, 129S1/SvIm), 129S2, 129S4, 129S5, 129S9/SvEvH, 129S6 (129/SvEvTac), 129S7, 129S8, 129T1, 129T2 (see, e.g., Festing et al. (1999) Revised nomenclature for strain 129 mice, Mammalian Genome 10:836, see also, Auerbach et al (2000) Establishment and Chimera Analysis of 129/SvEv- and C57BL/6-Derived Mouse Embryonic Stem Cell Lines). In a specific embodiment, the genetically modified mouse is a mix of an aforementioned

129 strain and an aforementioned C57BL/6 strain. In another specific embodiment, the mouse is a mix of aforementioned 129 strains, or a mix of aforementioned BL/6 strains. In a specific embodiment, the 129 strain of the mix is a 129S6 (129/SvEvTac) strain. In another embodiment, the mouse is a BALB strain, e.g., BALB/c strain. In yet another embodiment, the mouse is a mix of a BALB strain and another aforementioned strain.

5 [0066] The non-human animal is a mouse. Thus, the genetically modified animal is a mouse and the mouse comprises at an endogenous mouse CD3 locus a nucleotide sequence encoding an extracellular domain of a human CD3 protein. The mouse comprises a nucleic acid sequence encoding an extracellular domain of a human CD3 ϵ , a nucleic acid sequence encoding an extracellular domain of a human CD3 δ , and a nucleic acid sequence encoding an extracellular domain of a human CD3 γ . In some embodiments of the invention, the extracellular domain of the human CD3 ϵ comprises the sequence set forth in SEQ ID NO:33, the extracellular domain of the human CD3 δ comprises the sequence set forth in SEQ ID NO:34, and the extracellular domain of the human CD3 γ comprises the sequence set forth in SEQ ID NO:35. In some embodiments, the mouse comprises the sequence(s) encoding endogenous mouse CD3 signal sequence(s). In other embodiments, the mouse comprises the sequence(s) encoding human CD3 signal sequence(s).

10 [0067] The mouse of the invention expresses humanized CD3 protein(s). The mouse expresses humanized CD3 ϵ , humanized CD3 δ , and humanized CD3 γ proteins. In some embodiments of the invention, the mouse expresses a human CD3 ϵ extracellular domain and endogenous mouse CD3 ϵ transmembrane and cytoplasmic domains, a human CD3 δ extracellular domain and endogenous mouse CD3 δ transmembrane and cytoplasmic domains, and a human CD3 γ extracellular domain and endogenous mouse CD3 γ transmembrane and cytoplasmic domains. In some such embodiments, the mouse expresses humanized CD3 proteins wherein the humanized CD3 proteins are humanized CD3 ϵ set forth in SEQ ID NO:24, humanized CD3 δ set forth in SEQ ID NO:25, and humanized CD3 γ set forth in SEQ ID NO:26.

15 [0068] In some aspects of the invention, the genetically engineered mouse is an immunocompetent mouse. In some embodiments of the invention, the introduction of humanized CD3 protein(s) does not affect the mouse's immune system function. In some embodiments of the invention, the mouse comprises normal T and B cell ratio. In some embodiments of the invention, the mouse is capable of mounting a normal response to mouse infection. In some aspects, the mouse displays similar CD4+ to CD8+ cell ratio in the thymus as compared to a wild type mouse, e.g., a mouse that has not been genetically modified to express humanized CD3 protein(s). In some embodiments of the invention, the CD4+ to CD8+ cell ratio in the thymus of the mouse is within 30%, e.g., within 20%, e.g., within 15%, e.g., within 12%, e.g., within 10%, e.g., within 5%, e.g., within 2%, of the CD4+ to CD8+ cell ratio of a mouse that is not genetically modified to express humanized CD3 protein(s). In some aspects, the mouse displays similar T and B cell percentages in the spleen, lymph nodes, and peripheral blood as a wild type mouse, e.g., a mouse that is not genetically modified to express humanized CD3 protein(s). In some aspects, the mouse displays similar numbers of circulating white blood cells, lymphocytes, monocytes, neutrophils, eosinophils, and basophils as a wild type mouse, e.g., a mouse that is not genetically modified to express humanized CD3 protein(s).

20 [0069] Also provided herein are methods of making the genetically modified mouse described herein. In some examples, the method of making a genetically modified mouse wherein the animal expresses a humanized CD3 protein comprises introducing at an endogenous non-human animal CD3 locus a nucleic acid sequence encoding an extracellular domain of a human CD3 protein, wherein the human CD3 protein is selected from the group consisting of CD3 ϵ , CD3 δ , CD3 γ , CD3 ζ , and a combination thereof. If multiple human CD3 proteins are introduced, they can be introduced together on a single nucleic acid (as in the present Examples) or separately. If the latter, a single cell line (e.g., ES cell line) can undergo successive modifications until modified to include nucleic acids encoding each of the desired human CD3s. In one embodiment the animal does not comprise a functional extracellular domain of the corresponding non-human CD3 protein(s). The mouse comprises at an endogenous mouse CD3 locus a nucleic acid sequence encoding an extracellular domain of human CD3 ϵ , an extracellular domain of human CD3 δ , and an extracellular domain of human CD3 γ . In some embodiments, the extracellular domain of a human CD3 ϵ is set forth in SEQ ID NO:33, the extracellular domain of a human CD3 δ is set forth in SEQ ID NO:34, and the extracellular domain of a human CD3 γ is set forth in SEQ ID NO:35. In one embodiment, the animal does not comprise a functional extracellular domain of the corresponding non-human CD3 protein(s).

25 [0070] In some embodiments, the method of making a genetically modified mouse mouse of the invention comprises replacing at the endogenous CD3 locus a nucleotide sequence encoding the extracellular domain of a mouse CD3 protein(s) with a nucleotide sequence encoding an extracellular domain of a corresponding human CD3 protein(s). In one embodiment, the animal retains transmembrane and cytoplasmic domains of the mouse CD3 protein(s). In some embodiments, the replacement results in a chimeric protein(s) comprising an extracellular domain of a human CD3 protein(s) and transmembrane and cytoplasmic domains of corresponding endogenous mouse CD3 protein(s).

30 [0071] Nucleic acid(s) encoding human CD3 protein(s) are typically introduced into a cell, and a mouse is propagated from the cell. In some embodiments, the replacement method utilizes a targeting construct made using VELOCIGENE® technology, introducing the construct into ES cells, and introducing targeted ES cell clones into a mouse embryo using VELOCIMOUSE® technology, as described in the Examples.

35 [0072] In one embodiment, wherein the method comprises the replacement of the nucleotide sequence encoding the

extracellular domain of endogenous mouse CD3 ϵ with the nucleotide sequence encoding the extracellular domain of a human CD3 ϵ protein, the method comprises a replacement of partial sequence of endogenous mouse coding exons 2 to 4 of mouse CD3 ϵ gene with partial sequence of human coding exons 2 to 5 of human CD3 ϵ gene. In one embodiment, wherein the method comprises the replacement of the nucleotide sequence encoding the extracellular domain of endogenous mouse CD3 δ with the nucleotide sequence encoding the extracellular domain of a human CD3 δ , the method comprises a replacement of partial sequence of endogenous mouse coding exons 2 to 3 of mouse CD3 δ with the partial sequence of human coding exons 2 to 3 of human CD3 δ gene. In one embodiment, wherein the method comprises a replacement of the nucleotide sequence encoding the extracellular domain of endogenous mouse CD3 γ with the nucleotide sequence encoding the extracellular domain of human CD3 γ , the method comprises replacement of partial sequence of mouse coding exons 2 to 4 of mouse CD3 γ with the partial sequence of human coding exons 2 to 4 of human CD3 γ gene. In one embodiment of the invention, the replacement comprises the replacement of sequence of CD3 ϵ , CD3 δ , and CD3 γ . In such an embodiment, the replacement may be accomplished by creating a large targeting vector that incorporates the sequential genetic modification in all three loci and then introducing the large targeting vector into mouse ES cells to make a mouse, e.g., as described in Example 1.

[0073] Thus, disclosed herein is a large targeting vector for making a genetically modified animal of the invention. In one example, the large targeting vector comprises 5' and 3' mouse homology arms; a DNA fragment comprising the CD3 ϵ gene which comprises a replacement of partial sequence of mouse CD3 ϵ coding exons 2 to 4 with partial sequence of human CD3 ϵ coding exons 2 to 5; a DNA fragment comprising the CD3 δ gene which comprises a replacement of partial sequence of mouse CD3 δ coding exons 2 to 3 with partial sequence of human CD3 δ coding exons 2 to 3; a DNA fragment comprising the CD3 γ gene which comprises a replacement of partial sequence of mouse CD3 γ coding exons 2 to 4 with partial sequence of human CD3 γ coding exons 2 to 4; and a selection cassette.

[0074] A selection cassette is a nucleotide sequence inserted into a targeting construct to facilitate selection of cells (e.g., bacterial cells, ES cells) that have integrated the construct of interest. A number of suitable selection cassettes are known in the art (Neo, Hyg, Pur, CM, SPEC, etc.). In addition, a selection cassette may be flanked by recombination sites, which allow deletion of the selection cassette upon treatment with recombinase enzymes. Commonly used recombination sites are *loxP* and *Fr*t, recognized by Cre and Flp enzymes, respectively, but others are known in the art. A selection cassette may be located anywhere in the construct outside the coding region. In one example, the selection cassette is inserted upstream of human CD3 ϵ inserted sequence.

[0075] Upon completion of gene targeting, ES cells or genetically modified mice are screened to confirm successful incorporation of exogenous nucleotide sequence of interest or expression of exogenous polypeptide. Numerous techniques are known to those skilled in the art, and include (but are not limited to) Southern blotting, long PCR, quantitative PCR (e.g., real-time PCR using TAQMAN®), fluorescence *in situ* hybridization, Northern blotting, flow cytometry, Western analysis, immunocytochemistry, immunohistochemistry, etc. In one example, non-human animals (e.g., mice) bearing the genetic modification of interest can be identified by screening for loss of mouse allele and/or gain of human allele using a modification of allele assay described in Valenzuela et al. (2003) High-throughput engineering of the mouse genome coupled with high-resolution expression analysis, *Nature Biotech.* 21(6):652-659. Other assays that identify a specific nucleotide or amino acid sequence in the genetically modified animals are known to those skilled in the art.

[0076] Heterozygotes resulting from the above methods can be bred to generate homozygotes.

[0077] A method for making a chimeric human/non-human CD3 molecule is disclosed, comprising expressing in a single cell a chimeric CD3 protein from a nucleotide construct as described herein. In one example, the nucleotide construct is a viral vector; in a specific example, the viral vector is a lentiviral vector. In one example, the cell is selected from a CHO, COS, 293, HeLa, and a retinal cell expressing a viral nucleic acid sequence (e.g., a PERC.6™ cell).

[0078] A cell that expresses a chimeric human/non-human CD3 protein is disclosed. In one example, the cell comprises an expression vector comprising a chimeric CD3 sequence as described herein. In one example, the cell is selected from CHO, COS, 293, HeLa, and a retinal cell expressing a viral nucleic acid sequence (e.g., a PERC.6™ cell).

[0079] A chimeric CD3 molecule made by a mouse as described herein is also disclosed, wherein, in one example, the chimeric CD3 molecule comprises an amino acid sequence of all or substantially all of an extracellular domain of a human CD3 protein, and at least transmembrane and cytoplasmic domains from a non-human CD3 protein, e.g., mouse CD3 protein.

[0080] In addition to a genetically engineered mouse a non-human embryo (e.g., a rodent, e.g., a mouse or a rat embryo) is also disclosed, wherein the embryo comprises a donor ES cell that is derived from a non-human animal (e.g., a example, rodent, e.g., a mouse or a rat) as described herein. In one example, the embryo comprises an ES donor cell that comprises the chimeric CD3 gene, and host embryo cells.

[0081] Also disclosed is a tissue, wherein the tissue is derived from a non-human animal (e.g., a rodent, e.g., a mouse or a rat) as described herein, and expresses the chimeric CD3 protein.

[0082] In addition, a non-human cell isolated from a non-human animal as described herein is disclosed. In one example, the cell is an ES cell. In one example, the cell is a T cell. In one example, the cell is a CD8+ T cell. In another example, the cell is a CD4+ T cell.

[0083] In some examples, also disclosed herein are genetic loci comprising the nucleic acid sequences that encoding the humanized CD3 protein(s) described herein.

Mouse Model for Testing Human Therapies

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[0084] In some aspects, provided herein is a mouse model for testing CD3-targeted ("anti-CD3") therapeutic agents according to claim 8. In some embodiments, provided herein is a mouse model for testing anti-CD3 antigen-binding proteins. In some embodiments, provided herein is a mouse model for testing anti-CD3 antibodies. In some such embodiments, provided is a mouse model for testing anti-CD3 multispecific, e.g. bispecific antigen-binding proteins or anti-CD3 bispecific antibodies. As such, an anti-CD3 multispecific antigen-binding protein, e.g. an anti-CD3 bispecific antigen-binding protein, targets or specifically binds said humanized CD3 protein and at least one other antigen of interest. In various aspects, the mouse model for testing anti-CD3 bispecific antigen-binding proteins wherein the antigen-binding protein is capable of binding both CD3 and the antigen of interest comprises a nucleic acid sequence encoding a humanized CD3 protein, wherein the humanized CD3 protein comprises CD3 ϵ , CD3 δ , CD3 γ , and a combination thereof, and a cell expressing or comprising the antigen of interest. The mouse comprises a T cell expressing said humanized CD3 protein(s).

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[0085] In an embodiment, the testing of the monospecific or bispecific antigen-binding protein involves performing an assay or a study that allows determination of the effect of the antigen-binding protein on the T cell expressing said humanized CD3 protein. In another embodiment, the testing of the bispecific antigen-binding protein involves performing an assay or a study that allows determination of the effect of the antigen-binding protein on both the T cell expressing said humanized CD3 protein and the cell expressing or comprising the antigen of interest, or the interaction between said CD3-expressing T cell and the cell expressing or comprising the antigen of interest. In one embodiment, the testing of the monospecific or bispecific antigen-binding protein involves performing an assay or a study that allows determination of the effect of the T cell expressing said humanized CD3 protein on the cell expressing or comprising said antigen of interest. In one embodiment, such assay measures, e.g., the number of cells expressing the antigen of interest, immune response, cellular interactions, cellular cytotoxicity, cytokine release, cellular activation, cell proliferation, tumor growth or regression, changes in pathology, or the like. Various assays include but are not limited to measurements of complement-directed cytotoxicity (CDC), antibody-dependent cytotoxicity (ADCC), antibody-dependent cellular phagocytosis (ADCP), PBMC proliferation, CD69 activation, histological tissue analysis, analysis of tissue and cellular biomarkers (e.g., cells or tissue may be extracted from the mouse for the purpose of the assays, or analyzed by radiography, MRI, PET, SPECT, BLI, and fluorescence-based imaging modalities).

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[0086] In some embodiments of the invention, in such a mouse model, the antigen of interest has been introduced into said mouse. The antigen of interest may be introduced by several methods known to those skilled in the art. Some nonlimiting methods include transgenesis, injection, infection, tissue or cell transplantation. The antigen of interest or a fragment thereof (e.g., a fragment that is recognized by the antigen-binding protein being tested) can be targeted to, or expressed by, particular cell types. In some embodiments, the antigen of interest is a humanized antigen of interest encoded by the mouse genome.

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[0087] The antigen of interest may be a membrane-bound protein such that it is expressed only on cell surface. Alternatively, the antigen of interest or a fragment thereof (e.g., a fragment that is recognized by the antigen-binding protein being tested) may be displayed on the cell surface complexed with another protein or moiety. Some cell-surface antigens may associate with other proteins as co-receptor complexes, or bind or have affinity to extracellular molecules. Thus, the mouse model may be utilized to test bispecific antigen-binding molecules that interact with T cells in various cell systems.

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[0088] The mouse model expresses human CD3 ϵ , CD3 δ , CD3 γ extracellular domains. In one embodiment, the mouse expresses mouse transmembrane and cytoplasmic domain of CD3 ϵ , CD3 δ , and CD3 γ ; an in one embodiment, the transmembrane and cytoplasmic domains are endogenous mouse domains. In one embodiment, the mouse model expresses CD3 ϵ , CD3 δ , and CD3 γ , each comprising a human extracellular domain and mouse, e.g., endogenous mouse, transmembrane and cytoplasmic domains.

