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(57) Abstract: The present invention relates to novel anti-CD3 antibodies (as monoclonal antibodies or as used in other formats, such as bispecific or multi-specific formats) and compositions comprising such antibodies or cells activated, by such antibodies for use in treating disorders associated with CD3, such as human cancer therapy.



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ANTI-CD3 ANTIBODIES AND METHODS FOR THEIR USE

FIELD OF THE INVENTION

The present invention relates to novel anti-CD3 antibodies (as monoclonal antibodies or as
5 used in other formats, such as bispecific or multi-specific formats) and compositions comprising
such antibodies or cells activated by such antibodies for use in treating disorders associated with
CD3, such as human cancer therapy.

BACKGROUND OF THE INVENTION

10 The body's immune system serves as a defense against a variety of conditions, including,
e.g., injury, infection and neoplasia, and is mediated by two separate but interrelated systems: the
cellular and humoral immune systems. Generally speaking, the humoral system is mediated by
soluble products (antibodies or immunoglobulins) that have the ability to combine with and
neutralize products recognized by the system as being foreign to the body. In contrast, the cellular
15 immune system involves the mobilization of certain cells, termed T cells, that serve a variety of
therapeutic roles. T cells are lymphocytes that are derived from the thymus and circulate between
the tissues, lymphatic system and the circulatory system. They act against, or in response to, a
variety of foreign structures (antigens). In many instances these foreign antigens are expressed on
host cells as a result of neoplasia or infection. Although T cells do not themselves secrete
20 antibodies, they are usually required for antibody secretion by the second class of lymphocytes, B
cells (which derive from bone marrow). Critically, T cells exhibit extraordinary immunological
specificity so as to be capable of discerning one antigen from another).

A naive T cell, e.g., a T cell which has not yet encountered its specific antigen, is activated
when it first encounters a specific peptide:MHC complex on an antigen presenting cell. The
25 antigen presenting cell may be a B cell, a macrophage or a dendritic cell. When a naive T cell
encounters a specific peptide:MHC complex on an antigen presenting cell, a signal is delivered
through the T-cell receptor which induces a change in the conformation of the T cell's lymphocyte
function associated antigen (LFA) molecules, and increases their affinity for intercellular adhesion
molecules (ICAMs) present on the surface of the antigen presenting cell. The signal generated by
30 the interaction of the T cell with an antigen presenting cell is necessary, but not sufficient, to
activate a naive T cell. A second co-stimulatory signal is required. The naive T cell can be activated

only by an antigen-presenting cell carrying both a specific peptide MHC complex and a co-stimulatory molecule on its surface. Antigen recognition by a naive T cell in the absence of co-stimulation results in the T cell becoming anergic. The need for two signals to activate T cells and B cells such that they achieve an adaptive immune response may provide a mechanism for avoiding responses to self-antigens that may be present on an antigen presenting cell at locations in the system where it can be recognized by a T cell. Where contact of a T cell with an antigen presenting cell results in the generation of only one of two required signals, the T cell does not become activated and an adaptive immune response does not occur.

The efficiency with which humans and other mammals develop an immunological response to pathogens and foreign substances rests on two characteristics: the exquisite specificity of the immune response for antigen recognition, and the immunological memory that allows for faster and more vigorous responses upon re-activation with the same antigen (Portolés, P. et al. (2009) “*The TCR/CD3 Complex: Opening the Gate to Successful Vaccination*,” *Current Pharmaceutical Design* 15:3290-3300; Guy, C. S. et al. (2009) “*Organization of Proximal Signal Initiation at the TCR: CD3 Complex*,” *Immunol Rev.* 232(1):7-21). The specificity of the response of T-cells is mediated by the recognition of antigen (displayed on Antigen-Presenting Cells (APCs) by a molecular complex involving the T Cell Receptor (“TCR”) and the cell surface receptor ligand, CD3. The TCR is a covalently linked heterodimer of α and β chains (“TCR $\alpha\beta$ ”). These chains are class I membrane polypeptides of 259 (α) and 296 (β) amino acids in length. The CD3 molecule is a complex containing a CD3 γ chain, a CD3 δ chain, and two CD3 ϵ chains associated as three dimers ($\epsilon\gamma$, $\epsilon\delta$, $\zeta\zeta$) (Guy, C. S. et al. (2009) “*Organization of Proximal Signal Initiation at the TCR: CD3 Complex*,” *Immunol Rev.* 232(1):7-21; Call, M. E. et al. (2007) “*Common Themes In The Assembly And Architecture Of Activating Immune Receptors*,” *Nat. Rev. Immunol.* 7:841-850; Weiss, A. (1993) “*T Cell Antigen Receptor Signal Transduction: A Tale Of Tails And Cytoplasmic Protein-Tyrosine Kinases*,” *Cell* 73:209-212). The TCR and CD3 complex, along with the CD3 ζ chain zeta chain (also known as T-cell receptor T3 zeta chain or CD247) comprise the TCR complex (van der Merwe, P. A. etc. (epub Dec. 3, 2010) “*Mechanisms For T Cell Receptor Triggering*,” *Nat. Rev. Immunol.* 11:47-55; Wucherpfennig, K. W. et al. (2010) “*Structural Biology of the T-cell Receptor: Insights into Receptor Assembly, Ligand Recognition, and Initiation of Signaling*,” *Cold Spring Harb. Perspect. Biol.* 2:a005140). The complex is particularly

significant since it contains a large number (ten) of immunoreceptor tyrosine-based activation motifs (ITAMs).

In mature T cells, TCR/CD3 activation by foreign antigenic peptides associated to self-MHC molecules is the first step needed for the expansion of antigen-specific T cells, and their
5 differentiation into effector or memory T lymphocytes. These processes involve the phosphorylation of the immunoreceptor tyrosine-based activation motifs (ITAMs) of the TCR complex. Because the TCR complex has such a large number of ITAMS (10 in all), and these ITAMS are arrayed in tandem (due to the dimerization of the constituent chains), phosphorylation of the relevant tyrosine residues upon TCR ligation creates paired docking sites for proteins that
10 contain Src homology 2 (SH2) domains such as the ζ chain-associated protein of 70 kDa (ZAP-70), and thereby initiate an amplifying signaling cascade which leads to T-cell activation and differentiation (Guy, C. S. et al. (2009) "*Organization of Proximal Signal Initiation at the TCR: CD3 Complex*," Immunol Rev. 232(1):7-21).

The outcome of these processes is modulated by the intensity and quality of the antigen
15 stimulus, as well as by the nature of accompanying signals delivered by co-receptor and co-stimulatory surface molecules, or by cytokine receptors (Portoles, P. et al. (2009) "*The TCR/CD3 Complex: Opening the Gate to Successful Vaccination*," Current Pharmaceutical Design 15:3290-3300; Riha, P. et al. (2010) "*CD28 Co-Signaling In The Adaptive Immune Response*," Self/Nonself 1(3):231-240). Although TCR stimulation is a prerequisite for T-cell activation, it is
20 well recognized that engagement of co-stimulatory molecules, such as CD28, is necessary for full T-cell activation and differentiation (Guy, C. S. et al. (2009) "*Organization of Proximal Signal Initiation at the TCR:CD3 Complex*," Immunol Rev. 232(1):7-21).

Due to the fundamental nature of CD3 in initiating an anti-antigen response, monoclonal
25 antibodies against this receptor have been proposed as being capable of blocking or at least modulating the immune process and thus as agents for the treatment of inflammatory and/or autoimmune disease. Indeed, anti-CD3 antibodies were the first antibody approved for the human therapy (St. Clair E. W. (2009) "*Novel Targeted Therapies for Autoimmunity*," Curr. Opin. Immunol. 21(6):648-657). Anti-CD3 antibody (marketed as ORTHOCLONETM OKT3TM by Janssen-Cilag) has been administered to reduce acute rejection in patients with organ transplants
30 and as a treatment for lymphoblastic leukemia (Cosimi, A. B. et al. (1981) "*Use Of Monoclonal Antibodies To T-Cell Subsets For Immunologic Monitoring And Treatment In Recipients Of Renal*

Allografts,” N. Engl. J. Med. 305:308-314; Kung, P. et al. (1979) Monoclonal antibodies defining distinctive human T cell surface antigens,” Science 206:347-349; Vigerel, P. et al. (1986) “Prophylactic Use Of OKT3 Monoclonal Antibody In Cadaver Kidney Recipients. Utilization Of OKT3 As The Sole Immunosuppressive Agent,” Transplantation 41:730-733; Midtvedt, K. et al. 5 (2003) “Individualized T Cell Monitored Administration Of ATG Versus OKT3 In Steroid-Resistant Kidney Graft Rejection,” Clin. Transplant. 17(1):69-74; Gramatzki, M. et al. (1995) “Therapy With OKT3 Monoclonal Antibody In Refractory T Cell Acute Lymphoblastic Leukemia Induces Interleukin-2 Responsiveness,” Leukemia 9(3):382-390; Herold, K. C. et al. (2002) “Anti-CD3 Monoclonal Antibody In New-Onset Type 1 Diabetes Mellitus,” N. Engl. J. Med. 346:1692-1698; Cole, M. S. et al. (1997) “Human IgG2 Variants Of Chimeric Anti-CD3 Are Nonmitogenic to T cells,” J. Immunol. 159(7):3613-3621; Cole, M. S. et al. (1999) “Hum291, A Humanized Anti-CD3 Antibody, Is Immunosuppressive To T Cells While Exhibiting Reduced Mitogenicity in vitro,” Transplantation 68:563-571; U.S. Pat. Nos. 6,491,916; 5,585,097 and 6,706,265).

However, such anti-CD3 treatment has not proven to be specific enough to avoid side 15 effects (Ludvigsson, J. (2009) “The Role of Immunomodulation Therapy in Autoimmune Diabetes,” J. Diabetes Sci. Technol. 3(2):320-330). Repeated daily administration of OKT3 results in profound immunosuppression and provides effective treatment of rejection following renal transplantation. The in vivo administration of OKT3 results in both T cell activation and suppression of immune responses. However, the use of OKT3 has been hampered by a first toxic 20 dose reaction syndrome that is related to initial T-cell activation events and to the ensuing release of cytokines that occurs before immunosuppression of T cell responses. The reported side effects that follow the first and sometimes the second injection of this mouse monoclonal antibody include a “flu-like” syndrome consisting of high fever, chills, headache, and gastrointestinal symptoms (vomiting and diarrhea) and in severe cases pulmonary edema within hours of treatment has been 25 noted (Thistlethwaite, J. R. Jr. et al. (1988) “Complications and Monitoring of OKT3 Therapy,” Am. J. Kidney Dis. 11:112-119). This syndrome is believed to reflect OKT3-mediated cross-linking of the TCR/CD3 complex on the T cell surface and the resultant release of cytokines (e.g., tumor necrosis factor alpha (TNF α), interferon- γ , interleukins IL-2, IL-3, IL-4, IL-6, IL-10 and granulocyte-macrophage colony-stimulating factor (Masharani, U. B. et al. (2010) “Teplizumab 30 Therapy For Type 1 Diabetes,” Expert Opin. Biol. Ther. 10(3):459-465; Abramowicz, D. et al. (1989) “Release Of Tumor Necrosis Factor, Interleukin-2, And Gamma-Interferon In Serum After

Injection Of OKT3 Monoclonal Antibody In Kidney Transplant Recipients,” Transplantation 47:606-608; Ferran, C. et al. (1990) “*Cytokine-Related Syndrome Following Injection Of Anti-CD3 Monoclonal Antibody: Further Evidence For Transient In Vivo T Cell Activation,*” Eur. J. Immunol. 20:509-515; Hirsch, R. et al. (12989) “*Effects Of In Vivo Administration Of Anti-CD3 Monoclonal Antibody On T Cell Function In Mice. II. In Vivo Activation Of T Cells,*” J. Immunol. 142:737-743). The use of anti-CD3 antibodies is disclosed in U.S. Pat. Nos. 7,883,703; 7,728,114; 7,635,472; 7,575,923; and 7,381,903, and in United States Patent Publications Nos. 2010/0150918; 2010/0209437; 2010/0183554; 2010/0015142, 2008/0095766, 2007/0077246 and in PCT Publication WO2008/119567.

10

BRIEF SUMMARY OF THE INVENTION

The present invention relates to novel anti-CD3 antibodies (as monoclonal antibodies or as used in other formats, such as bispecific or multi-specific formats) and compositions comprising such antibodies or cells activated by such antibodies for use in treating disorders associated with CD3, such as human cancer therapy. The present invention also includes compositions comprising one or more of these peptides/antibodies, or fragments thereof, and/or immune cells that are modified to include and/or be activated by one or more of these antibodies, or fragments thereof, to treat a disease or condition, such as cancer. Antibodies of the present invention may similarly find use as a targeting arm of a bispecific or multi-specific format.

15

In one aspect, the present invention provides an anti-CD3 antibody or antibody fragment comprising a heavy chain CDR3 sequence (VH) comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59. Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

20

Additionally or alternatively, the present invention provides an anti-CD3 antibody or antibody fragment comprising a heavy chain CDR3 sequence (VH) comprising an amino acid

sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59 and a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175. Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175 and a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

In certain aspects, the present invention provides methods for treating a disorder, such as cancer, using one or more antibodies of the invention as described above, by, at least administering such antibodies to a subject, such as a human subject.

In certain aspects, the present invention provides immunoconjugates and/or compositions comprising such immunoconjugates, wherein said immunoconjugates comprise an anti-

CD3 antibody of the invention conjugated to another therapeutic agent, such as an anti-cancer agent. The invention further provides immunoconjugates comprising two or more different anti-CD3 antibodies or fragment thereof, wherein each different anti-CD3 antibody or fragment targets a different CD3 fragment or epitope.

5 A further aspect of the invention relates to a nucleic acid molecule having a nucleotide sequence that encodes an anti-CD3 antibody or fragment thereof, as disclosed herein, as well as expression vectors comprising such a polynucleotide and host cells that have been transfected with such an expression vector.

10 Aspects of the invention also provide methods for producing the anti-CD3 antibodies, fragments thereof, and compositions of the invention.

The present invention also provides methods for treating a disease in a human or animal subject, in particular treatment of cancer in humans, by administering an anti-CD3 antibody or composition of the invention to said subject. The invention also includes the use of one or more anti-CD3 antibodies of the invention for preparation of a medicament for use in treating a disease
15 in a human or animal, in particular for the treatment of cancer in humans.

In some embodiments, the antibodies herein are full length antibodies. In some embodiments, the antibodies are an IgA, an IgD, an IgE, an IgG, or an IgM antibody. In some embodiments, the anti-CD3 antibody is an IgG antibody (e.g., an IgG1, IgG2, or IgG3 antibody).

20 In some embodiments, the antibodies herein are an antibody fragment. In some embodiments, the antibodies are an Fv fragment, a Fab fragment, a F(ab')₂ fragment, a Fab' fragment, a Fab'-SH, an scFv (sFv) fragment, and an scFv-Fc fragment. In some embodiments, the bispecific antibody is an scFv fragment. In some embodiments, the antibodies herein are monoclonal, human, humanized, or chimeric.