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[0089] The antigen-binding protein binds both CD3 and the antigen of interest in the mouse model. In one embodiment, the antigen of interest is a human antigen. In one embodiment, the antigen of interest is a primate antigen, e.g., a cynomolgus monkey antigen. In one embodiment, the antigen-binding protein is capable of binding the same antigen of interest of both human and monkey origin. In one embodiment, the antigen-binding protein is capable of binding both human and monkey CD3.

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[0090] In one embodiment, the mouse model comprises a xenograft of a tumor expressing the antigen of interest. In one embodiment, the cell expressing or comprising the antigen of interest in said mouse is an immortalized cell, such as a tumor cell. Thus, the mouse model is utilized to test the activity of anti-CD3 bispecific antigen-binding proteins in blocking or affecting the tumor cell expressing the antigen of interest.

[0091] Thus, in the embodiment of the invention, wherein the cell expressing or comprising the antigen of interest is

a tumor cell, the antigen of interest may be a tumor-associated antigen (TAA). Various tumor antigens are listed in the database of T cell defined tumor antigens (van der Bruggen P, Stroobant V, Vigneron N, Van den Eynde B. Peptide database: T cell-defined tumor antigens. *Cancer Immun* 2013). Exemplary tumor associated antigens include but are not limited to ALK, BAGE proteins, BIRC5 (survivin), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, 5 CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, EGFR variant III, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, GAGE proteins, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5, EBV-derived LMP2, L1CAM, MAGE proteins, MLANA, MSLN, 10 MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), RAGE proteins, RET, RGS5, ROR1, SART1, SART3, SLAMF7, 15 SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMPRSS2, Thompson-nouvelle antigen, TNFRSF17, TYR, UPK3A, VTCN1, WT1. In one example, as described in Example 3 herein, the antigen of interest may be CD20, e.g., human or humanized CD20.

[0092] In another embodiment of the invention, the mouse model is used to determine if a candidate bispecific antigen-binding protein is capable of blocking or affecting an antigen of interest which is an infectious disease associated antigen.

15 In one embodiment of the invention, the mouse is infected with an infectious agent. In one embodiment of the invention, the infectious disease associated antigen is a viral antigen. In one aspect, the viral antigen is selected from the group consisting of HIV, hepatitis A, hepatitis B, hepatitis C, herpes virus (e.g., HSV-1, HSV-2, CMV, HAV-6, VZV, Epstein Barr virus), adenovirus, influenza virus, flavivirus, echovirus, rhinovirus, coxsackie virus, coronavirus, respiratory syncytial virus, mumps virus, rotavirus, measles virus, rubella virus, parvovirus, vaccinia virus, HTLV, dengue virus, papillomavirus, 20 molluscum virus, poliovirus, rabies virus, JC virus, ebola virus, and arboviral encephalitis virus antigen.

[0093] In another embodiment of the invention, wherein the antigen of interest is an infectious disease associated antigen, the antigen of interest is a bacterial antigen. In some aspects of the invention, the bacterial antigen is selected from the group consisting of chlamydia, rickettsia, mycobacteria, staphylococci, streptococci, pneumonococci, meningococci, gonococci, klebsiella, proteus, serratia, pseudomonas, legionella, diphtheria, salmonella, bacilli, cholera, 25 tetanus, botulism, anthrax, plague, leptospira, and Lyme disease bacterial antigen.

[0094] In some aspects of the invention, the CD3-based bispecific antigen binding protein is a human CD3 based antigen binding protein. In one embodiment, the antigen binding protein is an antibody, e.g., a human antibody, or an antigen-binding fragment thereof.

30 **[0095]** In some embodiments of the invention, the mouse model is an immunocompetent mouse model. In some embodiments of the invention, the mouse model allows for testing of efficacy and/or toxicity of an antigen-binding protein of interest. The measures of efficacy will depend on the antigen of interest being targeted by the bispecific agent. In some embodiments, the measure of efficacy is T cell killing of the cell expressing the antigen. In other embodiments, the measure of efficacy is neutralization of the virus. In other embodiment, the measure of efficacy may be viability of the animal. In yet another embodiment, the measure of efficacy may be elimination of cells expressing the antigen of interest, proliferation of T cells, production of cytokines (e.g., IFNg, TNFa, IL-1, IL-2, IL-10, IL4, IL-6, granzyme, perforin, etc.)

35 **[0096]** In some embodiments of the invention, the toxicity in the animal may be measured as an adverse event in the animal, e.g., change in body weight, appetite, digestive changes, changes in blood cell counts, splenomegaly, histological changes of the organs, change in liver enzyme function, changes in urinalysis, organ toxicity, hemorrhage, dehydration, loss of fur and scruffiness, or other signs of morbidity. One measure may be determination of antigen-binding protein 40 cross-reactivity with irrelevant antigens, which, in one embodiment, can be detected by organ histology, specifically detection of antigen-binding protein in tissues or cell types that are not known to express the antigen of interest.

Use of Genetically Modified Non-Human Animals

45 **[0097]** The invention also provides various methods of using the genetically modified mice described herein.

[0098] In one embodiment, provided herein is a method of screening therapeutic drug candidates that target an antigen of interest comprising (a) providing or receiving a genetically modified mouse comprising at its endogenous mouse CD3 locus a nucleic acid sequence encoding an extracellular domain of a human CD3 ϵ , CD3 δ , and CD3 γ , (b) introducing 50 into said genetically modified mouse an antigen of interest, (c) contacting said mouse with a drug candidate of interest, wherein the drug candidate is directed against the human CD3 and the antigen of interest, and (d) determining if the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest. The mouse expresses a functional humanized CD3 protein on the surface of its T cells. The genetically modified mouse comprises at the endogenous mouse CD3 locus a nucleic acid sequence encoding an 55 extracellular domain of human CD3 ϵ , an extracellular domain of human CD3 δ , and an extracellular domain of human CD3 γ . In one embodiment of the method described herein, the mouse does not comprise the nucleic acid sequence encoding a functional extracellular domain of the corresponding mouse protein. In some embodiments of the method, the extracellular domain(s) of the human CD3 protein(s) is operably linked to the transmembrane and cytoplasmic

domain(s) of the corresponding endogenous mouse CD3 protein(s). In various such embodiments of the methods, the extracellular domain of a human CD3 ϵ is set forth in SEQ ID NO:33, the extracellular domain of a human CD3 δ is set forth in SEQ ID NO:34, and the extracellular domain of a human CD3 γ is set forth as SEQ ID NO:35. In various embodiment of the methods described here, the mouse may express a humanized CD3 ϵ protein set forth in SEQ ID NO:24, a humanized CD3 δ protein set forth in SEQ ID NO:25, and a humanized CD3 γ set forth in SEQ ID NO:26.

[0099] In various embodiments of the method described herein, introduction of the antigen of interest into the genetically modified mouse described herein may be accomplished by any methods known to those skilled in the art, which may include, without limitation, transgenesis, injection, infection, tissue or cell transplantation. As such, introduction may be achieved by expressing in the mouse the antigen of interest, which can comprise genetically modifying said mouse to express the antigen of interest. Alternatively, introduction may comprise introduction into said mouse a cell expressing the antigen of interest, e.g., as in cell or tissue transplantation. Introduction may also comprise infecting said mouse with the antigen of interest, e.g., as in bacterial or viral infection. In one embodiment, the antigen of interest may be a human antigen of interest. In another embodiment, it may be a bacterial or a viral antigen of interest.

[0100] The antigen of interest may be a tumor-associated antigen, as described in detail above. The antigen may also be an infectious disease associated antigen, e.g., a bacterial or a viral antigen, as described in detail above.

[0101] In various embodiments of the methods of screening a therapeutic drug candidate, the drug candidate may be an antigen-binding protein, e.g., an antibody, e.g., a bispecific antibody. In various aspects, such drug candidate is capable of binding both human CD3 and the antigen of interest. The antigen of interest may be a human antigen. The antigen of interest may also be a primate, e.g., a monkey, antigen. Thus, the drug candidate used for screening may be capable of binding both a human antigen and a corresponding primate antigen, in addition to binding human CD3. The drug candidate may also be capable of binding primate, e.g., monkey, CD3. Thus, the drug candidate may be capable of binding both human and primate, e.g., monkey, CD3; and also, in one embodiment, be capable of binding a human antigen of interest. In another embodiment, the antigen of interest may be a bacterial or a viral antigen, and the drug candidate may be capable of binding both the human and primate, e.g., monkey, CD3 and the bacterial or viral antigen of interest.

[0102] In some aspects, the therapeutic candidate is an antibody, which is a human antibody. In other aspects, it may be a humanized antibody. For example, the therapeutic candidate may be an antibody generated in VELOCIMMUNE® mice (U.S. Patent Number 8,502,018, thus, the initial antibody candidate may comprise a human variable region and a mouse constant region. The mouse constant region of the antibody candidate may be reengineered to be of human origin by expressing the human variable region selected in VELOCIMMUNE® mice in operable linkage with a human constant region.

[0103] In various embodiments of the methods described herein, the therapeutic candidate is capable of reducing, eliminating, or preventing a disease. In one embodiment, the disease is a tumor, and the therapeutic candidate is capable of reducing, eliminating, or preventing tumor growth as compared to an agent that does not target the antigen of interest. In such an embodiment of the method, determination whether the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest can be performed using a tumor volume assay, a tumor cell killing assay, induction of apoptotic markers in tumors, reduction in blood vessel growth in tumors, infiltration of immune cells into tumors, etc. In another embodiment, the disease is an infectious disease, and a therapeutic candidate is capable reducing, eliminating, or preventing a bacterial or a viral infection as compared to an agent that does not target the antigen of interest. In such an embodiment of the method, determination whether the drug candidate is efficacious in preventing, reducing or eliminating cells characterized by the presence or expression of the antigen of interest can be performed using a measure of bacterial or viral titers, induction of apoptotic markers in infected cells, etc.

[0104] Other methods of use of the humanized CD3 mice of the present invention are also disclosed. For example, the non-human animal, e.g., a humanized CD3 mouse, disclosed herein may be used to study the mechanism of drug action. Prior to the development of the present animal, it was difficult to study the mechanism of drug action as such studies are not typically conducted in humans and primates, and often require an immunocompetent animal model. Understanding drug action mechanism can lead to development of better antibodies. In various embodiments of the invention, the humanized CD3 mouse is an immunocompetent mouse. For example, the humanized CD3 mouse of the invention, which comprises a healthy normal immune system with intact development and complete complement of all immune cell types and intact immune signaling pathways, can be used to study the effects of various therapeutic candidates on specific cell types, cytokines, chemokines, etc. The mouse can then be used to answer mechanistic questions relating to drug candidate function.

[0105] In addition, the humanized CD3 mice can be used in methods that involve testing the effect of bispecific anti-CD3 drug candidates on tumor grafts. Previously developed mouse models were immunocompromised mouse models to allow for proper human tumor engraftment. Humanized CD3 mouse is fully immunocompetent and allows introduction and growth of tumor cells expressing the antigen of interest, so full affect on the immune response can be studied, included but not limited to answering mechanistic questions, early toxicity questions, early efficacy questions, etc.

[0106] In yet other embodiments, the humanized CD3 mouse can be used to study the effects of combination drug therapies in animal models, specifically combination drug therapies, e.g., where one drug is an antigen-binding protein that binds CD3 and another drug is an agent that has previously been approved for a particular indication. Specific questions related to the dosing of the drugs and its effects can be addressed in an animal model prior to any human trials.

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EXAMPLES

[0107] The following examples are provided so as to describe to those of ordinary skill in the art how to make and use methods and compositions of the invention, and are not intended to limit the scope of what the inventors regard as their invention. Efforts have been made to ensure accuracy with respect to numbers used (e.g., amounts, temperature, etc.) but some experimental errors and deviations should be accounted for. The Examples do not include detailed descriptions of conventional methods that would be well known to those of ordinary skill in the art (molecular cloning techniques, etc.). Unless indicated otherwise, parts are parts by weight, molecular weight is average molecular weight, temperature is indicated in Celsius, and pressure is at or near atmospheric.

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Example 1. Construction of Humanized CD3 Locus

Example 1.1. Construction of Humanized CD3 $\gamma\delta\epsilon$

[0108] The mouse CD3 locus was humanized by construction of unique targeting vectors from human and mouse bacterial artificial chromosomes (BAC) DNA using VELOCIGENE® technology (see, e.g., US Patent No. 6,586,251 and Valenzuela et al. (2003) High-throughput engineering of the mouse genome couple with high-resolution expression analysis. *Nat. Biotech.* 21(6): 652-659). DNA from mouse BAC bMQ-425K11 was modified by homologous recombination to replace the genomic DNA encoding portions of mouse CD3 ϵ , CD3 δ , and CD3 γ genes (mouse CD3 genes located within close proximity to one another on chromosome 9) with corresponding portions of CD3 ϵ , CD3 δ , and CD3 γ genes derived from human BAC RP11-414G21 (human CD3 genes are located within close proximity to one another on chromosome 11), respectively.

[0109] Specifically, to generate humanized CD3 $\gamma\delta\epsilon$ mice, the mouse BAC was first modified by replacing 714 bp of mouse Cd3d sequence (corresponding to partial sequence of mouse coding exons 2-3 of Cd3d gene) with 939 bp of human CD3D sequence (corresponding to partial sequence of human coding exons 2-3 of CD3D gene) in a single targeting event using a targeting vector comprising a Spec cassette using mouse homology arms.

[0110] Mouse BAC comprising a replacement of partial sequence of mouse coding exons 2-3 of CD3d gene with corresponding human sequence was subsequently modified by replacement of 1,738bp of mouse Cd3g sequence (corresponding to partial sequence of mouse coding exons 2-4 of Cd3g gene) with 1,639 bp of human CD3G sequence (corresponding to partial sequence of human coding exons 2-4 of CD3G gene) also in a single targeting event using another Spec cassette-containing vector and mouse homology arms.

[0111] Finally, the BAC comprising the replacement of mouse CD3d and CD3g genes with corresponding human genes was further modified by replacing 6,213bp mouse CD3e sequence with 6,817bp of human sequence (corresponding to replacement of partial sequence of mouse coding exons 2 to 4 of mouse CD3e gene with partial sequence of human coding exons 2 to 5 of human CD3E gene). A 4,996bp floxed neomycin cassette was inserted upstream of human CD3E sequence knock-in.

[0112] The resulting humanized large targeting vector for insertion into ES cells is depicted in FIG. 2A, with A, B, C, D, E, F, and G indicating various mouse/human or mouse/NEO cassette or human/NEO cassette junctions. The sequences at the junctions are depicted in Table 1 below.

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Table 1. Junctional Sequences of the Large Targeting Vector

Sequence designation in FIG. 1	Junction	Sequence	SEQ ID NO:
5	A 5' mouse Cd3e/ Xhol/ (loxP) cassette	CGACTTCTTGA CTT CTATTGTTAACACTGTGCATTACATCGAATGCTAGAAGTTCCCTCGTCCCCTGCTCCTGAATTGCCCTGGATCCTCTGCTTGATGCCCTGTAGGAAACGTCCTTCTCTGGTATAGAAATGACTG/ CTC GAG/ATAAC TCGTATAATGTATGC TATACGAAGTTAT ATGCATGGCCTCGCGGCCGTTGGCCCTCCGC	1
10	B 3' cassette (loxP) Ice UI// human CD3E	TGTATCTTATCATGTCTGGAA ATAAC TTCGTATAATGTATGCTATACGAAG TTAT GCTAGTAACTATAACGGTCCTAAGGTAGCGAGCTAGC//CTTCCACAGACACCAATGTTAAAATGGAGGC TTGGGGGCAAAATTCTTTGCTATG TCTCTAGTCGCAAAAAATGGTCC TAACTTTTCTGACTCCTGCTTGTCAAAAATTGTGGGCTCATAGTTAATGC	2
15	C 3' human CD3E/ mouse Cd3e	AGGGGAGAATGGCCTTCATGCACTCCCTCCTACCTCCAGGGCTTGTGTTTTCCTTGCTTAGTGATTCCCCCTCTCCCAACCCCCACCCCCACAGTGTGTGAGAACTGCATGGAGATGGATGTGATGTCGGTG/ GCCATAATCATCATTGTTGAC ATCTGTATCACTCTGGCTTGCTGAT GGTCA TTTATTACTGGAGCAAGAATA GGAAGGCCAAGGCCAAGCCT	3
20	D 3' mouse Cd3d/ human CD3D	GAAAGAGAGAGTCTTCTGCTAACTAACCCCCAGAAGGCCTTCGGTCTCATGTCCTGCAAAGCAGTAGACGCCAAA GCCAGGAGCAGAGTTGCGATGAGGTC AATGAAGATGACACC/AGCCACGGTG GCTGGATCCAGCTCCACACAGCTCTG GCACACTGTGGGGGAAGGGAGGAGAGAGAGAGGTTGAGAGCCTTAAAGATCAGGAACCATCCT	4
25	E 5' human CD3D/ Sgr DI/ mouse Cd3d	CAAGAGAGACAGAAGTCACAAGAAAAAGCCTTCAGAAAGTCCCCACCAACTGCAGGGGTCAAGGGGGACATGAGGATGCCATTCAAG/ CGTCGACG /AGCGTAGGCAGCTTATTGCTCTGCATACTTACAGACCATTTGTGTAGTAAGGGACATGATGCCGAGTGA AA AGGGCAGGAGCAA CCAGAGGGAGATTTCAGGAAGTTCTCAGGGACTCGAGGTTCGTGA	5
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(continued)

Sequence designation in FIG. 1	Junction	Sequence	SEQ ID NO:
5	F 5' mouse Cd3g/ AsisI /human CD3G	GAAGCCCCACCCAGAAAGGTAGGACAA AGATCATAGTCATATTACTTCATCCA GGAGAGAAACACAGACACAGCCATTGC CTTGGCCATCATCTCTCCATCTTGA CCTCACGTGATCATG/ GCGATCGC/GA GTGATTAGTCTACAATCCGGAAA AAGTATAGATACTACCATTTCATGGA TTTGGATCTTCTTCATCTTGGCCTCA AATAACCATG	6
10	G 3' human CD3G/mouse Cd3g	GCATTATTGCAGACAGGCAGGACAAAA CGAACCCAGGAAAAACAACCTTCGCAAC CTGAAGGTTGTCTCTCCTTTCCCTA CAGTGTGTCAAGAACTGCATTGA ATGCAGGCCACCATACT/GGCTTATC TTCGCTGAGGTCA CTTGCTCTGGGTATATCTCATTGCG GGACAGGATGGACAATA ACCTGTCTTA A	7
15			
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[0113] The targeted BAC DNA was used to electroporate mouse ES cells comprising a deletion in mouse CD3 locus to create modified ES cells for generating mice that express humanized CD3 ϵ , CD3 δ , and CD3 γ on the surface of their T cells. ES cells containing insertions of human CD3 ϵ , CD3 δ , and CD3 γ sequences were identified by a quantitative TAAQMAN™ assay (see, e.g., Lie and Petropoulos, 1998. Curr. Opin. Biotechnology 9:43-48. Specific primer sets and probes were designed for detecting insertion of human sequences (gain-of-allele, GOA) and deletion of mouse sequences (loss-of-allele, LOA). Table 2 identifies the names and locations of each of primers/probe sets used in the quantitative PCR assays.

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Table 2: Primers/Probe Pairs Used for Genotyping

Gene	Sequence Name	Assay	Fwd Primer	Probe (BHQ)	Rev Primer
Mouse Cd3e	968 mTU	LOA	CCTCTGCCATG TAGGTTCTG TAC (SEQ ID NO:9)	TGCCGTGATGT TTGTTCAATGA CCAAA (SEQ ID NO:10)	GTTCCTGAG AAAGGGT TCTTAAGTG (SEQ ID NO:11)
Mouse Cc3g	7164 mTD	LOA	CCAGGGCTACT TGCTGTTCTG (SEQ ID NO:12)	TGGGCTTACCAT CCAGGGACGA (SEQ ID NO:13)	GCTACTCTTC CCACAAACTG CTTAG (SEQ ID NO:14)
Human CD3E	7170 hTU	GOA	CCAGCAGTAAG TTCCACTGTT TAG (SEQ ID NO:15)	TGTAGAAATGG CTGTGACCCAGCA (SEQ ID NO:16)	GGGCTGTGTT GCAGTATGAC (SEQ ID NO:17)
Human CD3D	928 hTU	GOA	ACCGTGCAAGT TCATTATCGAAG (SEQ ID NO:18)	ACGTGCTTCCGT AACCTTTGGGT (SEQ ID NO:19)	TCTCACATCCA GAAGCCCTATC (SEQ ID NO:20)
Human CD3G	7164 hTD	GOA	CGAGGGATGTA TCAGTGTAAAG GA (SEQ ID NO:21)	CACAGAACAAAGT CAAAACACTCC AAGTG (SEQ ID NO:22)	GCTCACCAAGA CAGCAAATACTG (SEQ ID NO:23)

[0114] Targeted ES cells described above were used as donor ES cells and introduced into an 8-cell stage mouse embryo by the VELOCIMOUSE® method (see, e.g., US Pat. No. 7,294,754 and Poueymirou et al. (2007) F0 generation mice that are essentially fully derived from the donor gene-targeted ES cells allowing immediate phenotypic analyses (Nature Biotech. 25(1):91-99). VELOCIMICE® (F0 mice fully derived from the donor ES cell) independently bearing a humanized CD3 genes were identified by genotyping using a modification of allele assay (see above) that detects the presence of the unique human CD3 gene sequences.

[0115] The selection cassette may be removed by methods known by the skilled artisan. For example, ES cells bearing the humanized CD3 locus may be transfected with a construct that expresses Cre in order to remove floxed cassette. The selection cassette may optionally be removed by breeding to mice that express Cre recombinase. Optionally, the selection cassette is retained in the mice. The mouse/human junction of the humanized CD3 ϵ allele after selection cassette removal (depicted as A-B in FIG. 2B), is presented in Table 3 below. The remaining junction sequences are the same as in the targeting vector and are presented in Table 1 above.

Table 3. Junctional Sequences of the Humanized Allele

Sequence designation in FIG. 1B	Junction	Sequence	SEQ ID NO:
A-B	5'mouse Cd3e/Xhol/Lox/IceUI //human CD3E	CGACTTCTTGACTCTATTGTTAAA CACTGTGCATTACATCGAATGCTAGA AGTTTCCTCGTCCGCTTCCTCGAA TTGCCTGGGATCCTCTGCTTGATGCC TGTAGGAAACGTCCTTCCCTGTGGTAT AGAAATGACTG/ CTCGAG/ATAACTTC GTATAATGTATGCTATACGAAGTTAT/ GCTAGTAACATAACGGTCCTAAGGTA GCGAGCTAGC / CTTCCACAGACACCA ATGTTAAAATGGAGGCTGGGGCAA AATTCTTGCTATGTCTCTAGTCGT CAAAAAATGGTCTAACTTTCTGAC TCCTGCTTGTCAAAAATTGTGGGCTCA TAGTTAATGC	8

[0116] The sequence of the resulting humanized CD3 ϵ , CD3 δ , and CD3 γ proteins is depicted in FIG. 3 and included in the sequence listing. Additionally, alignment of mouse-human sequences and junctions at the 5' and 3' of inserted human sequence are shown in FIG. 4 as * and **, respectively. GenBank Protein Accession Numbers for CD3 ϵ , CD3 δ , and CD3 γ proteins are summarized below in Table 4.

Table 4 GenBank Protein Accession Numbers

Protein Name	Mouse Accession # (SEQ ID NO)	Human Accession # (SEQ ID NO)
CD3 ϵ	NP_031674 (SEQ ID NO:27)	NP_000724 (SEQ ID NO:28)
CD3 δ	NP_038515 (SEQ ID NO:29)	NP_000723 (isoform A) (SEQ ID NO:30)
CD3 γ	NP_033980 (SEQ ID NO:31)	NP_000064 (SEQ ID NO:32)

Example 2. Characterization of Humanized CD3 Mice

Example 2.1. Immune Cell Development in Humanized CD3 Mice

[0117] Immune cell development in the thymus and periphery of human CD3 $\epsilon\delta\gamma$ mice was assessed using fluorescence-activated cell sorting (FACS) analysis and differential cell counting. Thymus, spleen and lymph nodes were harvested from cohorts of wildtype (WT, no human CD3 $\gamma\delta\epsilon$), heterozygous (Het, one hCD3 $\gamma\delta\epsilon$ allele) and homozygous (Ho, two hCD3 $\gamma\delta\epsilon$ alleles) mice. Peripheral blood was obtained by cardiac puncture or retro-orbital bleed into EDTA coated Microtainer tubes (BD). Single cell suspensions were prepared from the spleen, LN and thymus using mechanical disruption, and red blood cells were removed from the spleen, thymus and whole blood by lysis with AKC Lysis buffer.

Cells were incubated for 10 minutes at room temperature with purified antibodies to CD16/CD32 (FcBlock) to block non-specific binding via Fc receptors, and then incubated for 30 minutes at 4°C with a cocktail of directly conjugated antibodies to T and B cell markers. Cells were washed twice with cold PBS containing 1% BSA, resuspended in buffer and analyzed by flow cytometry on a FACSCanto II™ flow cytometer (BD Biosciences). Thymocytes were identified first by forward and side scatter gating, and then by gating on the B220- population. In the periphery, T cells were identified as CD45+/TCRb+/B220-, and B cells were identified as CD45+/TCRb-/B220+. Absolute counts were obtained on a Hemavet 950FS Hematology Analyzer.

[0118] As demonstrated in FIGS. 5A and 5B, humanized CD3 $\gamma\delta\epsilon$ mice appeared to have normal thymocyte development and normal T cell and B cell ratios in thymus, peripheral blood, and spleen. Additionally, T and B cell percentages appeared normal in lymph nodes, and absolute cell counts for spleen and lymph nodes (data not shown) were within normal range. CD4 and CD8 cell numbers in the blood were similar between the WT, Het, and Ho mice. Circulating white blood cells, lymphocytes, monocytes, neutrophils, eosinophils, and basophils all appeared within normal range (data not shown). Thus, normal immune cell development is observed in the humanized CD3 $\epsilon\delta\gamma$ mice.

[0119] In order to determine whether the humanized CD3 $\epsilon\delta\gamma$ mice exhibited a polyclonal V β CD4+ and CD8+ T cell repertoire, splenocytes were isolated from four humanized and five strain-matched control mice and examined for TCR V β usage. Spleens were harvested and single cell splenocytes prepared as described above. Cells were incubated for 10 minutes at room temperature with purified antibodies to CD16/CD32 (FcBlock; Biolegend) to block non-specific binding via Fc receptors, and then resuspended in a cocktail of directly conjugated antibodies to mouse CD4 (Biolegend) and mouse CD8 (Biolegend). The directly-conjugated antibodies to the TCR V β were then added to the individual wells and incubated for 30 minutes at 4°C. Cells were washed with cold PBS and incubated with a viability dye (LIVE/DEAD Fixable Aqua Dead cell stain, Life Technologies) for 15 minutes at room temperature. Cells were washed with cold PBS containing 2% FBS then resuspended in buffer and fixed with BD Stabilization buffer before being analyzed by flow cytometry on a LSR Fortessa™ flow cytometer (BD Biosciences). CD4 and CD8 T cells were identified first by forward and side scatter gating, and then by gating on the live population. CD4 T cells (CD4+CD8-) and CD8 T cells (CD4-CD8+) were then examined for TCR V β usage.

[0120] As can be seen from FIG. 5C, V β repertoire used by both CD4 and CD8 T cells in the humanized CD3 $\epsilon\delta\gamma$ mice shows polyclonality, with usage not significantly different from the strain-matched control mice.

Example 2.2. T Cell Response to Infection in Humanized CD3 Mice

[0121] To determine whether the humanized CD3 mice (humanized CD3 $\gamma\delta\epsilon$ mice) exhibited normal response to infection, the ability of humanized mice to clear lymphocytic choriomeningitis virus (LCMV) was tested. LCMV is a mouse tropic virus, where the fate of infection depends on the viral strain. Infection with Armstrong strain results in an acute infection, where mice can quickly mount a T cell response against the virus and clear the infection in about a week. On the other hand, Clone 13 virus cannot be cleared, and T cells become "exhausted" and chronic infection is established. As both chronic and acute infections depend on T cell activity, LCMV is an ideal model to test for T cell function.

[0122] 6-8 week old humanized CD3 or strain matched control mice were infected with 2x10⁵ ffu of Armstrong i.p. and/or 2x10⁶ ffu of Clone 13 i.v. for Clone 13 infection, two weeks after infection spleens were harvested and virus titers were measured by plaque assay. Viral titers were similar in both control and huCD3 mice (FIG. 6A), indicating that CD3 humanization did not have an effect on the T-cell exhaustion phenotype, as T-cells can control the virus to similar levels in both strains of mice. For the Armstrong strain infection, two weeks after initial Armstrong strain infection, mice were challenged with Clone 13 and two weeks after Clone 13 challenge viral titers were measured in spleens. No virus was detected in either control or humanized CD3 mice (FIG. 6B). The data suggests that the acute Armstrong infection was cleared. In addition, this demonstrates that T-cell memory that was elicited from the Armstrong infection was sufficient to protect mice from the subsequent Clone 13 infection in both strains of mice.

Example 3. Humanized CD3 Mice as a Model for Testing Anti-CD3-Based Therapeutic Candidates

Example 3.1. Humanized CD3 Mouse for Testing Cynomolgus Monkey Cross-Reactive Anti-Human CD3 Antibodies

[0123] The ability of different human restricted or cynomolgus cross-reactive anti-CD3 antibodies to bind splenocytes from wild type (WT) or humanized CD3 $\gamma\delta\epsilon$ (Ho = homozygous, Het = heterozygous) mice was tested using fluorescence-activated cell sorting (FACS) analysis.

[0124] Freshly isolated splenocytes (2x10⁵ per well) were incubated with anti-CD3 antibodies (15 ug/ml) for 30 minutes at 4°C. Post incubation, cells were washed twice and appropriate secondary antibodies (e.g. fluorescent-tagged PE anti-human IgG and directly conjugated antibodies to T cell markers) were added and incubated for an additional 30 minutes at 4°C, then washed twice. The following antibodies were used: ah/mfCD3-2 and ah/mfCD3-1 are two antibodies that recognize both human and monkey CD3; ahCD3-2 and ahCD3-1 are two antibodies that only recognize human CD3,

amCD3-2C11 is an antibody that recognizes mouse CD3 only, control human IgG is an unrelated control antibody, and 2nd only is a secondary antibody only control. Cells were washed twice with cold PBS containing 1% BSA, resuspended in buffer and analyzed by flow cytometry on a FACSCanto II™ flow cytometer (BD Biosciences). T cells were identified as CD45+/TCRb+/B220-. Anti-mCD3-2C11 engineered to contain hlgG1 was used to identify T cells on WT mouse splenocytes.

[0125] As demonstrated in FIG. 7, anti-CD3 antibodies that recognized only human CD3 were able to bind CD3 on the surface of splenocytes from humanized CD3 $\gamma\delta\epsilon$ mice; similarly anti-CD3 antibodies that recognized human and monkey CD3 were able to bind CD3 on the surface of humanized CD3 $\gamma\delta\epsilon$ mice. Thus, mice humanized for all three CD3 ϵ , CD3 δ , and CD3 γ are relevant for early pre-clinical studies of CD3-based drug candidates which can be followed up by efficacy and toxicity studies in cynomolgus monkeys.