25 In some embodiments, the antibodies further comprise an Fc region. In some embodiments, the antibodies comprise one or more heavy chain constant domains, wherein the one or more heavy chain constant domains are selected from a first CH1 domain, a first CH2 domain, a first CH3 domain, a second CH1 domain, a second CH2 domain, and a second CH3 domain. In some embodiments, one or more heavy constant chain domains are paired with another heavy chain constant domain.

In some embodiments, the antibodies further comprise a glycosylation site mutation. In some embodiments, the mutation reduces effector function. In some embodiments, the mutation is a substitution mutation.

5

DETAILED DESCRIPTION OF THE INVENTION

The T cell receptor (TCR) binds to antigens (Ags) displayed by major histocompatibility complexes (MHCs) and plays critical roles in T cell function. But the TCR does not possess intracellular signaling by itself. Instead, TCR non-covalently associates with the Cluster of Differentiation 3 (CD3) complex and triggers intracellular signaling through immunoreceptor tyrosine-based activation motifs (ITAM) of CD3. The CD3 T cell co-receptor helps to activate both the cytotoxic T cell (CD8⁺ naive T cells) and also T helper cells (CD4⁺ naive T cells). It consists of a protein complex and is composed of four distinct chains. In mammals, the complex contains a CD3 γ chain, a CD3 δ chain, and two CD3 ϵ chains. These chains associate with the T cell receptor (TCR) and the CD3 ζ chain (zeta-chain) to generate an activation signal in T lymphocytes. The TCR, ζ -chain, and CD3 γ , δ , and ϵ chains together constitute the TCR complex. The CD3 four-chain complex then forms CD3 $\epsilon\gamma$, CD3 $\epsilon\delta$ and $\delta\delta$ dimers in 1: 1: 1 stoichiometry.

CD3 is initially expressed in the cytoplasm of pro-thymocytes, the stem cells from which T cells arise in the thymus. The pro-thymocytes differentiate into common thymocytes, and then into medullary thymocytes, and it is at this latter stage that CD3 antigen begins to migrate to the cell membrane. The antigen is found bound to the membranes of all mature T cells, and in virtually no other cell type, although it does appear to be present in small amounts in Purkinje cells.

This high specificity, combined with the presence of CD3 at all stages of T cell development, makes it a useful immunohistochemical marker for T cells in tissue sections. The antigen remains present in almost all T cell lymphomas and leukemias, and can therefore be used to distinguish them from superficially similar B cell and myeloid neoplasms. Some antibodies against CD3 ϵ chain have been shown to activate TCR-CD3 complex, possibly by clustering CD3 complex on T cells. In addition, bispecific antibodies targeting both CD3 and a tumor specific antigen have been studied for redirected tumor eradication by T cells. Because CD3 is required for T cell activation, drugs (often monoclonal antibodies) that target it are being investigated as immunosuppressant therapies (e.g., oteelixizumab) for type 1 diabetes and other autoimmune diseases.

The present invention is directed to novel peptides (e.g., antibodies and antibody fragments) that bind to CD3. The present invention also includes compositions comprising one or more of these peptides/antibodies, or fragments thereof, and/or immune cells that are modified to include and/or be activated by one or more of these antibodies, or fragments thereof, to treat a disease or condition, such as cancer.

The presently disclosed antibodies may provide treatments that are far more effective than current therapies than present CD3 treatments. The presently disclosed anti-CD3 antibodies of the invention may be included as part of a treatment regime, which may include, for example, providing two or more such antibodies, and/or in combination with other treatments such as chemotherapy.

Definitions

The term “antibody” or “antibody molecule” describes a functional component of serum and is often referred to either as a collection of molecules (antibodies or immunoglobulin) or as one molecule (the antibody molecule or immunoglobulin molecule). An antibody is capable of binding to or reacting with a specific antigenic determinant (the antigen or the antigenic epitope), which in turn may lead to induction of immunological effector mechanisms. An antibody is generally considered as monospecific, and a composition of antibodies may be monoclonal (i.e., consisting of identical antibody molecules) or polyclonal (i.e., a plurality of different antibodies that may react with the same or different epitopes on the same antigen or on distinct/different antigens). An antibody has a unique structure enabling it to bind specifically to its corresponding antigen, and all natural antibodies have the same overall basic structure of two identical light chains and two identical heavy chains.

As used herein, “antibody” or “antibodies” may include chimeric and single chain antibodies, as binding fragments of antibodies, such as Fab, Fv fragments or single chain Fv (scFv) fragments, and multimeric forms, e.g., dimeric IgA molecules or pentavalent IgM. Antibodies of the invention may be of human or non-human origin, for example a murine or other rodent-derived antibody, or a chimeric, humanized or reshaped antibody based e.g., on a murine antibody.

A heavy chain of an antibody typically includes a heavy chain variable region (VH) and a heavy chain constant region. The heavy chain constant region usually comprises three domains, referred to as CH1, CH2 and CH3. An antibody light chain includes a light chain variable region

(VL) and a light chain constant region. The light chain constant region includes a single domain, referred to as CL. The VH and VL regions are subdivided into regions of hypervariability (“hypervariable regions”), which may be hypervariable in sequence and/or in looped structure. These regions are also referred to as complementarity determining regions (CDRs), which are interspersed with regions that are more conserved, termed framework regions (FRs). Each VH and VL typically includes three CDRs and four FRs, arranged from the amino terminus to the carboxy terminus in the following order: FR1, CDR1, FR2, CDR2, FR3, CDR3, FR4. Amino acid residues in the variable regions are often numbered using a standardized numbering method known as the Kabat numbering scheme (Kabat et al. (1991) Sequences of Proteins of Immunological Interest, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, Md., USA).

The antibody identifiers found in the tables of this application, e.g., “70701_06A03A”, refer to the specific antibodies.

As used herein, an antibody or fragment “derived from” or “based on” an antibody means that the “derived” antibody comprises, depending on the particular context, one of the following: the heavy chain CDR3 sequence of said specified antibody; the heavy chain CDR3 sequence and the light chain CDR3 sequence of said specified antibody; the heavy chain CDR1, CDR2 and CDR3 sequences and light chain CDR1, CDR2 and CDR3 sequences of said specified antibody; or the heavy chain variable region sequence and the light chain variable region sequence of said specified antibody, or a humanized variant of said heavy chain variable region sequence and/or light chain variable region sequence, or a heavy chain and/or light chain variable region sequence having at least 80%, 85%, 90% or 95% sequence identity, such as at least 96%, 97%, 98% or 99% sequence identity, with the respective heavy chain and light chain variable region sequences.

An antibody that is derived from or based on a specified antibody described herein will generally bind the same CD3 epitope as said specified antibody and will preferably exhibit substantially the same activity as said specified antibody.

The specificity of an antibody's interaction with a target antigen is driven, primarily, by the amino acid residues located in the six CDRs of the heavy and light chain. The amino acid sequences within CDRs are more variable between individual antibodies than sequences outside of CDRs. Since CDR sequences are responsible for most antibody-antigen interactions, antibodies that mimic the properties of a specific naturally occurring antibody, or any specific antibody with

a given amino acid sequence, can be expressed by constructing expression vectors that express CDR sequences from the specific antibody grafted into framework sequences from a different antibody. This permits “humanization” of a non-human antibody, which will nonetheless substantially maintain the binding specificity and affinity of the original antibody. Nonetheless, in preferred aspects, the anti-CD3 antibodies are human antibodies.

A “chimeric antibody” means an antibody that includes one or more regions from one antibody and one or more regions from one or more different antibodies. A “chimeric antibody” is typically an antibody that is partially of human origin and partially of non-human origin. Chimeric antibodies may be preferred over non-human antibodies, as they have been shown to reduce the risk of a human anti-antibody response. A chimeric antibody may include an antibody in which the variable region sequences are murine sequences derived from immunization of a mouse, while the constant region sequences are human. In the case of a chimeric antibody, the non-human parts, which often include the framework regions of the variable region sequences, may be further altered in order to humanize the antibody.