Example 3.2. T Cell Activation in Humanized CD3 Mice

[0126] The ability of anti-human CD3 antibodies to elicit immune response in humanized CD3 mice was tested. Mice humanized for CD3 $\gamma\delta\epsilon$ (n of 2/group), were injected intraperitoneally with 10 ug of different human restricted or cynomolgus cross-reactive anti-CD3 antibodies (all hlgG1). To obtain cellular composition and plasma cytokine levels, blood was drawn into EDTA coated Microtainer tubes (BD) from the retro-orbital sinus starting 2 hours post injection. The number of peripheral T and B cells was assessed by FACS. Briefly, 50 ul whole blood was incubated for 30 minutes at 4°C with a cocktail of directly conjugated antibodies to T and B cell markers. Red blood cells were removed by lysis with AKC Lysis buffer, and the labeled cells were washed one time with cold PBS containing 1% BSA. After washing, the cells were re-suspended in cold buffer and analyzed by flow cytometry on a FACSCANTO II™ flow cytometer (BD Biosciences). T cells were identified as live CD45+/TCRb+/B220-, and B cells were identified as live CD45+/TCRb-/B220+. Absolute cell counts were determined by adding a known quantity of CountBright TM Absolute Counting Beads. Plasma cytokine levels were assessed using a Mouse ProInflammatory 7-Plex Ultra-Sensitive Kit (Meso-Scale Discovery) from blood obtained 2 hours post injection.

[0127] As demonstrated in FIG. 8A, injection of 10 ug of anti-CD3 antibodies induced a transient T and B cell depletion, which was largely restored by day 4 after initial antibody treatment. Additionally, injection of anti-CD3 antibodies (both anti-CD3 antibodies recognizing only human CD3 (ahCD3-1 and ahCD3-3) and anti-CD3 antibodies recognizing both human and monkey CD3 (ah/mfCD3-1 and ah/mfCD3-2)) induced cytokine production in CD3 $\gamma\delta\epsilon$ humanized mice (Fig. 8B).

[0128] In addition, the ability of anti-human CD3, anti-human/cynomolgus CD3, or anti-mouse antibodies to induce proliferation of splenocytes obtained from wild type or humanized CD3 $\gamma\delta\epsilon$ mice was assessed using ATP catalyzed quantification (CellTiter Glo®). The activation of mouse splenocytes results in the release of cytokines, which drive cellular proliferation. Proliferation data was acquired using the following protocol: splenocytes (5x10⁵/well) derived from wild type (WT) or humanized homozygous CD3 $\gamma\delta\epsilon$ (hCD3 $\gamma\delta\epsilon$ Ho) were added to 96 well plates which had been coated overnight at 4°C with decreasing amounts of human restricted, cynomolgus cross-reactive, or murine specific anti-CD3 antibodies. 500 ng/ml anti-mouse CD28 was added to the cultures, and the plates were incubated for 72 h at 37°C. Following incubation, CellTiter Glo® was added and luminescence was measured using a VICTOR X5 multi-label plate reader (PerkinElmer). The EC50 of cell viability (ATP catalyzed quantification) was determined using Prism (GraphPad Software, San Diego, CA). Values were calculated using a 4-parameter non-linear regression analysis.

[0129] As demonstrated in FIG. 9, splenocytes from humanized CD3 $\gamma\delta\epsilon$ mice were induced to proliferate by cynomolgus monkey-crossing CD3 antibodies.

[0130] A summary of various properties of WT and CD3 $\gamma\delta\epsilon$ mice are presented in FIG. 10. As can be seen, lymphocytes from CD3 $\gamma\delta\epsilon$ mice are able to bind anti-human CD3 antibodies and respond to anti-human CD3 antibodies, particularly those that are known to cross-react with monkey CD3, which is an important aspect for therapeutic agents as preclinical studies on drug candidates are often conducted in large animals such as cynomolgus monkeys.

Example 3.3. Tumor Depletion Studies in Humanized CD3 Mouse

[0131] Mice doubly humanized for both CD3 (humanized CD3 $\gamma\delta\epsilon$ mice described above) and CD20 were produced by crossing mice humanized at CD3 locus with mice humanized at CD20 locus. The resultant animals expressed both humanized proteins. Specifically, to produce humanized CD20 mice, the entire mouse Ms4a1 (Cd20) coding region from the 2nd amino acid (first being Met which is in common) to 167bp downstream 3' untranslated region, spanning 9312bp (Murine Chr. 19) was replaced with the corresponding CD20 human coding region from the 2nd amino acid to 107bp downstream 3'untranslated region, spanning 8482bp (Human Chr.11). Both mouse and human CD20 have six exons. The animals used in the experiment described below were homozygous for the replacements at both CD3 and CD20 loci and produced by crossing mice modified at the individual loci.

[0132] Humanized CD3/CD20 mice were implanted subcutaneously with 2x10⁵ B16F10.9 melanoma tumor cells trans-

duced with human CD20. Starting at Day 0 (day of tumor transplantation), mice were treated intraperitoneally 2 times per week with either vehicle (PBS; n=5), 0.4 mg/kg control Ab 2 (control antibody that does not display cross-reactivity to CD20 antigen; n=5), 0.4 mg/kg of Ab 1 (anti-CD3/CD20 bispecific antibody, see WO2014121087A1, published August 7, 2014, N=5), or 0.004 mg/kg Ab 1 (n=5). Tumor volumes were measured as indicated in Fig. 11A. Mice were sacrificed when tumors reached volume of greater than about 1500 mm³. As demonstrated in Fig. 11A, treatment with Ab 1 delayed tumor growth when treatment was initiated simultaneously with tumor transplantation.

[0133] In a separate experiment, ability of Ab 1 to inhibit tumor growth in an already established tumor was also tested (Fig. 11B). Humanized CD3/CD20 mice were implanted subcutaneously with 2x10⁵ B16F10.9 melanoma tumor cells expressing human CD20. On day 10 post tumor implantation, mice were randomized based on tumor size and organized into the following treatment groups, 5 mice in each group: vehicle (PBS), 4 mg/kg control Ab 2 (control antibody that does not display cross-reactivity to CD20 antigen), 4 mg/kg of Ab 1, or 0.4 mg/kg Ab 1. All mice were treated i.p. 2 times a week. Mice were sacrificed when tumors reached volume of greater than about 1500 mm³. As demonstrated in Fig. 11B, treatment with Ab 1 delayed tumor growth of already established tumors, demonstrating that the humanized CD3 mice are advantageous for early drug candidate studies.

Equivalents

[0134] Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents of the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims.

[0135] If an accession number or other citation is associated with different content at different times, the content in effect at the effective filing date of the application is meant, the effective filing date being the filing date of the earliest priority application referencing the citation, or if none, the actual filing date.

[0136] Unless otherwise apparent from the context any embodiment, aspect, element, feature, step or the like can be combined with any other.

SEQUENCE LISTING

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Asn Ile Thr Trp Phe Lys Asp Gly Lys Met Ile Gly Phe Leu Thr Glu
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Asp Lys Lys Lys Trp Asn Leu Gly Ser Asn Ala Lys Asp Pro Arg Gly
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Met Tyr Gln Cys Lys Gly Ser Gln Asn Lys Ser Lys Pro Leu Gln Val
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20

Tyr Tyr Arg Met Cys Gln Asn Cys Ile Glu Leu Asn Ala Ala Thr Ile
 85 90 95

30

Claims

35 1. A genetically modified mouse comprising
at an endogenous mouse CD3 ϵ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 ϵ protein
comprising an extracellular domain of a human CD3 ϵ protein operably linked to mouse CD3 ϵ transmembrane and
cytoplasmic domains,
40 at an endogenous mouse CD3 δ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 δ protein
comprising an extracellular domain of human CD3 δ operably linked to mouse CD3 δ transmembrane and cytoplasmic
domains, and
45 at an endogenous mouse CD3 γ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 γ protein
comprising an extracellular domain of human CD3 γ operably linked to mouse CD3 γ transmembrane and cytoplasmic
domains,
50 wherein the mouse expresses on the surface of its T cells a functional humanized CD3 complex comprising the
chimeric human/mouse CD3 ϵ , CD3 δ , and CD3 γ proteins and wherein the endogenous mouse CD3 ϵ , CD3 δ , and
CD3 γ loci are genetically modified not to express functional extracellular domains of endogenous mouse CD3 ϵ ,
CD3 δ , and CD3 γ proteins.

55 2. A genetically modified mouse embryonic stem (ES) cell comprising
at an endogenous mouse CD3 ϵ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 ϵ protein
comprising an extracellular domain of a human CD3 ϵ protein operably linked to mouse CD3 ϵ transmembrane and
cytoplasmic domains,
at an endogenous mouse CD3 δ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 δ protein
comprising an extracellular domain of human CD3 δ operably linked to mouse CD3 δ transmembrane and cytoplasmic
domains, and
55 at an endogenous mouse CD3 γ locus, a nucleic acid sequence encoding a chimeric human/mouse CD3 γ protein
comprising an extracellular domain of human CD3 γ operably linked to mouse CD3 γ transmembrane and cytoplasmic

domains

3. The mouse of claim 1 or the mouse ES cell of claim 2,

5 wherein the nucleic acid sequence encoding the chimeric CD3 ϵ protein replaces the nucleic acid sequence
encoding the corresponding endogenous mouse CD3 ϵ protein,
wherein the nucleic acid sequence encoding the chimeric CD3 δ protein replaces the nucleic acid sequence
encoding the corresponding endogenous mouse CD3 δ protein, and
wherein the nucleic acid sequence encoding the chimeric CD3 γ protein replaces the nucleic acid sequence
10 encoding the corresponding endogenous mouse CD3 γ protein.

15 4. The mouse or mouse ES cell of any one of the preceding claims wherein extracellular domain of the human CD3 ϵ
protein comprises the sequence of SEQ ID NO:33, the extracellular domain of the human CD3 δ protein comprises
the sequence of SEQ ID NO:34, and the extracellular domain of the human CD3 γ protein comprises the sequence
of SEQ ID NO:35.

20 5. The mouse or mouse ES cell of any of the preceding claims, which is heterozygous for the modified endogenous
mouse CD3 loci.

25 6. The mouse or mouse ES cell of any of claims 1-4, which is homozygous for the modified endogenous mouse CD3 loci.

30 7. A method of making a genetically modified mouse as defined in any preceding claim, comprising:

35 a) introducing into the genome of an embryonic stem (ES) cell of the mouse
at an endogenous mouse CD3 ϵ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 ϵ protein
comprising an extracellular domain of human CD3 ϵ operably linked to mouse CD3 ϵ transmembrane and cyto-
plasmic domains,
at an endogenous mouse CD3 δ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 δ protein
comprising an extracellular domain of human CD3 δ operably linked to mouse CD3 δ transmembrane and cyto-
plasmic domains, and
at an endogenous mouse CD3 γ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 γ protein
comprising an extracellular domain of human CD3 γ operably linked to mouse CD3 γ transmembrane and cyto-
plasmic domains; and
b) generating the genetically modified mouse from the cell, optionally, wherein the cell is a single ES cell, and
the single ES cell is introduced into a mouse embryo to generate a mouse.

40 8. A method of making a genetically modified mouse ES cell as defined in any of claims 2-6, comprising:

45 introducing into the genome of the ES cell
at an endogenous mouse CD3 ϵ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 ϵ protein
comprising an extracellular domain of human CD3 ϵ operably linked to mouse CD3 ϵ transmembrane and cyto-
plasmic domains,
at an endogenous mouse CD3 δ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 δ protein
comprising an extracellular domain of human CD3 δ operably linked to mouse CD3 δ transmembrane and cyto-
plasmic domains, and
at an endogenous mouse CD3 γ locus a nucleic acid sequence encoding a chimeric human/mouse CD3 γ protein
comprising an extracellular domain of human CD3 γ operably linked to mouse CD3 γ transmembrane and cyto-
plasmic domains.

50 9. The method of claim 7 or 8,

 wherein the nucleic acid sequence encoding the chimeric CD3 ϵ protein replaces the nucleic acid sequence encoding
the corresponding endogenous mouse CD3 ϵ protein,
 wherein the nucleic acid sequence encoding the chimeric CD3 δ protein replaces the nucleic acid sequence encoding
the corresponding endogenous mouse CD3 δ protein, and
 wherein the nucleic acid sequence encoding the chimeric CD3 γ protein replaces the nucleic acid sequence encoding
55 the corresponding endogenous mouse CD3 γ protein.

10. A mouse model for testing a CD3-based bispecific antigen-binding protein, wherein the antigen-binding protein is

capable of binding both CD3 and a non-mouse antigen of interest, comprising:

5 a) a mouse whose genome is genetically modified such that it expresses on the surface of its T cells a functional humanized CD3 complex comprising chimeric human/mouse CD3 ϵ , CD3 δ , and CD3 γ proteins, wherein the chimeric human/mouse CD3 ϵ protein comprises an extracellular domain of a human CD3 ϵ protein operably linked to transmembrane and cytoplasmic domains of a mouse CD3 ϵ protein, wherein the chimeric human/mouse CD3 δ protein comprises an extracellular domain of a human CD3 δ protein operably linked to transmembrane and cytoplasmic domains of a mouse CD3 δ protein, and wherein the chimeric human/mouse CD3 γ protein comprises an extracellular domain of a human CD3 γ protein operably linked to transmembrane and cytoplasmic domains of a mouse CD3 γ protein, and
10 b) a cell expressing or comprising the non-mouse antigen of interest, optionally wherein the antigen of interest is a human antigen of interest.

15 11. A method of screening an antigen binding protein that binds human CD3 and an antigen of interest, the method comprising:

20 a. Introducing the antigen of interest into a genetically modified mouse as defined in one of claims 1 and 3-6 or made according to the method of claim 7 or 9,
b. contacting the mouse with the antigen-binding protein, and
c. determining if the antigen-binding protein is efficacious in activating mouse T cells that express the functional CD3 complex.

25 12. The method of claim 11, wherein the step of introducing comprises genetically modifying the mouse to express the antigen of interest; infecting the mouse with the antigen of interest, optionally by performing viral or bacterial infection; and/or introducing into said mouse a cell expressing the antigen of interest.

13. The mouse model or the method of claim 10 or 12, wherein the cell is a tumor cell or a bacterial cell.

30 14. The mouse model or the method of any one of claims 10-13, wherein the mouse is an immunocompetent mouse.

35 15. The mouse model or method of any one of claims 10-14, wherein the antigen of interest is

40 (a) a tumor associated antigen, which optionally may be selected from the group consisting of ALK, BAGE proteins, BIRC5 (survivin), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, EGFR variant III, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, GAGE proteins, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5, EBV-derived LMP2, L1CAM, MAGE proteins, MLANA, MSLN, MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), RAGE proteins, RET, RGS5, ROR1, SART1, SART3, SLAMF7, SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMPRSS2, Thompson-nouvelle antigen, TNFRSF17, TYR, UPK3A, VTCN1, and WT1,
45 (b) an infectious disease antigen, which optionally may be a viral antigen, such as HIV; hepatitis A; hepatitis B; hepatitis C; herpes virus such as HSV-1, HSV-2, CMV, HAV-6, VZV, and Epstein Barr virus; adenovirus; influenza virus; flavivirus; echovirus; rhinovirus; coxsackie virus; coronavirus; respiratory syncytial virus; mumps virus; rotavirus; measles virus; rubella virus; parvovirus; vaccinia virus; HTLV; dengue virus; papillomavirus; molluscum virus; poliovirus; rabies virus; JC virus; ebola virus; and arboviral encephalitis virus antigen; or
50 (c) a bacterial antigen, which optionally may be chlamydia, rickettsia, mycobacteria, staphylococci, streptococci, pneumonococci, meningococci, gonococci, klebsiella, proteus, serratia, pseudomonas, legionella, diphtheria, salmonella, bacilli, cholera, tetanus, botulism, anthrax, plague, leptospira, and Lyme disease bacterial antigen.

55 16. The method of any one of claims 11-15, wherein the antigen-binding protein is

a) an antibody, a bispecific antibody, or a bispecific antigen-binding protein, and/or
b) is capable of recognizing a monkey CD3 protein.

55 17. The method of claim 13, wherein the antigen-binding protein is

a) capable of reducing, eliminating, or preventing tumor growth as compared to an agent that does not target

the antigen of interest and wherein the step of determining optionally comprises a tumor volume assay or a T cell mediated tumor cell killing assay, or
 b) capable of reducing, eliminating, or preventing bacterial or viral infection as compared to an agent that does not target the antigen of interest and wherein the step of determining optionally comprises the measurement of viral or bacterial titers.