In preferred aspects, the presently disclosed antibodies of the invention are derived from transgenic mice that contain human antibody gene segments, such that the antibodies are human antibodies derived by hybridoma technology from transgenic mice.

The terms “heavy chain variable region sequence” and “light chain variable region sequence” and similar terms as used herein with reference to any specific amino acid sequence encompass not only that specific sequence, but also any recombinant antibodies, human antibodies, including those derived from transgenic mice that contain human antibody gene segments, such that the antibodies are human antibodies derived by hybridoma technology, and/or humanized variants thereof.

As used herein, a reference to a heavy chain variable region sequence or a light chain variable region sequence with a particular minimum level of sequence identity compared to a specified heavy chain or light chain variable region sequence.

A “recombinant antibody” is an antibody that is expressed from a cell or cell line transfected with an expression vector (or possibly more than one expression vector, typically two expression vectors) comprising the coding sequence of the antibody, where said coding sequence is not naturally associated with the cell.

A “vector” is a nucleic acid molecule into which a nucleic acid sequence can be inserted for transport between different genetic environments and/or for expression in a host cell. A vector that carries regulatory elements for transcription of the nucleic acid sequence (at least a suitable promoter) is referred to as an “an expression vector”. The terms “plasmid” and “vector” may be used interchangeably. Expression vectors used in the context of the present invention may be of any suitable type known in the art, for example, a viral vectors or plasmids.

It is well-known in the art that antibodies exist as different isotypes, such as the human isotypes IgG1, IgG2, IgG3, IgG4, IgA1 and IgA2, or the murine isotypes IgG1, IgG2a, IgG2b, IgG3 and IgA. An antibody of the invention may be of any isotype.

In certain aspects, compositions of the invention include antibody compositions comprising a plurality of individual anti-CD3 antibodies, e.g., 2, 3, 4, 5, 6, 7, 8, 9, 10, or 10 or more different CD3 antibodies.

A “CDR” or “complementarity determining region” refers to the “hypervariable” regions found in the variable domains of an antibody that are primarily responsible for determining the antibody's binding specificity. Each of the heavy and light chains of an antibody contain three CDR regions, referred to as CDR1, CDR2 and CDR3, of which CDR3 shows the greatest variability.

An “epitope” describes a part of a larger molecule (e.g., antigen or antigenic site) having antigenic or immunogenic activity in an animal. An epitope having immunogenic activity is a portion of a larger molecule that elicits an antibody response. An epitope having antigenic activity is a portion of a larger molecule to which an antibody immune-specifically binds. Antigenic epitopes are not necessarily immunogenic. An antigen is a substance to which an antibody or antibody fragment specifically binds, such as a toxin, virus, bacteria, protein or DNA. An antigen or antigenic site may have more than one epitope and may be capable of stimulating an immune response.

Epitopes may be linear or conformational. A linear epitope generally consists of about 6 to 10 adjacent amino acids on a protein molecule that are recognized by an antibody. In contrast, a conformational epitope consists of amino acids that are not arranged sequentially, but where an antibody recognizes a particular three-dimensional structure. When a protein molecule folds into a three-dimensional structure, the amino acids forming the epitope are juxtaposed, enabling the antibody to recognize the conformational epitope. In a denatured protein, only linear epitopes

are recognized. A conformational epitope, by definition, must be on the outside of the folded protein.

The term “distinct epitopes” refers to the fact that when two different antibodies of the invention bind distinct epitopes, there is less than 100% competition for antigen binding, preferably less than 80% competition for antigen binding, more preferably less than 50% competition for antigen binding, and most preferably as little competition as possible, such as less than about 25% competition for antigen binding.

Antibodies capable of competing with each other for binding to the same antigen may bind the same or overlapping epitopes or may have a binding site in the close vicinity of one another, so that competition is mainly caused by steric hindrance. An analysis for “distinct epitopes” of antibody pairs may be performed by methods known in the art, for example by way of binding experiments under saturating antibody conditions using either FACS (fluorescence activated cell sorting) or other flow cytometry analysis on cells expressing CD3 and individual fluorescent labeled antibodies, or by Surface Plasmon Resonance (SPR) using CD3 antigens attached to a flow cell surface.

Antibodies binding to different epitopes on the same antigen can have varying effects on the activity of the antigen to which they bind, depending on the location of the epitope. An antibody binding to an epitope in an active site of the antigen may block the function of the antigen completely, whereas another antibody binding at a different epitope may have no or little effect on the activity of the antigen alone. Such antibodies may, however, still activate complement and thereby result in the elimination of the antigen, and may result in synergistic effects when combined with one or more antibodies binding at different epitopes on the same antigen.

“Immunoglobulin” is a collective designation of the mixture of antibodies found in blood or serum, but may also be used to designate a mixture of antibodies derived from other sources.

The term “cognate V_H and V_L coding pair” describes an original pair of V_H and V_L coding sequences contained within or derived from the same antibody-producing cell. Thus, a cognate V_H and V_L pair represents the V_H and V_L pairing originally present in the donor from which such a cell is derived. The term “an antibody expressed from a V_H and V_L coding pair” indicates that an antibody or an antibody fragment is produced from a vector, plasmid or other polynucleotide containing the V_H and V_L coding sequence. When a cognate V_H and V_L coding pair is expressed, either as a complete antibody or as a stable fragment thereof, they preserve the binding affinity

and specificity of the antibody originally expressed from the cell they are derived from. A library of cognate pairs is also termed a repertoire or collection of cognate pairs, and may be kept individually or pooled.

5 By “protein” or “polypeptide” is meant any chain of amino acids, regardless of length or post-translational modification. Proteins can exist as monomers or multimers, comprising two or more assembled polypeptide chains, fragments of proteins, polypeptides, oligopeptides, or peptides.

10 The term “head-to-head promoters” refers to a promoter pair being placed in close proximity so that transcription of two gene fragments driven by the promoters occurs in opposite directions. Head-to-head promoters are also known as bi-directional promoters.

The term “transfection” is herein used as a broad term for introducing foreign DNA into a cell. The term is also meant to cover other functional equivalent methods for introducing foreign DNA into a cell, such as e.g., transformation, infection, transduction or fusion of a donor cell and an acceptor cell.

15 As used herein, CD3 is intended to include variants, isoforms and species homologs of CD3. Preferably, binding of an antibody of the invention to CD3 inhibits the growth of cells expressing CD3. In certain aspects, this inhibition is caused by inhibiting formation of heteromeric complexes between CD3 and other ErbB family members.

20 As used herein, the term “inhibits growth” (e.g., referring to cells) is intended to include any measurable decrease in the proliferation (increase in number of cells) or metabolism of a cell when contacted with an anti-CD3 antibody as compared to the growth of the same cells in the absence of an anti-CD3 antibody, e.g., inhibition of growth of a cell culture by at least about 10%, and preferably more, such as at least about 20% or 30%, more preferably at least about 40% or 50%, such as at least about 60%, 70%, 80%, 90%, 99% or even 100%.

25 The term “treatment” as used herein refers to administration of an anti CD3 antibody, antibody composition of the invention, or composition of immune cells that express or are activated by a CD3 antibody or fragment thereof, in a sufficient amount to ease, reduce, ameliorate or eradicate (cure) symptoms or disease states.

30 The percent identity between two sequences, e.g., variable region sequences, refers to the number of identical positions shared by the sequences (calculated as # of identical positions/total # of positions×100), taking into account gaps that must be introduced for optimal alignment of the

two sequences. The comparison of sequences and determination of percent identity between two sequences may be accomplished using readily available software. Suitable software programs are available from various sources, both for online use and for download, and for alignment of both protein and nucleotide sequences. One suitable program is ClustalW (Thompson et al. (1994) Nucleic Acids Res. 11; 22(22):4673-80), available from www.clustal.org.

An “acceptor human framework” for the purposes herein is a framework comprising the amino acid sequence of a light chain variable domain (VL) framework or a heavy chain variable domain (VH) framework derived from a human immunoglobulin framework or a human consensus framework, as defined below. An acceptor human framework “derived from” a human immunoglobulin framework or a human consensus framework may comprise the same amino acid sequence thereof, or it may contain amino acid sequence changes. In some embodiments, the number of amino acid changes are 10 or less, 9 or less, 8 or less, 7 or less, 6 or less, 5 or less, 4 or less, 3 or less, or 2 or less. In some embodiments, the VL acceptor human framework is identical in sequence to the VL human immunoglobulin framework sequence or human consensus framework sequence.

“Affinity” refers to the strength of the sum total of noncovalent interactions between a single binding site of a molecule (e.g., an antibody) and its binding partner (e.g., an antigen). Unless indicated otherwise, as used herein, “binding affinity” refers to intrinsic binding affinity, which reflects a 1:1 interaction between members of a binding pair (e.g., antibody and antigen). The affinity of a molecule X for its partner Y can generally be represented by the dissociation constant (Kd). Affinity can be measured by common methods known in the art, including those described herein.

An “affinity matured” antibody refers to an antibody with one or more alterations in one or more hypervariable regions, compared to a parent antibody, which does not possess such alterations, such alterations resulting in an improvement in the affinity of the antibody for antigen.

The terms “anti-CD3 antibody” and “an antibody that binds to CD3” refer to an antibody that is capable of binding CD3 with sufficient affinity such that the antibody is useful as a diagnostic and/or therapeutic agent in targeting CD3. In one embodiment, the extent of binding of an anti-CD3 antibody to an unrelated, non-CD3 protein is less than about 10% of the binding of the antibody to CD3 as measured, e.g., by a radioimmunoassay (RIA).

In certain embodiments, an antibody that binds to CD3 has a dissociation constant (Kd) of $\leq 1 \mu\text{M}$, $\leq 100 \text{ nM}$, $\leq 10 \text{ nM}$, $\leq 1 \text{ nM}$, $\leq 0.1 \text{ nM}$, $\leq 0.01 \text{ nM}$, or $\leq 0.001 \text{ nM}$ (e.g., 10^{-6} M or less, e.g., from 10^{-8} M to 10^{-13} M , e.g., from 10^{-9} M to 10^{-13} M). In preferred aspects, this affinity range is the “optimal affinity range”, which retains anti-tumor activity but has reduced toxicity due to reduced cytokine release. In certain preferred aspects, the anti-CD3 antibody has an affinity in the range of 30 nM to 40 nM. In preferred aspects, the anti-CD3 antibody has an affinity in the range of 30-40 nM as measured by alanine scanning of the HC CDR3 of the antibody.

In certain embodiments, an anti-CD3 antibody binds to an epitope of CD3 that is conserved among CD3s from different species.

The term “cluster of differentiation 3” or “CD3,” as used herein, refers to any native CD3 from any vertebrate source, including mammals such as primates (e.g., humans) and rodents (e.g., mice and rats), unless otherwise indicated, including, for example, CD3 ϵ , CD3 γ , CD3 α , and CD3 β chains. The term encompasses “full-length,” unprocessed CD3 (e.g., unprocessed or unmodified CD3 ϵ or CD3 γ), as well as any form of CD3 that results from processing in the cell. The term also encompasses naturally occurring variants of CD3, including, for example, splice variants or allelic variants. CD3 includes, for example, human CD3 ϵ protein (NCBI RefSeq No. NP-000724), which is 207 amino acids in length, and human CD3 γ protein (NCBI RefSeq No. NP-000064), which is 182 amino acids in length.

The “class” of an antibody refers to the type of constant domain or constant region possessed by its heavy chain. There are five major classes of antibodies: IgA, IgD, IgE, IgG, and IgM, and several of these may be further divided into subclasses (isotypes), e.g., IgG1, IgG2, IgG3, IgG4, IgA1, and IgA2. The heavy chain constant domains that correspond to the different classes of immunoglobulins are called α , δ , ϵ , γ , and μ , respectively.

“Effector functions” refer to those biological activities attributable to the Fc region of an antibody, which vary with the antibody isotype. Examples of antibody effector functions include: C1 q binding and complement dependent cytotoxicity (CDC); Fc receptor binding; antibody-dependent cell-mediated cytotoxicity (ADCC); phagocytosis; down regulation of cell surface receptors (e.g., B cell receptor); and B cell activation.

An “effective amount” of a compound, for example, an anti-CD3 antibody of the invention or a composition (e.g., pharmaceutical composition) thereof, is at least the minimum amount required to achieve the desired therapeutic or prophylactic result, such as a measurable

improvement or prevention of a particular disorder (e.g., a cell proliferative disorder, e.g., cancer). An effective amount herein may vary according to factors such as the disease state, age, sex, and weight of the patient, and the ability of the antibody to elicit a desired response in the individual. An effective amount is also one in which any toxic or detrimental effects of the treatment are outweighed by the therapeutically beneficial effects. For prophylactic use, beneficial or desired results include results such as eliminating or reducing the risk, lessening the severity, or delaying the onset of the disease, including biochemical, histological and/or behavioral symptoms of the disease, its complications and intermediate pathological phenotypes presenting during development of the disease. For therapeutic use, beneficial or desired results include clinical results such as decreasing one or more symptoms resulting from the disease, increasing the quality of life of those suffering from the disease, decreasing the dose of other medications required to treat the disease, enhancing effect of another medication such as via targeting, delaying the progression of the disease, and/or prolonging survival. In the case of cancer or tumor, an effective amount of the drug may have the effect in reducing the number of cancer cells; reducing the tumor size; inhibiting (i.e., slow to some extent or desirably stop) cancer cell infiltration into peripheral organs; inhibit (i.e., slow to some extent and desirably stop) tumor metastasis; inhibiting to some extent tumor growth; and/or relieving to some extent one or more of the symptoms associated with the disorder. An effective amount can be administered in one or more administrations. For purposes of this invention, an effective amount of drug, compound, or pharmaceutical composition is an amount sufficient to accomplish prophylactic or therapeutic treatment either directly or indirectly. As is understood in the clinical context, an effective amount of a drug, compound, or pharmaceutical composition may or may not be achieved in conjunction with another drug, compound, or pharmaceutical composition. Thus, an "effective amount" may be considered in the context of administering one or more therapeutic agents, and a single agent may be considered to be given in an effective amount if, in conjunction with one or more other agents, a desirable result may be or is achieved.

The term "Fc region" herein is used to define a C-terminal region of an immunoglobulin heavy chain that contains at least a portion of the constant region. The term includes native sequence Fc regions and variant Fc regions. In one embodiment, a human IgG heavy chain Fc region extends from Cys226, or from Pro230, to the carboxyl-terminus of the heavy chain. However, the C-terminal lysine (Lys447) of the Fc region may or may not be present. Unless

otherwise specified herein, numbering of amino acid residues in the Fc region or constant region is according to the EU numbering system, also called the EU index, as described in Kabat et al., Sequences of Proteins of Immunological Interest, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, Md., 1991.

5 “Framework” or “FR” refers to variable domain residues other than hypervariable region residues. The FR of a variable domain generally consists of four FR domains: FR1, FR2, FR3, and FR4. Accordingly, the HVR and FR sequences generally appear in the following sequence in VH (or VL): FR1-H1(L)-FR2-H2(L2)-FR3-H3(L3)-FR4.