Patentansprüche

10 1. Genetisch modifizierte Maus, umfassend:

eine Nucleinsäuresequenz an einem endogenen Maus-CD3 ϵ -Locus, die für ein chimäres Mensch/Maus-CD3 ϵ -Protein kodiert, das eine extrazelluläre Domäne eines menschlichen CD3 ϵ -Proteins umfasst, die operabel an eine Maus-CD3 ϵ -Transmembran- und -Zytoplasmadomäne gebunden ist,
 15 eine Nucleinsäuresequenz an einem endogenen Maus-CD3 δ -Locus, die für ein chimäres Mensch/Maus-CD3 δ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 δ umfasst, die operabel an eine Maus-CD3 δ -Transmembran- und -Zytoplasmadomäne gebunden ist, und
 20 eine Nucleinsäuresequenz an einem endogenen Maus- CD3 γ -Locus, die für ein chimäres Mensch/Maus-CD3 γ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 γ umfasst, die operabel an eine Maus-CD3 γ -Transmembran- und -Zytoplasmadomäne gebunden ist,
 25 wobei die Maus auf der Oberfläche ihrer T-Zellen einen funktionellen humanisierten CD3-Komplex exprimiert, der die chimären Mensch/Maus-CD3 ϵ -, -CD3 δ - und -CD3 γ -Proteine umfasst, und wobei die endogenen Maus-CD3 ϵ -, -CD3 δ - und -CD3 γ -Loci genetisch so modifiziert sind, dass sie die funktionellen extrazellulären Domänen von endogenen Maus-CD3 ϵ -, -CD3 δ - und -CD3 γ -Proteinen nicht exprimieren.

25 2. Genetisch modifizierte embryonische Maus-Stamm- (Maus-ES-) Zelle, die Folgendes umfasst:

eine Nucleinsäuresequenz an einem endogenen Maus-CD3 ϵ -Locus, die für ein chimäres Mensch/Maus-CD3 ϵ -Protein kodiert, das eine extrazelluläre Domäne eines menschlichen CD3 ϵ -Proteins umfasst, die operabel an eine Maus-CD3 ϵ -Transmembran- und -Zytoplasmadomäne gebunden ist,
 30 eine Nucleinsäuresequenz an einem endogenen Maus-CD3 δ -Locus, die für ein chimäres Mensch/Maus-CD3 δ -Protein kodiert, eine extrazelluläre Domäne von menschlichem CD3 δ umfasst, die operabel an eine Maus-CD3 δ -Transmembran- und -Zytoplasmadomäne gebunden ist, und
 35 eine Nucleinsäuresequenz an einem endogenen Maus- CD3 γ -Locus, die für ein chimäres Mensch/Maus-CD3 γ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 γ umfasst, die operabel an eine Maus-CD3 γ -Transmembran- und -Zytoplasmadomäne gebunden ist.

40 3. Maus nach Anspruch 1 oder Maus-ES-Zelle nach Anspruch 2,

wobei die Nucleinsäuresequenz, die für das chimäre CD3 ϵ -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 ϵ -Protein kodiert,
 45 wobei die Nucleinsäuresequenz, die für das chimäre CD3 δ -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 δ -Protein kodiert, und
 wobei die Nucleinsäuresequenz, die für das chimäre CD3 γ -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 γ -Protein kodiert.

50 4. Maus oder Maus-ES-Zelle nach einem der vorangegangenen Ansprüche, wobei die extrazelluläre Domäne des menschlichen CD3 ϵ -Proteins die Sequenz von SEQ ID NO: 33 umfasst, die extrazelluläre Domäne des menschlichen CD δ -Proteins die Sequenz von SEQ ID NO: 34 umfasst und die extrazelluläre Domäne des menschlichen CD3 γ -Proteins die Sequenz von SEQ ID NO: 35 umfasst.

55 5. Maus oder Maus-ES-Zelle nach einem der vorangegangenen Ansprüche, die in Bezug auf die modifizierten endogenen Maus-CD3-Loci heterozygot ist.

6. Maus oder Maus-ES-Zelle nach einem der Ansprüche 1 bis 4, die in Bezug auf die modifizierten endogenen Maus-CD3-Loci homozygot ist.

7. Verfahren zur Herstellung einer genetisch modifizierten Maus wie in einem der vorangegangenen Ansprüche definiert, das Folgendes umfasst:

5 a) das Einführen einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 ε -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 ε umfasst, die operabel an die Maus-CD3 ε -Transmembran- und -Zytoplasmadomäne gebunden ist, in das Genom einer embryonischen Stamm- (ES-) Zelle der Maus an einem endogenen Maus-CD3 ε -Locus,

10 das Einführen einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 δ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 δ umfasst, die operabel an die Maus-CD3 δ -Transmembran- und -Zytoplasmadomäne gebunden ist, in das Genom einer embryonischen Stamm- (ES-) Zelle der Maus an einem endogenen Maus-CD3 δ -Locus, und

15 das Einführen einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 γ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 γ umfasst, die operabel an die Maus-CD3 γ -Transmembran- und -Zytoplasmadomäne gebunden ist, in das Genom einer embryonischen Stamm- (ES-) Zelle der Maus an einem endogenen Maus-CD3 γ -Locus; und

15 b) das Bilden der genetisch modifizierten Maus aus der Zelle, wobei die Zelle gegebenenfalls eine einzelne ES-Zelle ist und die einzelne ES-Zelle in einen Mäuseembryo eingeführt wird, um eine Maus zu bilden.

8. Verfahren zur Herstellung einer genetisch modifizierten Maus-ES-Zelle wie in einem der Ansprüche 2 bis 6 definiert, das Folgendes umfasst:

das Einführen in das Genom der ES-Zelle:

20 einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 ε -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 ε umfasst, die operabel an die Maus-CD3 ε -Transmembran- und -Zytoplasmadomäne gebunden ist, an einem endogenen Maus-CD3 ε -Locus,

25 einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 δ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 δ umfasst, die operabel an die Maus-CD3 δ -Transmembran- und -Zytoplasmadomäne gebunden ist, an einem endogenen Maus-CD3 δ -Locus, und

30 einer Nucleinsäuresequenz, die für ein chimäres Mensch/Maus-CD3 γ -Protein kodiert, das eine extrazelluläre Domäne von menschlichem CD3 γ umfasst, die operabel an die Maus-CD3 γ -Transmembran- und -Zytoplasmadomäne gebunden ist, an einem endogenen Maus-CD3 γ -Locus.

9. Verfahren nach Anspruch 7 oder 8,

wobei die Nucleinsäuresequenz, die für das chimäre CD3 ε -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 ε -Protein kodiert,

wobei die Nucleinsäuresequenz, die für das chimäre CD3 δ -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 δ -Protein kodiert, und

35 wobei die Nucleinsäuresequenz, die für das chimäre CD3 γ -Protein kodiert, die Nucleinsäuresequenz ersetzt, die für das entsprechende endogene Maus-CD3 γ -Protein kodiert.

10. Mausmodell zum Testen eines CD3-basierten, bispezifischen, Antigen-bindenden Proteins, wobei das Antigen-bindende Protein dazu in der Lage ist, CD3 sowie ein Nicht-Maus-Antigen von Interesse zu binden, das Folgendes umfasst:

40 a) eine Maus, deren Genom genetisch so modifiziert ist, dass sie auf der Oberfläche ihrer T-Zellen einen funktionellen humanisierten CD3-Komplex exprimiert, der chimäre Mensch/Maus-CD3 ε -, -CD3 δ - und -CD3 γ -Proteine umfasst,

45 wobei das chimäre Mensch/Maus-CD3 ε -Protein eine extrazelluläre Domäne von menschlichem CD3 ε umfasst, die operabel an die Maus-CD3 ε -Transmembran- und -Zytoplasmadomäne gebunden ist,

wobei das chimäre Mensch/Maus-CD3 δ -Protein eine extrazelluläre Domäne von menschlichem CD3 δ umfasst, die operabel an die Maus-CD3 δ -Transmembran- und -Zytoplasmadomäne gebunden ist, und

50 wobei das chimäre Mensch/Maus-CD3 γ -Protein eine extrazelluläre Domäne von menschlichem CD3 γ umfasst, die operabel an die Maus-CD3 γ -Transmembran- und -Zytoplasmadomäne gebunden ist; und

b) eine Zelle, die das Nicht-Maus-Antigen von Interesse exprimiert oder umfasst, wobei das Antigen von Interesse gegebenenfalls ein menschliches Antigen von Interesse ist.

11. Verfahren zum Screenen eines Antigen-bindenden Proteins, das menschliches CD3 und ein Antigen von Interesse bindet, wobei das Verfahren Folgendes umfasst:

55 a. das Einführen des Antigens von Interesse in eine genetisch modifizierte Maus, die wie in einem der Ansprüche 1 und 3 bis 6 definiert ist oder die durch ein Verfahren nach Anspruch 7 oder 9 hergestellt wurde,

- b. das Kontaktieren der Maus mit dem Antigen-bindenden Protein und
- c. das Bestimmen, ob das Antigen-bindende Protein in Bezug auf die Aktivierung von Maus-T-Zellen, die den funktionellen CD3-Komplex exprimieren, wirksam ist.

5 **12.** Verfahren nach Anspruch 11, wobei der Schritt des Einführens genetisches Modifizieren der Maus, um das Antigen von Interesse zu exprimieren; Infizieren der Maus mit dem Antigen von Interesse, gegebenenfalls durch Herbeiführen einer viralen oder bakteriellen Infektion; und/oder Einführen einer Zelle, die das Antigen von Interesse exprimiert, in die Maus umfasst.

10 **13.** Mausmodell oder Verfahren nach Anspruch 10 oder 12, wobei die Zelle eine Tumorzelle oder eine Bakterienzelle ist.

14. Mausmodell oder Verfahren nach einem der Ansprüche 10 bis 13, wobei die Maus eine immunkompetente Maus ist.

15 **15.** Mausmodell oder Verfahren nach einem der Ansprüche 10 bis 14, wobei das Antigen von Interesse Folgendes ist:

- (a) ein tumorassoziiertes Antigen, das gegebenenfalls aus der aus den folgenden bestehenden Gruppe ausgewählt ist:

20 ALK, BAGE-Proteinen, BIRC5 (Survinin), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, EGFR Variante III, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, GAGE-Proteinen, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5, EBV-abgeleitetem LMP2, L1CAM, MAGE-Proteinen, MLANA, MSLN, MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1 B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), RAGE-Proteinen, RET, RGS5, ROR1, SART1, SART3, SLAMF7, SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMPRSS2, Thompson-nouvelle-Antigen, TNFRSF17, TYR, UPK3A, VTCN1 und WT1,

25 (b) ein Antigen einer Infektionskrankheit, das gegebenenfalls ein Virusantigen ist, wie z.B. HIV-; Hepatitis-A-; Hepatitis-B-; Hepatitis-C-; Herpesvirus- wie HSV-1-, HSV-2-, CMV-, HAV-6-, VZV- und Epstein-Barr-Virus-; Adenovirus-; Influenzavirus-; Flavivirus-; Echovirus-; Rhinovirus-; Coxsackie-Virus-, Coronavirus-; Respiratorisches-Syncytial-Virus-; Mumpsvirus-; Rotavirus-; Masernvirus-; Rötelnvirus-; Parvovirus-; Vaccinia-Virus-; HT-LV-; Dengue-Virus-; Papillomavirus-; Molluscum-Virus-; Poliovirus-; Tollwutvirus-; JC-Virus-; Ebolavirus-; und Arbovirales-Enzephalitisvirus-Antigen; oder

30 (c) ein bakterielles Antigen, das gegebenenfalls ein bakterielles Antigen für Chlamydia, Rickettsia, Mycobacteria, Staphylokokken, Streptokokken, Pneumokokken, Meningokokken, Gonokokken, Klebsiellen, Proteus, Serratia, Pseudomonas, Legionellen, Diphtherie, Salmonellen, Bacilli, Cholera, Tetanus, Botulismus, Anthrax, Pest, Lepospira und Borreliose ist.

35 **16.** Verfahren nach einem der Ansprüche 11 bis 15, wobei das Antigen-bindende Protein

- a) ein Antikörper, ein bispezifischer Antikörper oder ein bispezifisches Antigenbindendes Protein ist und/oder
- b) dazu in der Lage ist, ein Affen-CD3-Protein zu erkennen.

40 **17.** Verfahren nach Anspruch 13, wobei das Antigen-bindende Protein

45 a) dazu in der Lage ist, Tumorwachstum im Vergleich zu einem Mittel, das nicht auf das Antigen von Interesse abzielt, zu reduzieren, zu stoppen oder zu verhindern, wobei der Schritt des Bestimmens gegebenenfalls einen Tumorvolumentest oder einen T-Zellen-vermittelten Tumorzellentötungstest umfasst; oder

50 b) dazu in der Lage ist, eine bakterielle oder virale Infektion im Vergleich zu einem Mittel, das nicht auf das Antigen von Interesse abzielt, zu reduzieren, zu stoppen oder zu verhindern, wobei der Schritt des Bestimmens gegebenenfalls das Messen von Virus- oder Bakterientitern umfasst.

Revendications

55 **1.** Souris génétiquement modifiée comprenant

au niveau d'un locus CD3 ϵ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 ϵ humaine/murine chimérique comprenant un domaine extracellulaire d'une protéine CD3 ϵ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 ϵ murine,

au niveau d'un locus CD3 δ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 δ

humaine/murine chimérique comprenant un domaine extracellulaire de CD3 δ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 δ murine,
 5 au niveau d'un locus CD3 γ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 γ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 γ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 γ murine,
 dans laquelle la souris exprime sur la surface de ses lymphocytes T un complexe CD3 humanisé fonctionnel comprenant les protéines CD3 ϵ , CD3 δ et CD3 γ humaines/murines chimériques et dans laquelle les loci CD3 ϵ , CD3 δ et CD3 γ murins endogènes sont génétiquement modifiés pour ne pas exprimer les domaines extracellulaires fonctionnels des protéines CD3 ϵ , CD3 δ et CD3 γ murines endogènes.

10 2. Cellule souche embryonnaire (ES) de souris génétiquement modifiée comprenant au niveau d'un locus CD3 ϵ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 ϵ humaine/murine chimérique comprenant un domaine extracellulaire d'une protéine CD3 ϵ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 ϵ murine,
 15 au niveau d'un locus CD3 δ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 δ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 δ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 δ murine,
 au niveau d'un locus CD3 γ murin endogène, une séquence d'acide nucléique codant pour une protéine CD3 γ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 γ humaine lié de manière fonctionnelle 20 aux domaines transmembranaires et cytoplasmiques de CD3 γ murine.

25 3. Souris selon la revendication 1 ou cellule ES de souris selon la revendication 2, dans laquelle la séquence d'acide nucléique codant pour la protéine CD3 ϵ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 ϵ murine endogène correspondante,
 dans laquelle la séquence d'acide nucléique codant pour la protéine CD3 δ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 δ murine endogène correspondante, et
 dans laquelle la séquence d'acide nucléique codant pour la protéine CD3 γ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 γ murine endogène correspondante.

30 4. Souris ou cellule ES de souris selon l'une quelconque des revendications précédentes, dans laquelle le domaine extracellulaire de la protéine CD3 ϵ humaine comprend la séquence de SEQ ID NO : 33, le domaine extracellulaire de la protéine CD3 δ humaine comprend la séquence de SEQ ID NO : 34, et le domaine extracellulaire de la protéine CD3 γ humaine comprend la séquence de SEQ ID NO : 35.

35 5. Souris ou cellule ES de souris selon l'une quelconque des revendications précédentes, qui est hétérozygote pour les loci CD3 murins endogènes modifiés.

40 6. Souris ou cellule ES de souris selon l'une quelconque des revendications 1 à 4, qui est homozygote pour les loci CD3 murins endogènes modifiés.