10 The terms “full-length antibody,” “intact antibody,” and “whole antibody” are used herein interchangeably to refer to an antibody having a structure substantially similar to a native antibody structure or having heavy chains that contain an Fc region as defined herein.

15 A “human antibody” is one which possesses an amino acid sequence which corresponds to that of an antibody produced by a human or a human cell or derived from a non-human source that utilizes human antibody repertoires or other human antibody-encoding sequences. This definition of a human antibody specifically excludes a humanized antibody comprising non-human antigen-binding residues. Human antibodies can be produced using various techniques known in the art, including phage-display libraries. Hoogenboom and Winter, *J. Mol. Biol.*, 227:381 (1991); Marks et al., *J. Mol. Biol.*, 222:581 (1991). Also available for the preparation of human monoclonal antibodies are methods described in Cole et al., *Monoclonal Antibodies and Cancer Therapy*, Alan R. Liss, p. 77 (1985); Boerner et al., *J. Immunol.* 147(1):86-95 (1991). See also van Dijk and van de Winkel, *Curr. Opin. Pharmacol.*, 5: 368-74 (2001). Human antibodies can be prepared by administering the antigen to a transgenic animal that has been modified to produce such antibodies in response to antigenic challenge, but whose endogenous loci have been disabled, e.g., immunized xenomice (see, e.g., U.S. Pat. Nos. 6,075,181 and 6,150,584 regarding XENOMOUSE™ technology). See also, for example, Li et al., *Proc. Natl. Acad. Sci. USA*, 103:3557-3562 (2006) regarding human antibodies generated via a human B-cell hybridoma technology.

20 For example, in preferred aspects, technologies such as AlivaMab Mouse technology are used to produce the anti-CD3 antibodies described herein. Generally, AlivaMab Mouse technology is used to generate panels of monoclonal antibodies (mAbs) against a human antigen of interest, such as those expressed by or associated with tumor cells, e.g., variants of HER2.

The AlivaMab Mouse is a transgenic mouse that produces chimeric human-mouse monoclonal antibodies comprising fully human Fab and upper hinge regions and mouse middle hinge and Fc regions. Optimized constant domains facilitate the generation and identification of antibodies that retain structure-function characteristics. Antibodies produced using AlivaMab
5 Mouse technology possess biophysical properties, which are predictive and comparable to that of fully human antibody counterparts.

Antibodies produced by AlivaMab Kappa Mice include a chimeric immunoglobulin heavy (IgH) chain and a human immunoglobulin kappa (IgK) light chain. Antibodies produced by AlivaMab Lambda Mice include a chimeric IgH chain and a human immunoglobulin lambda (IgK)
10 light chain. The chimeric IgH chain of the AlivaMab Mouse antibodies include a human variable region comprising a human variable heavy (VH) domain, a human diversity heavy (DH) domain, and a human joining heavy (JH) domain, a human constant heavy 1 (CH1) domain, a human upper hinge region (except for O μ , which is naturally missing an upper hinge region), a mouse middle hinge region, a mouse CH2 domain, and a mouse CH3 domain.

When a lead candidate antibody is discovered, the human heavy chain variable region is readily appended to a fully human constant region while maintaining the antigen-binding characteristics of the parent chimeric antibody that were developed *in vivo* in the AlivaMab Mouse. In one embodiment, the human heavy chain variable region, CH1 and, optionally, upper hinge region of the chimeric antibody are appended to human hinge, a human CH2 domain and a human
20 CH3 domain in order to produce a fully human antibody.

Portions of variable regions from the antibodies produced from AlivaMab Mouse technology may include all or a combination of the complementarity determining regions (CDRs) of the VH and/or VL. The variable regions may be formatted with constant regions, either native or modified for various desired effector functions, in a standard antibody structure (two heavy chains with two light chains). The variable regions may also be formatted as multi-specific antibodies, e.g., bispecific antibodies binding to two different epitopes or to two different antigens. The variable regions may also be formatted as antibody fragments, e.g., single-domain antibodies comprising a single VH or VL, Fabs or Fab'2. The antibodies may also be used as antibody-drug conjugates, or carry other additions such as small molecule toxins, biologic toxins, cytokines,
30 oligopeptides, or RNAs to increase therapeutic modality and/or increase safety.

Methods for producing the anti-CDR3 antibodies of the invention using AlivaMab mouse technology may include immunizing AlivaMab Kappa Mice and AlivaMab Lambda Mice with an antigen of interest. Generally, within two weeks, the mice are sacrificed and terminal materials collected. Spleens and lymph nodes may be prepared and fused with myeloma cells (such as CRL-
5 2016 cells) using a PEG based method as generally described in “Antibodies: A Laboratory Manual” (Harlow and Lane 1988 CSH Press) to establish hybridomas.

Hybridomas may be grown in 384-well tissue culture plates and supernatants from individual wells were screened by ELISA for production of antibodies recognizing the antigen of interest. Positive wells are then transferred to 48-well plates, expanded, and supernatants were
10 collected for antigen binding confirmation by ELISA. Positive supernatants may also be counter-screened against a non-related histidine-tagged protein. Hybridoma lines each from AlivaMab Kappa Mice and AlivaMab Lambda Mice are confirmed to bind to the antigen specifically by ELISA and are picked at random and single-cell cloned into 96-well plates. They are grown into colonies and the supernatant from these individual colonies is screened by ELISA to re-confirm
15 monoclonal antibody binding to the antigen of interest. These supernatants are then screened by FACS to confirm binding to the native antigen expressed on cells.

AlivaMab Mouse technology and methods for producing antibodies using such technologies can be found in WO 2010/039900 and WO 2011/123708, which are incorporated herein in their entirety.

A “human consensus framework” is a framework which represents the most commonly occurring amino acid residues in a selection of human immunoglobulin VL or VH framework sequences. Generally, the selection of human immunoglobulin VL or VH sequences is from a subgroup of variable domain sequences. Generally, the subgroup of sequences is a subgroup as in
20 Kabat et al., Sequences of Proteins of Immunological Interest, Fifth Edition, NIH Publication 91-3242, Bethesda Md. (1991), vols. 1-3. In one embodiment, for the VL, the subgroup is subgroup kappa I as in Kabat et al., supra. In one embodiment, for the VH, the subgroup is subgroup III as
25 in Kabat et al., supra.

A “humanized” antibody refers to a chimeric antibody comprising amino acid residues from non-human HVRs and amino acid residues from human FRs. In certain embodiments, a
30 humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the HVRs (e.g., CDRs) correspond to those of a non-

human antibody, and all or substantially all of the FRs correspond to those of a human antibody. A humanized antibody optionally may comprise at least a portion of an antibody constant region derived from a human antibody. A “humanized form” of an antibody, e.g., a non-human antibody, refers to an antibody that has undergone humanization.

5 To “humanize” an antibody means that an antibody of wholly or partially of non-human origin, for example a murine antibody obtained from immunization of mice with an antigen of interest or a chimeric antibody based on such a murine antibody, can have amino acids replaced, particularly in the framework regions and constant domains of the heavy and light chains, to avoid or minimize an immune response in humans. It is known that all antibodies have the potential for
10 eliciting a human anti-antibody response, which correlates to some extent with the degree of “humanness” of the antibody in question.

Non-human antibodies tend to be more immunogenic than human antibodies. Chimeric antibodies, where the foreign (usually rodent) constant regions have been replaced with sequences of human origin, have been shown to be less immunogenic than antibodies of fully foreign origin,
15 and the most development efforts in therapeutic antibodies are trending towards the use of humanized or fully human antibodies. Preferably, chimeric antibodies or other antibodies of non-human origin are humanized to reduce the risk of a human anti-antibody response. For chimeric antibodies, humanization may include, for example, modification of the framework regions of the variable region sequences. Amino acid residues of a CDR may often not be altered during
20 humanization, although in certain cases it may be desirable to alter individual CDR amino acid residues, for example to remove a glycosylation site, a deamidation site or an undesired cysteine residue.

Numerous methods for humanization of an antibody sequence are known in the art. A commonly used method is CDR grafting, which may involve identification of human germline
25 gene counterparts to murine variable region genes and grafting of the murine CDR sequences into this framework. Since CDR grafting reduces the chance for binding specificity and affinity and the biological activity of a CDR grafted non-human antibody, back mutations are often introduced at selected positions of the CDR grafted antibody to retain the binding specificity and affinity. Amino acid residues that for back mutations may include those that are located at the surface of
30 an antibody molecule. Another humanization technique for CDR grafting and back mutation is

resurfacing, in which non-surface exposed residues of non-human origin are retained, while surface residues are altered to human variants.

Humanized antibodies and methods of making them are reviewed, e.g., in Almagro and Fransson, *Front. Biosci.* 13:1619-1633 (2008), and are further described, e.g., in Riechmann et al., *Nature* 332:323-329 (1988); Queen et al., *Proc. Nat'l Acad. Sci. USA* 86:10029-10033 (1989); U.S. Pat. Nos. 5,821,337, 7,527,791, 6,982,321, and 7,087,409; Kashmiri et al., *Methods* 36:25-34 (2005) (describing specificity determining region (SDR) grafting); Padlan, *Mol. Immunol.* 28:489-498 (1991) (describing "resurfacing"); Dall'Acqua et al., *Methods* 36:43-60 (2005) (describing "FR shuffling"); and Osbourn et al., *Methods* 36:61-68 (2005) and Klimka et al., *Br. J. Cancer*, 83:252-260 (2000) (describing the "guided selection" approach to FR shuffling).

Human framework regions that may be used for humanization include but are not limited to: framework regions selected using the "best-fit" method (see, e.g., Sims et al. *J. Immunol.* 151:2296 (1993)); framework regions derived from the consensus sequence of human antibodies of a particular subgroup of light or heavy chain variable regions (see, e.g., Carter et al. *Proc. Natl. Acad. Sci. USA*, 89:4285 (1992); and Presta et al. *J. Immunol.*, 151:2623 (1993)); human mature (somatically mutated) framework regions or human germline framework regions (see, e.g., Almagro and Fransson, *Front. Biosci.* 13:1619-1633 (2008)); and framework regions derived from screening FR libraries (see, e.g., Baca et al., *J. Biol. Chem.* 272:10678-10684 (1997) and Rosok et al., *J. Biol. Chem.* 271:22611-22618 (1996)).

The term "hypervariable region" or "HVR" as used herein refers to each of the regions of an antibody variable domain which are hypervariable in sequence ("complementarity determining regions" or "CDRs") and/or form structurally defined loops ("hypervariable loops") and/or contain the antigen-contacting residues ("antigen contacts"). Generally, antibodies comprise six HVRs: three in the VH (H1, H2, H3), and three in the VL(L1, L2, L3). Unless otherwise indicated, HVR residues and other residues in the variable domain (e.g., FR residues) are numbered herein according to Kabat et al., *supra*.

An "immunoconjugate" is an antibody conjugated to one or more heterologous molecule(s), including but not limited to a cytotoxic agent.

A "subject" or an "individual" is a mammal. Mammals include, but are not limited to, domesticated animals (e.g., cows, sheep, cats, dogs, and horses), primates (e.g., humans and non-

human primates such as monkeys), rabbits, and rodents (e.g., mice and rats). In certain embodiments, the subject or individual is a human.

An “isolated” antibody is one which has been separated from a component of its natural environment. In some embodiments, an antibody is purified to greater than 95% or 99% purity as determined by, for example, electrophoretic (e.g., SDS-PAGE, isoelectric focusing (IEF), capillary electrophoresis) or chromatographic (e.g., ion exchange or reverse phase HPLC). For review of methods for assessment of antibody purity, see, e.g., Flatman et al., *J. Chromatogr. B* 848:79-87 (2007).

An “isolated” nucleic acid refers to a nucleic acid molecule that has been separated from a component of its natural environment. An isolated nucleic acid includes a nucleic acid molecule contained in cells that ordinarily contain the nucleic acid molecule, but the nucleic acid molecule is present extrachromosomally or at a chromosomal location that is different from its natural chromosomal location.

“Isolated nucleic acid encoding an anti-CD3 antibody” refers to one or more nucleic acid molecules encoding antibody heavy and light chains (or fragments thereof), including such nucleic acid molecule(s) in a single vector or separate vectors, and such nucleic acid molecule(s) present at one or more locations in a host cell.

The term “monoclonal antibody” as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical and/or bind the same epitope, except for possible variant antibodies, e.g., containing naturally occurring mutations or arising during production of a monoclonal antibody preparation, such variants generally being present in minor amounts. In contrast to polyclonal antibody preparations, which typically include different antibodies directed against different determinants (epitopes), each monoclonal antibody of a monoclonal antibody preparation is directed against a single determinant on an antigen. Thus, the modifier “monoclonal” indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies and is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies to be used in accordance with the present invention may be made by a variety of techniques, including but not limited to the hybridoma method, recombinant DNA methods, phage-display methods, and methods utilizing transgenic

animals containing all or part of the human immunoglobulin loci, such methods and other exemplary methods for making monoclonal antibodies being described herein.

“Native antibodies” refer to naturally occurring immunoglobulin molecules with varying structures. For example, native IgG antibodies are heterotetrameric glycoproteins of about 150,000
5 Daltons, composed of two identical light chains and two identical heavy chains that are disulfide-bonded. From N- to C-terminus, each heavy chain has a variable region (VH), also called a variable heavy domain or a heavy chain variable domain, followed by three constant domains (CH1, CH2, and CH3). Similarly, from N- to C-terminus, each light chain has a variable region (VL), also called a variable light domain or a light chain variable domain, followed by a constant light (CL)
10 domain. The light chain of an antibody may be assigned to one of two types, called kappa (K) and lambda (A), based on the amino acid sequence of its constant domain.

The term “package insert” is used to refer to instructions customarily included in commercial packages of therapeutic products, that contain information about the indications, usage, dosage, administration, combination therapy, contraindications, and/or warnings
15 concerning the use of such therapeutic products.

The term “protein,” as used herein, refers to any native protein from any vertebrate source, including mammals such as primates (e.g., humans) and rodents (e.g., mice and rats), unless otherwise indicated. The term encompasses “full-length,” unprocessed protein as well as any form of the protein that results from processing in the cell. The term also encompasses naturally
20 occurring variants of the protein, e.g., splice variants or allelic variants.

The term “variable region” or “variable domain” refers to the domain of an antibody heavy or light chain that is involved in binding the antibody to antigen. The variable domains of the heavy chain and light chain (VH and VL, respectively) of a native antibody generally have similar structures, with each domain comprising four conserved framework regions (FRs) and three
25 hypervariable regions (HVRs). (*See, e.g., Kindt et al. Kuby Immunology, 6th ed., W.H. Freeman and Co., page 91 (2007).*) A single VH or VL domain may be sufficient to confer antigen-binding specificity. Furthermore, antibodies that bind a particular antigen may be isolated using a VH or VL domain from an antibody that binds the antigen to screen a library of complementary VL or VH domains, respectively. *See, e.g., Portolano et al., J. Immunol. 150:880-887 (1993); Clarkson
30 et al., Nature 352:624-628 (1991).*

As used herein, “administering” is meant a method of giving a dosage of a compound (e.g., an anti-CD3 antibody of the invention or a nucleic acid encoding an anti-CD3 antibody of the invention) or a composition (e.g., a pharmaceutical composition, e.g., a pharmaceutical composition including an anti-CD3 antibody of the invention) to a subject. The compositions
5 utilized in the methods described herein can be administered, for example, intramuscularly, intravenously, intradermally, percutaneously, intraarterially, intraperitoneally, intralesionally, intracranially, intraarticularly, intraprostatically, intrapleurally, intratracheally, intranasally, intravitreally, intravaginally, intrarectally, topically, intratumorally, peritoneally, subcutaneously, subconjunctivally, intravesicularly, mucosally, intrapericardially, intraumbilically, intraocularly,
10 orally, topically, locally, by inhalation, by injection, by infusion, by continuous infusion, by localized perfusion bathing target cells directly, by catheter, by lavage, in cremes, or in lipid compositions. The method of administration can vary depending on various factors (e.g., the compound or composition being administered and the severity of the condition, disease, or disorder being treated).