45 7. Procédé de préparation d'une souris génétiquement modifiée telle que définie dans l'une quelconque des revendications précédentes, comprenant :
 a) l'introduction dans le génome d'une cellule souche embryonnaire (ES) de la souris au niveau d'un locus CD3 ϵ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 ϵ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 ϵ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 ϵ murine,
 au niveau d'un locus CD3 δ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 δ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 δ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 δ murine, et
 50 au niveau d'un locus CD3 γ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 γ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 γ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 γ murine ; et
 b) la génération de la souris génétiquement modifiée à partir de la cellule, éventuellement, dans lequel la cellule 55 est une cellule ES unique, et la cellule ES unique est introduite dans un embryon de souris pour générer une souris.

8. Procédé de préparation d'une cellule ES de souris génétiquement modifiée telle que définie dans l'une quelconque

des revendications 2 à 6, comprenant :

5 l'introduction dans le génome de la cellule ES
 au niveau d'un locus CD3 ϵ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 ϵ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 ϵ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 ϵ murine,
 10 au niveau d'un locus CD3 δ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 δ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 δ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 δ murine, et
 15 au niveau d'un locus CD3 γ murin endogène, d'une séquence d'acide nucléique codant pour une protéine CD3 γ humaine/murine chimérique comprenant un domaine extracellulaire de CD3 γ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques de CD3 γ murine.

20 9. Procédé selon la revendication 7 ou 8,
 15 dans lequel la séquence d'acide nucléique codant pour la protéine CD3 ϵ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 ϵ murine endogène correspondante,
 dans lequel la séquence d'acide nucléique codant pour la protéine CD3 δ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 δ murine endogène correspondante, et
 20 dans lequel la séquence d'acide nucléique codant pour la protéine CD3 γ chimérique remplace la séquence d'acide nucléique codant pour la protéine CD3 γ murine endogène correspondante.

25 10. Modèle de souris pour tester une protéine de liaison à l'antigène bispécifique basée sur CD3, dans lequel la protéine de liaison à l'antigène est capable de se lier à la fois à CD3 et à un antigène non murin d'intérêt, comprenant :

30 a) une souris dont le génome est génétiquement modifié de sorte qu'il exprime sur la surface de ses lymphocytes T un complexe CD3 humanisé fonctionnel comprenant les protéines CD3 ϵ , CD3 δ et CD3 γ humaines/murines chimériques,
 dans lequel la protéine CD3 ϵ humaine/murine chimérique comprend un domaine extracellulaire d'une protéine CD3 ϵ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques d'une protéine CD3 ϵ murine,
 35 dans lequel la protéine CD3 δ humaine/murine chimérique comprend un domaine extracellulaire d'une protéine CD3 δ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques d'une CD3 δ murine, et
 dans lequel la protéine CD3 γ humaine/murine chimérique comprend un domaine extracellulaire d'une protéine CD3 γ humaine lié de manière fonctionnelle aux domaines transmembranaires et cytoplasmiques d'une protéine CD3 γ murine, et
 40 b) une cellule exprimant ou comprenant l'antigène non murin d'intérêt, éventuellement dans lequel l'antigène d'intérêt est un antigène humain d'intérêt.

45 11. Procédé de criblage d'une protéine de liaison à l'antigène qui se lie à CD3 humain et un antigène d'intérêt, le procédé comprenant :

a. l'introduction de l'antigène d'intérêt dans une souris génétiquement modifiée telle de définie dans l'une des revendications 1 et 3 à 6 ou préparée selon le procédé selon la revendication 7 ou 9,
 50 b. la mise en contact de la souris avec la protéine de liaison à l'antigène, et
 c. la détermination si la protéine de liaison à l'antigène est efficace pour activer les lymphocytes T de la souris qui expriment le complexe CD3 fonctionnel.

55 12. Procédé selon la revendication 11, dans lequel l'étape d'introduction comprend la modification génétique de la souris pour exprimer l'antigène d'intérêt ; l'infection de la souris avec l'antigène d'intérêt, éventuellement en réalisant une infection virale ou bactérienne ; et/ou l'introduction dans ladite souris d'une cellule exprimant l'antigène d'intérêt.

13. Modèle de souris ou procédé selon la revendication 10 ou 12, dans lequel la cellule est une cellule tumorale ou une cellule bactérienne.

55 14. Modèle de souris ou procédé selon l'une quelconque des revendications 10 à 13, dans lequel la souris est une souris immunocompétente.

15. Modèle de souris ou procédé selon l'une quelconque des revendications 10 à 14, dans lequel l'antigène d'intérêt est

5 (a) un antigène associé à une tumeur, qui peut être éventuellement choisi dans le groupe constitué par ALK, les protéines BAGE, BIRC5 (survivine), BIRC7, CA9, CALR, CCR5, CD19, CD20 (MS4A1), CD22, CD27, CD30, CD33, CD38, CD40, CD44, CD52, CD56, CD79, CDK4, CEACAM3, CEACAM5, CLEC12A, EGFR, le variant III de l'EGFR, ERBB2 (HER2), ERBB3, ERBB4, EPCAM, EPHA2, EPHA3, FCRL5, FLT3, FOLR1, les protéines GAGE, GD2, GD3, GPNMB, GM3, GPR112, IL3RA, KIT, KRAS, LGR5, LMP2 dérivée d'EBV, L1CAM, les protéines MAGE, MLANA, MSLN, MUC1, MUC2, MUC3, MUC4, MUC5, MUC16, MUM1, ANKRD30A, NY-ESO1 (CTAG1B), OX40, PAP, PAX3, PAX5, PLAC1, PRLR, PMEL, PRAME, PSMA (FOLH1), les protéines RAGE, RET, RGS5, ROR1, SART1, SART3, SLAMF7, SLC39A6 (LIV1), STEAP1, STEAP2, TERT, TMRSS2, l'antigène Thompson-nouvelle, TNFRSF17, TYR, UPK3A, VTCN1 et WT1,

10 (b) un antigène de maladie infectieuse, qui peut être éventuellement un antigène viral, tel que le VIH ; l'hépatite A ; l'hépatite B ; l'hépatite C ; le virus de l'herpès tel que le virus VHS-1, VHS-2, CMV, VHA-6, VZV et d'Epstein Barr ; l'adénovirus ; le virus de la grippe ; le flavivirus ; l'échovirus ; le rhinovirus ; le virus coxsackie ; le coronavirus ; le virus respiratoire syncytial ; le virus ourlien ; le rotavirus ; le virus de la rougeole ; le virus de la rubéole ; le parvovirus ; le virus de la vaccine ; l'HTLV ; le virus de la dengue ; le papillomavirus ; le virus molluscum ; le poliovirus ; le virus de la rage ; le virus JC ; le virus Ebola ; et l'antigène du virus de l'encéphalite arbovirale ; ou

15 (c) un antigène bactérien, qui peut être éventuellement une chlamydia, une rickettsie, une mycobactérie, les staphylocoques, les streptocoques, les pneumocoques, les méningocoques, les gonocoques, une klebsiella, un proteus, une serratia, un pseudomonas, une légionelle, la diphtérie, une salmonelle, les bacilles, le choléra, le tétanos, le botulisme, l'anthrax, la peste, un leptospire et l'antigène bactérien de la maladie de Lyme.

16. Procédé selon l'une quelconque des revendications 11 à 15, dans lequel la protéine de liaison à l'antigène est

25 a) un anticorps, un anticorps bispécifique ou une protéine de liaison à l'antigène bispécifique, et/ou
b) capable de reconnaître une protéine CD3 de singe.

17. Procédé selon la revendication 13, dans lequel la protéine de liaison à l'antigène est

30 a) capable de réduire, éliminer ou empêcher une croissance tumorale par comparaison à un agent qui ne cible pas l'antigène d'intérêt et dans lequel l'étape de détermination comprend éventuellement un dosage de volume tumoral ou un dosage de destruction de cellules tumorales médiée par les lymphocytes T, ou
35 b) capable de réduire, éliminer ou empêcher une infection bactérienne ou virale par comparaison à un agent qui ne cible pas l'antigène d'intérêt et dans lequel l'étape de détermination comprend éventuellement la mesure des titres viraux ou bactériens.

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FIG. 1

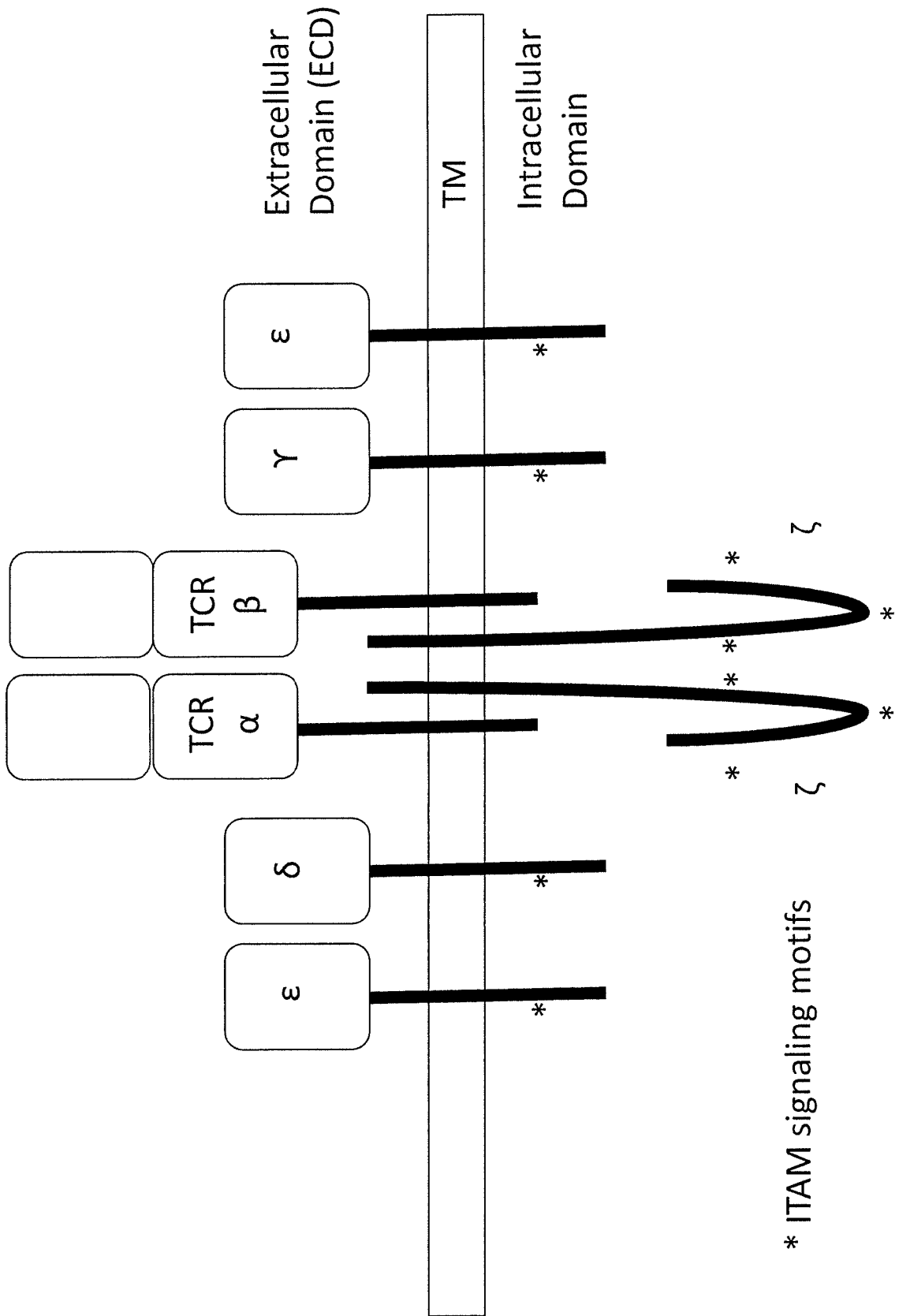


FIG. 2A

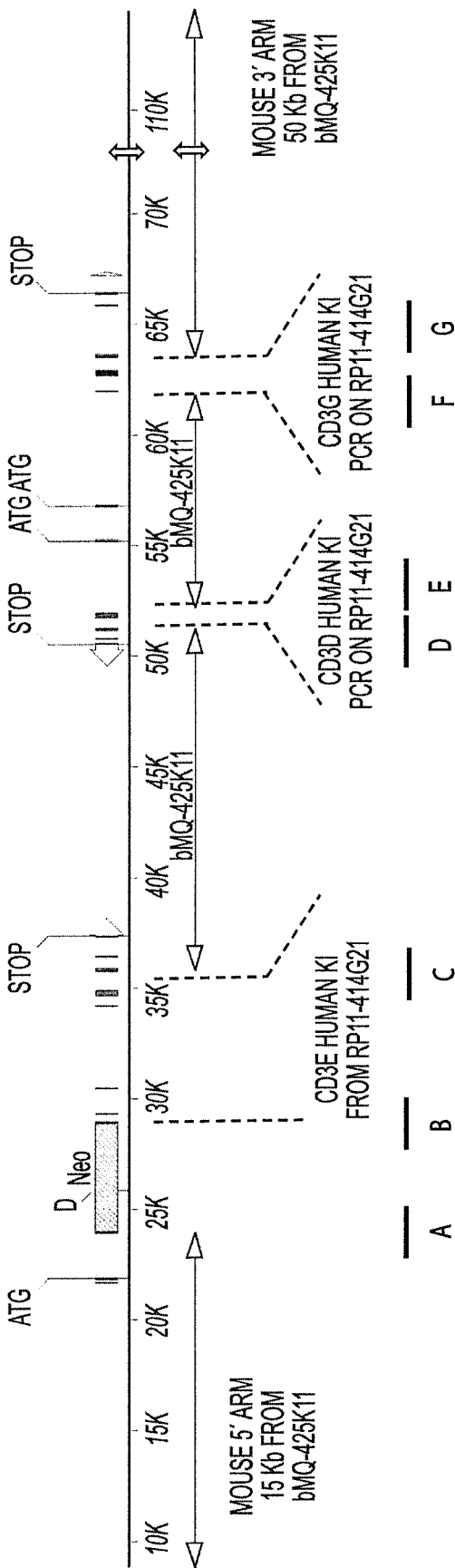


FIG. 2B

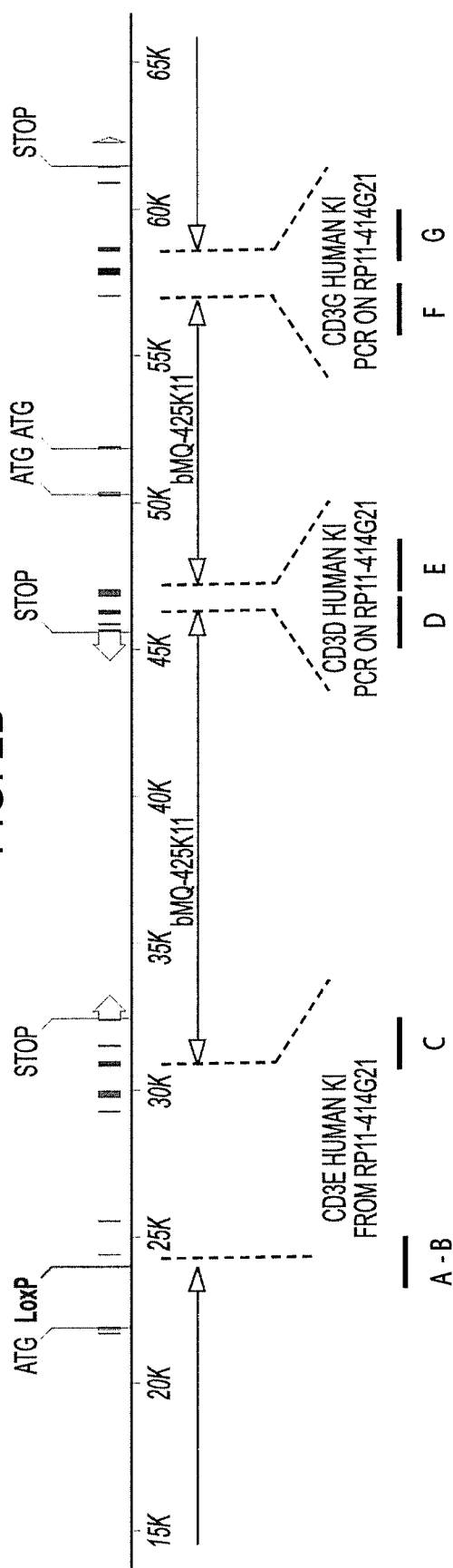


FIG. 3
CD3e humanized protein
(SEQ ID NO: 24)

MRWNTFWGILCLSLLAvgvwGQDGNEEMGGITQTPYKvSISGTTVILTCPQ
YPGSEILWQHNDKNIGGDEDDKNIGSDEDHLSLKEFSELEEQSGGYYVCYPRGSK
PEDANFYLRLARVCENCMEMDVMSVAIIVDICITLGLGLLMVYYWSKKNRKA
KAK
AKPVTRGTTGAGGSRPRGQQNKERPPVPNPDVEPIRKGQRDLYSGLNQRAV*

CD3d humanized protein
(SEQ ID NO:25)

MEHSGILASLILIAVLPQVSPFKIPIEELEDRvFVNCTSITWVEGTvGTLLSDTTRL
DLGKRILDPRGIYRCNGTDIYKDKESTVQVHYRMCQSCVELDPATVAGVIFIDLIA
TLLALGVYCFAGHETGRPSGAAEVQALLKNEQLYQPLRDDREDTQYSRLGGNWP
RNKKS*

CD3g humanized protein
(SEQ ID NO:26)

MEQRKGLAGLFLVISLLQGTLAQSIKGHNHLVKVYDYQEDGSVLLTCDAEAKNIT
WFKDGMIGFLTEDKKWNLGSNAKDPRGMYQCKGSQNKSKPLQVYYRMC
QNCIELNAATISGFIFAEVISIFFLALGVYLIAGQDGVRQSRASDIKQTLLQNEQLY
QPLKDREYDQYSHLQGNQLRKK*

Underscored residues coded by introduced human exons

FIG. 4

mCd3e/hCD3E protein alignment

mCD3ep	1 MRRNTFWGILCILSLLAVGTQDDAEN	1 LEYKVSISGTSVLETCPLDSDENLKWEKNGQELP	-QKHDK- -----	85
hCD3ep	1 MQSGTHWRLVGLCLLSSVWQGDGNEEMGGITQTPYKVSISGTTVILTCPOYPGSEILWQHNDKNIGGDEDKNTGSDDEHLSLKEFSELEQSYYVCYP	100	*****	100
mCD3ep	86 PAS -- NKNNTYLYLKARVCEYCVEDLTAVAILIVDICLITGLMVIYWSKNRKAZAKPEVTRGTAGGSRPRGONKERPPVNPDYEPIRKGQRDLYS	182	*****	182
hCD3ep	101 RSKPKEADNFYLYLPEAVCENCMEMDVMSTAVIVIDCITGGIILVIVYWSKNRKAZAKPEVTRGAGGGRQGONKERPPVNPDYEPIRKGQRDLYS	200	*****	200
mCD3ep	183 GLNQRAV 189			
hCD3ep	201 GLNQRRI 207			

mCd3d/hCD3D protein alignment

mCd3g/hCD3G protein alignment

FIG. 5A

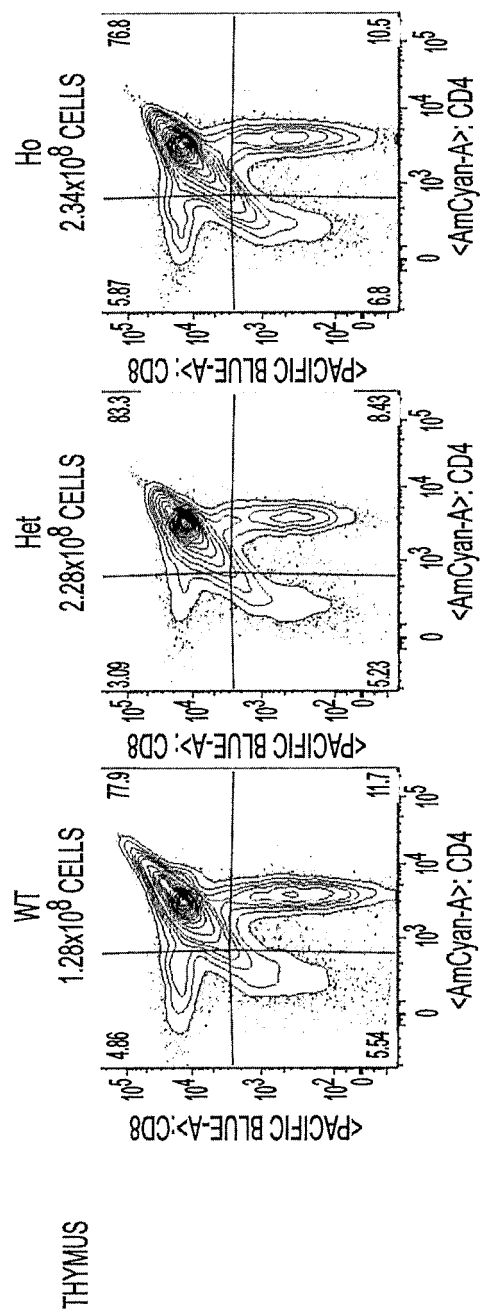


FIG. 5B

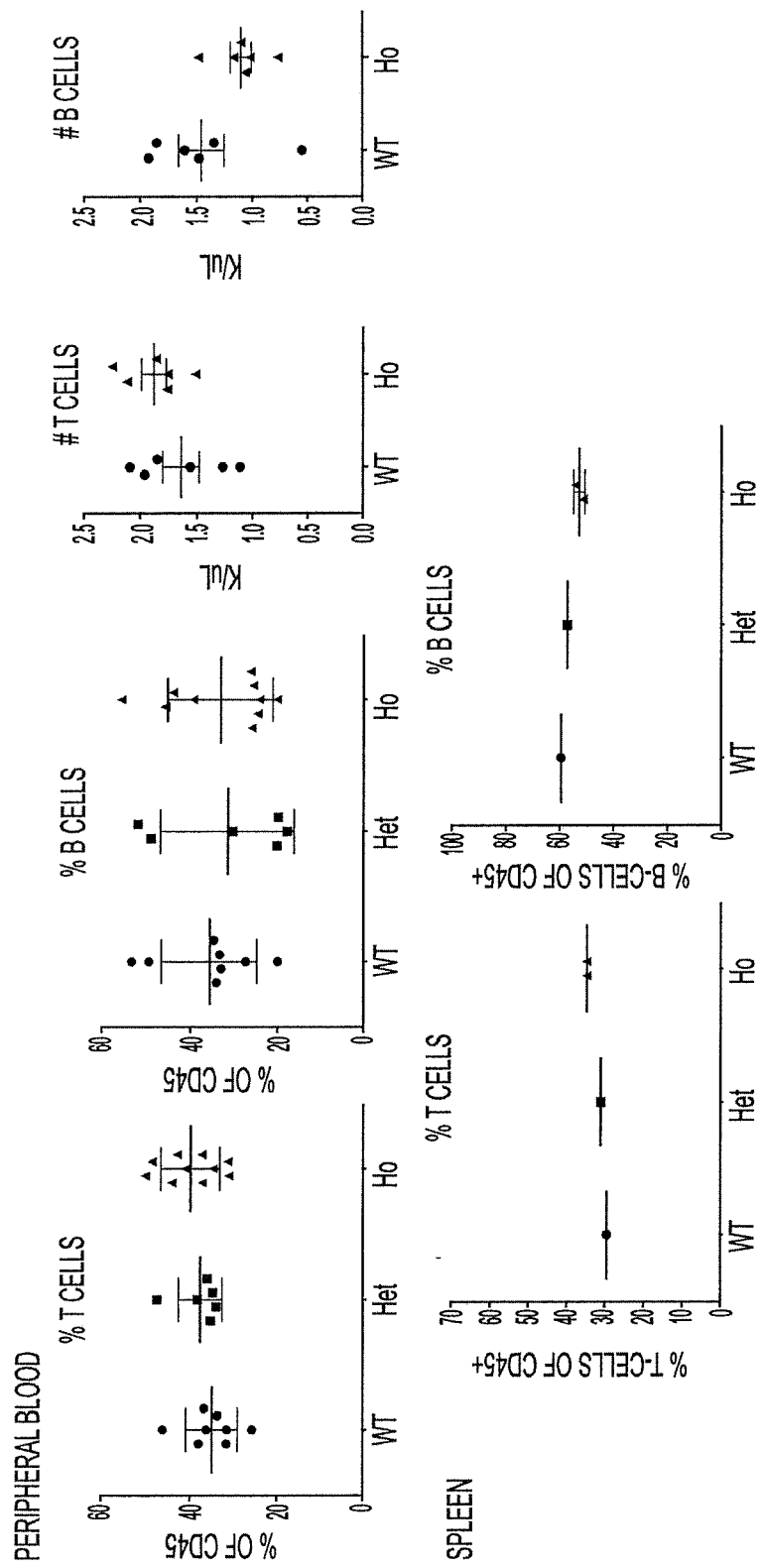
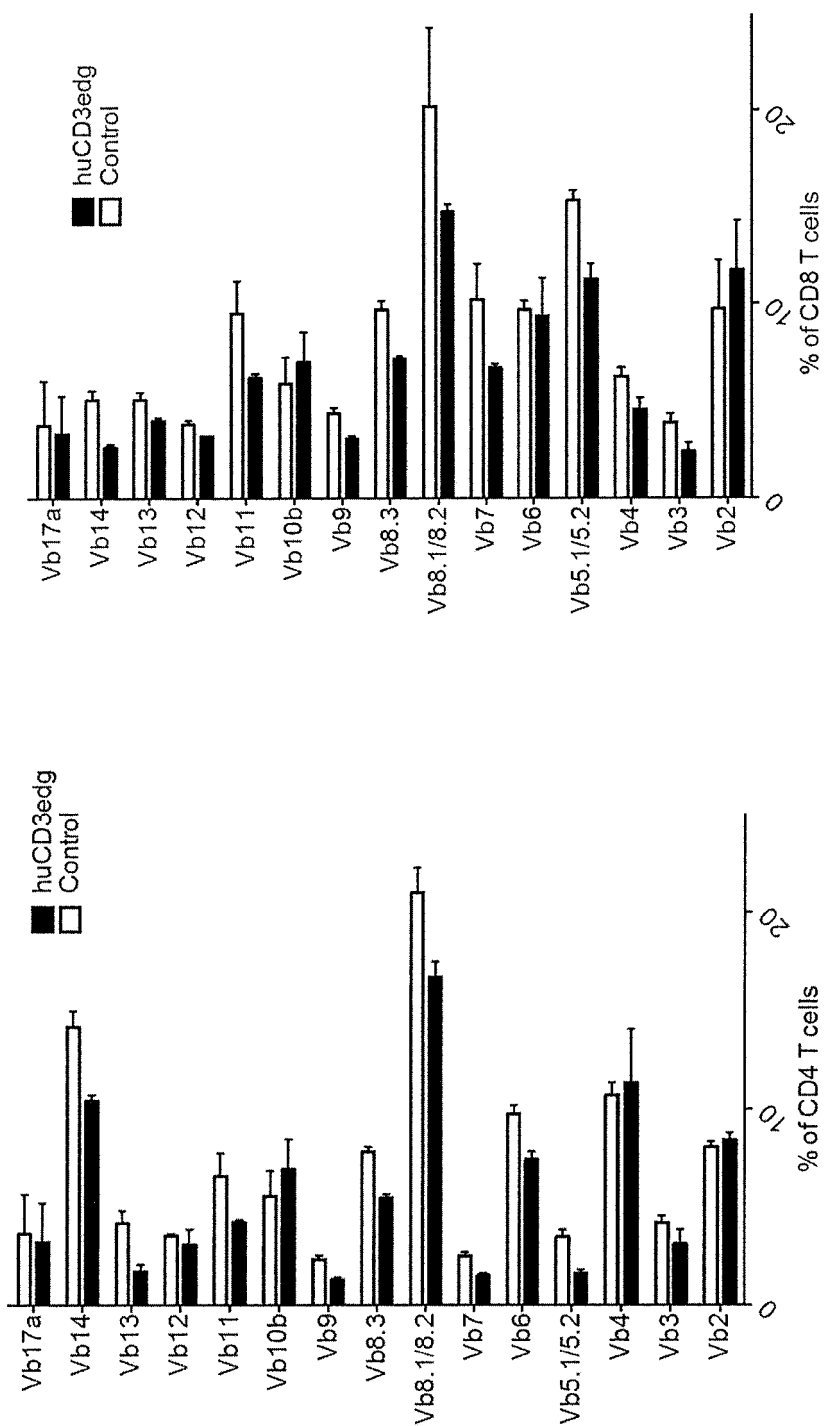


FIG. 5C



Humanized CD3 Mice Infected with LCMV Clone 13
with or without Previous LCMV Armstrong Infection
Spleen titers (ffu/mL)

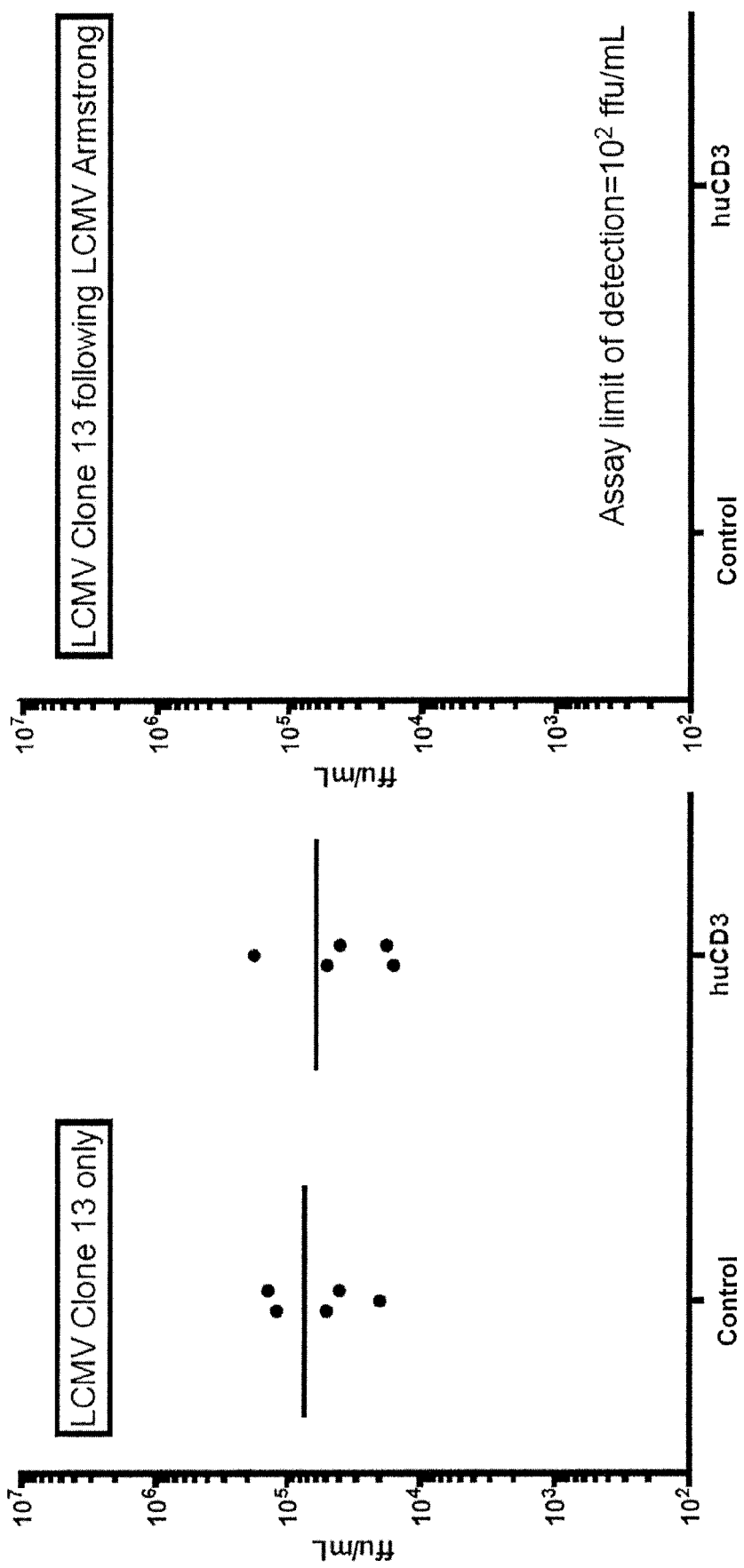
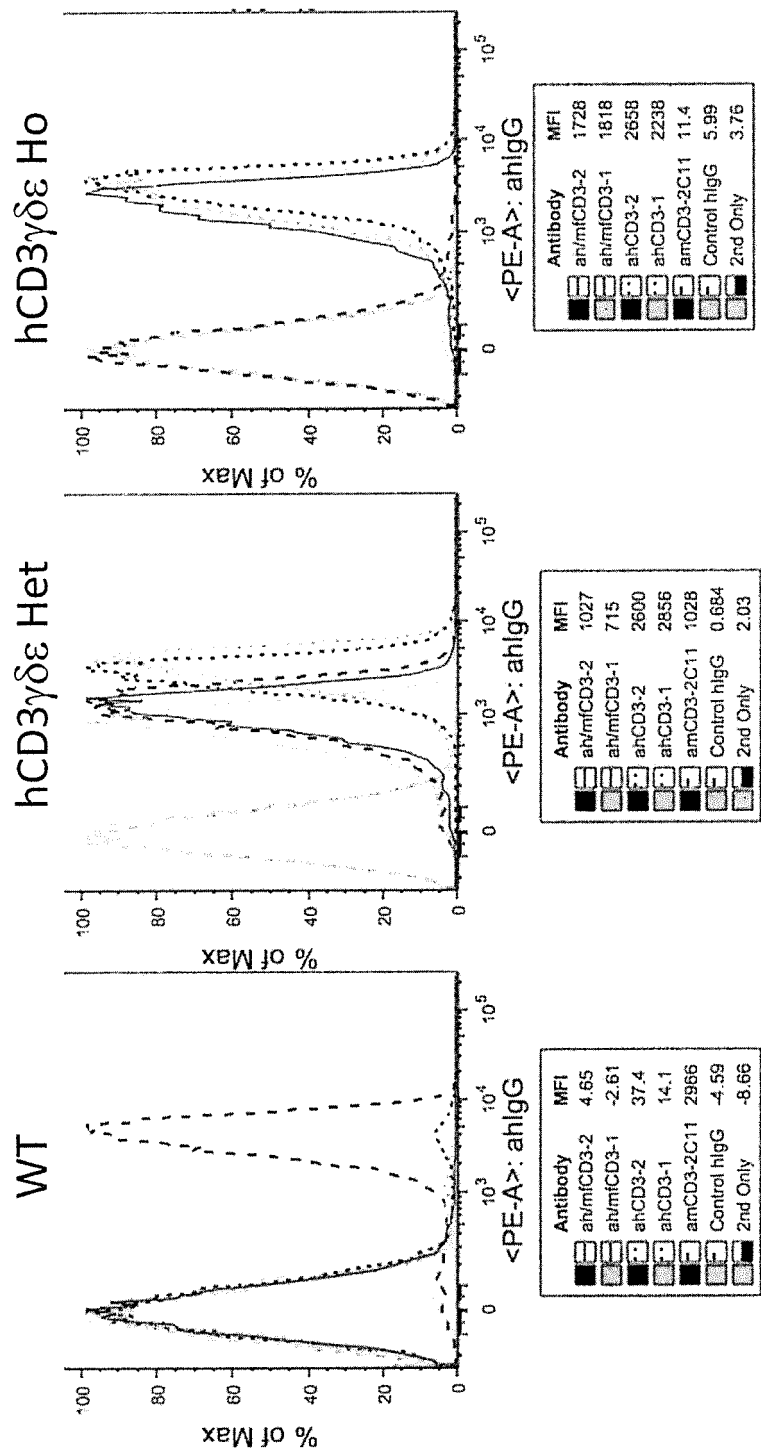


FIG. 6A

FIG. 6B

FIG. 7



- ah/mf = anti-human CD3 antibodies that cross-react to cynomolgus monkey CD3
- ah = anti-human CD3 antibodies that do not cross-react to cynomolgus monkey CD3

FIG. 8A

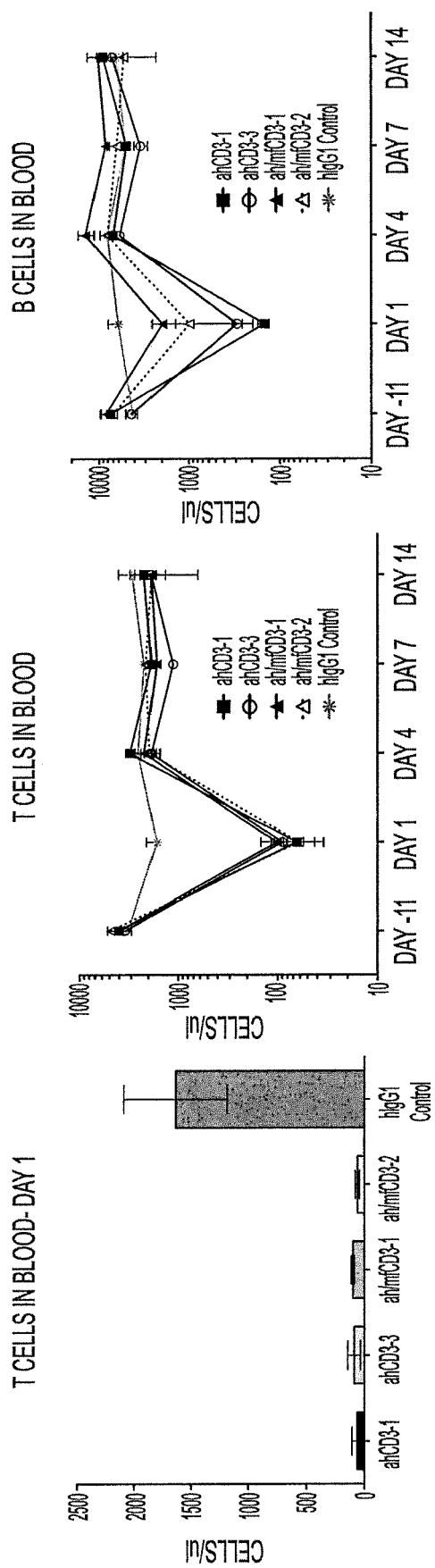


FIG. 8B

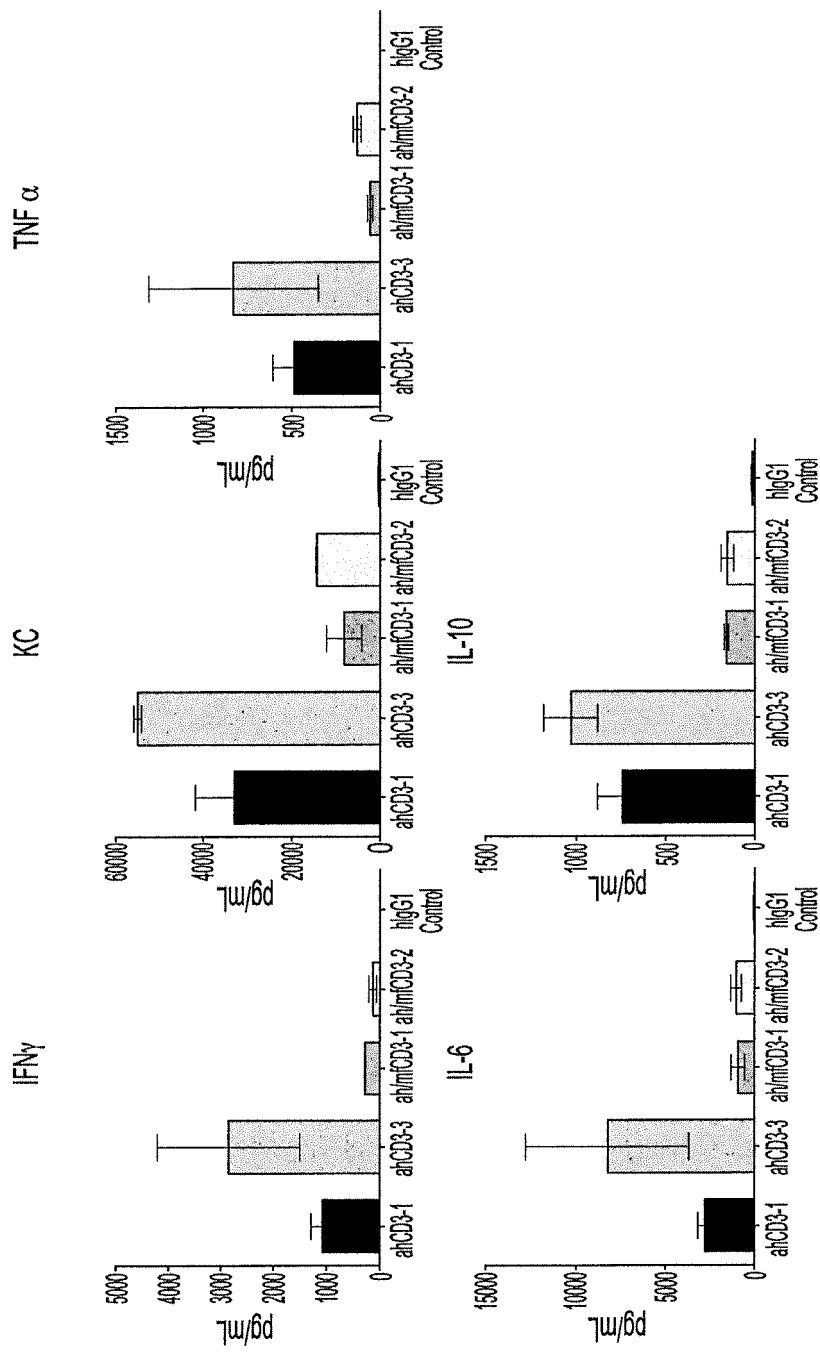


FIG. 9

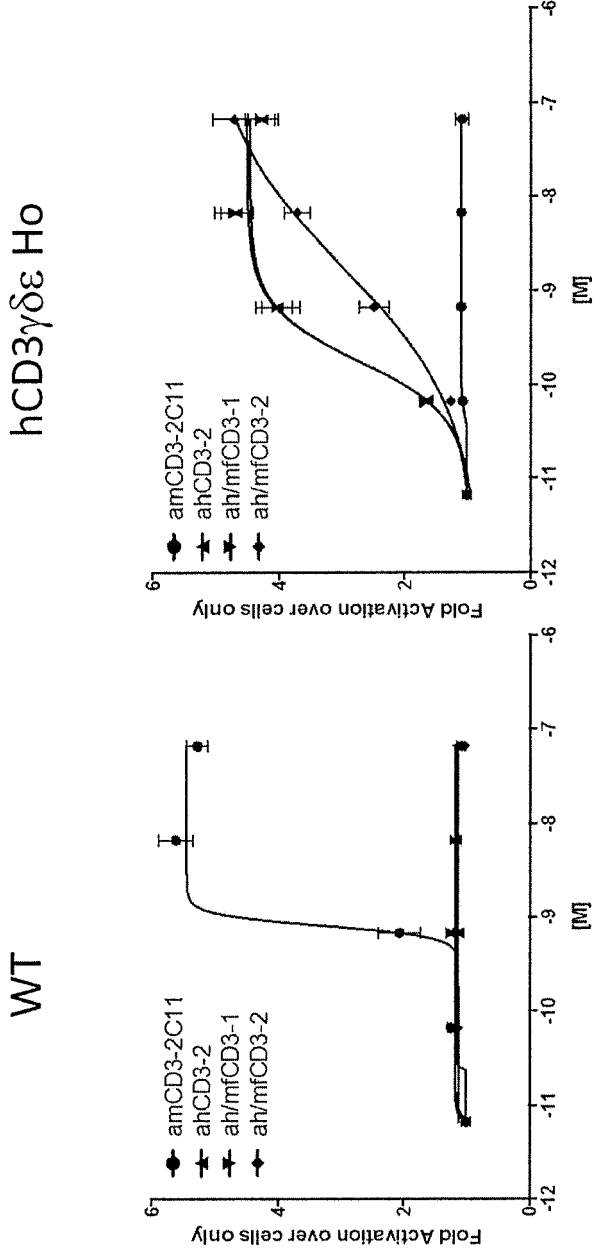
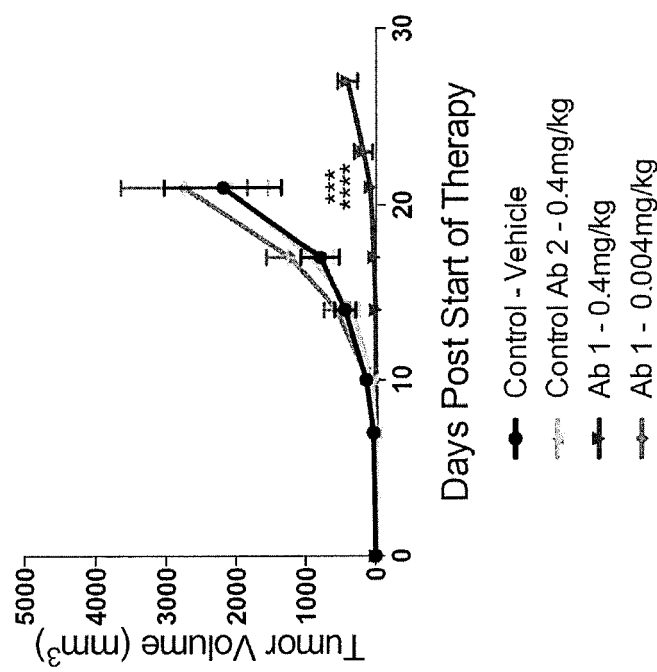


FIG. 10

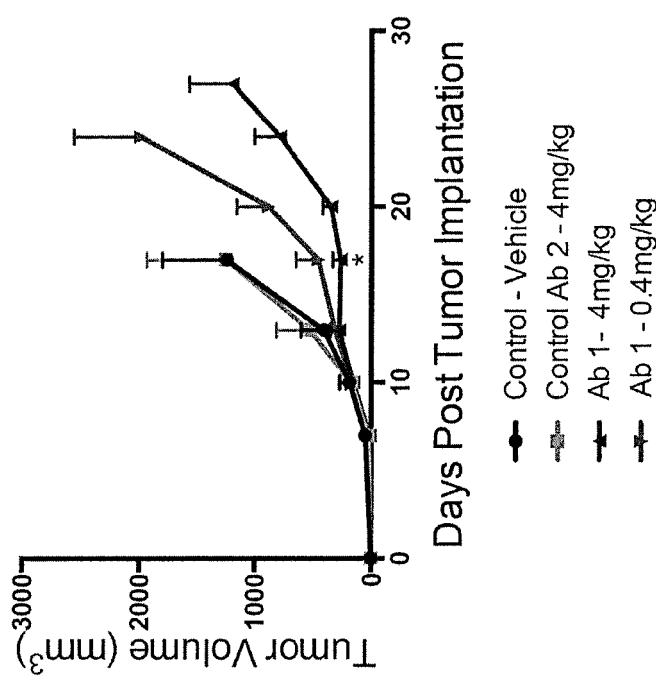
Genotype	ECD	T cell development	Responds to anti-mCD3	Binds anti-hCD3	Responds to anti-hCD3
WT	Mouse	Normal	Yes	No	No
hCD3gde-homozygous	Human	Normal	No	Yes, all antibodies tested, including cynomolgus monkey cross-reacting antibodies	Yes, all antibodies tested, including cynomolgus monkey cross-reacting antibodies

FIG. 11A



**** - compared to vehicle; ** - compared to Control Ab 2 control
 Data represents the composite data from $n=5$ mice per group. Data are expressed as mean (SEM) and
 were analysed using analysis of variance (ANOVA) and post hoc tests to probe significant effects
 (Tukey's for two-way ANOVA).

FIG. 11B



* - compared to vehicle and antibody control
Data represents the composite data from n=5 mice per group. Data are expressed as mean (SEM) and were analysed using analysis of variance (ANOVA).

REFERENCES CITED IN THE DESCRIPTION

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