As used herein, “somatic hypermutation” or “SHM” refers to the mutation of a polynucleotide sequence initiated by, or associated with the action of the Activation-Induced
15 Cytidine Deaminase (AID), a functional AID mutant, uracil glycosylase and/or error prone polymerases on that polynucleotide sequence. As used herein, the term includes mutagenesis that occurs as a consequence of the error prone repair, including mutagenesis mediated by the mismatch
20 repair machinery and related enzymes.

SHM is generally initiated by targeting AID to rearranged V(D)J and switch regions of Ig genes. The mutation rate of this programmed mutagenesis is a million-fold higher than the non-AID targeted genome of B cells. AID is a processive enzyme that binds single-stranded DNA and deaminates cytosines in DNA. Cytosine deamination generates highly mutagenic deoxy-uracil (U)
25 in the DNA of the Ig loci. Mutagenic processing of uracil through the DNA damage response produces the entire spectrum of base substitutions, which characterizes SHM at and around an initial U lesion. At least five, identified mutagenic DNA damage response pathways are known to generate a well-defined SHM spectrum of C/G transitions, C/G transversions, and A/T mutations around this initial lesion. These pathways include (1) replication opposite template U generates
30 transitions at C/G, (2) UNG2-dependent translesion synthesis (TLS) generates transversions at C/G, (3) a hybrid pathway comprising non-canonical mismatch repair (ncMMR) and UNG2-

dependent TLS generates transversions at C/G, (4) ncMMR generates mutations at A/T, and (5) UNG2- and PCNA Ubiquitination (PCNA-Ub)-dependent mutations at A/T. Specific strand-biases of SHM spectra arise as a consequence of a biased AID targeting, ncMMR, and anti-mutagenic repriming. By elucidating the amino acid and/or nucleotide sequences of the CDR3 variable regions and/or one of CDR3 heavy chain (HC) and light chain (LC or λ) variable regions of the anti-CD3 antibodies disclosed herein, the present inventors identified a series of “clusters” or “motifs” within the sequences. These clusters represent convergent somatic hypermutations (SHM) in the variable region sequences. Clustering may provide insight into the functionally related sequences and the diversity of the total population of antibodies and their variable region sequences. Sequences that are descendants from the same parent B cell or convergently evolved the sequences in the same cluster should be functionally more related than sequences belonging to other clusters. Convergent SHMs are likely functionally related mutations, e.g., they share a specific affinity for CD3. These SHM may inform the development of recombinant anti-CD3 antibodies with improved properties, such as specific binding for CD3 fragments.

15

Selected Embodiments

One aspect of the invention relates to various novel anti-CD3 antibodies and fragments thereof.

The presently disclosed antibodies may provide treatments that are far more effective than current therapies. The presently disclosed CD3 antibodies of the invention may be included as part of a treatment regime, which may include, for example, providing two or more such antibodies, and/or in combination with other treatments such as chemotherapy.

In one aspect, the invention relates to novel CD3 antigen binding peptides, which may be antibodies and/or fragments thereof. In certain aspects, the antibodies and/or fragments thereof bind to a CD3 fragment having an amino acid sequence of SEQ ID NO: 1, and/or a sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 1. In certain aspects, the antibodies and/or fragments thereof bind to a CD3 fragment having an amino acid sequence of SEQ ID NO: 234, and/or a sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 234. In certain aspects, the antibodies and/or fragments thereof bind to a CD3 fragment having an amino acid sequence of SEQ ID NO: 235, and/or a

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sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 235.

In certain aspects, the present invention provides an anti-CD3 antibody or antibody fragment comprising a heavy chain CDR3 sequence (VH) comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59. Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

Additionally or alternatively, the present invention provides an anti-CD3 antibody or antibody fragment comprising a heavy chain CDR3 sequence (VH) comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59 and a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175. Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

Additionally or alternatively, an anti-CD3 antibody or antibody fragment of the present invention may comprise a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175 and a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

Additionally or alternatively, an anti-CD3 antibody or antibody of the present comprises a heavy chain variable region and/or a light chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.1 and/or Clust 8.2, as set forth in Tables 3-4.

In certain aspects, the present invention provides compositions, including therapeutic compositions, comprising an anti-CD3 antibody or antibody fragment as described herein. In certain aspects, the present invention provides compositions, including therapeutic compositions, two or more of the CD3 antibodies disclosed herein. Certain compositions of the invention include a plurality of different CD3 antibodies, as disclosed herein, wherein each different antibody binds to a distinct CD3 epitope or fragment.

In certain aspects, the present invention provides methods for treating breast cancer using compositions comprising one or more CD3 antibodies, as described herein. In certain aspects, administration of such a composition results in reduced CD3 and /or HER2 expression, CD3 and /or HER2 receptor internalization, and/or ligand-induced phosphorylation of HER3.

In certain aspects, the present invention provides immunoconjugates and/or compositions comprising such immunoconjugates, wherein said immunoconjugates comprise a CD3 antibody of the invention conjugated to another therapeutic agent, such as an anti-cancer agent. The invention

further provides immunoconjugates comprising two or more different CD3 antibodies or fragment thereof, wherein each different CD3 antibody or fragment targets a different CD3 fragment or epitope.

5 A further aspect of the invention relates to a nucleic acid molecule having a nucleotide sequence that encodes a CD3 antibody or fragment thereof, as disclosed herein, as well as expression vectors comprising such a polynucleotide and host cells that have been transfected with such an expression vector.

Aspects of the invention also provide methods for producing the CD3 antibodies, fragments thereof, and compositions of the invention.

10 The present invention also provides methods for treating a disease in a human or animal subject, in particular treatment of cancer in humans, by administering an anti-CD3 antibody or composition of the invention to said subject. The invention also includes the use of one or more anti-CD3 antibodies of the invention for preparation of a medicament for use in treating a disease in a human or animal, in particular for the treatment of cancer in humans.

15 Another embodiment of this aspect of the invention relates to an antibody composition comprising at least first and second anti-CD3 antibodies, wherein the first and second antibodies bind distinct epitopes of CD3, said first and second antibodies independently comprising a heavy chain CDR3 sequence (VH) comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least
20 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59. Additionally or alternatively, said first and second anti-CD3 antibodies comprise a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least
25 99% sequence identity with any of SEQ ID NOS: 60-117.

Another aspect of the invention relates to nucleic acid molecules comprising a nucleotide sequence that encodes an antibody, VL variable region sequence, VH variable region sequence, VL CDR3 sequence, and/or VH CDR3 sequence as set forth herein, and/or a sequence having an amino acid sequence comprising a sequence having at least 85%, at least 90% at least 95%; at least
30 96%, at least 97%, at least 98% or at least 99% with any of SEQ ID NOS: 2-233.

A further aspect of the invention relates to an expression vector comprising a nucleic acid molecule as defined above. As noted above, expression vectors for use in the context of the present invention may be of any suitable type known in the art, e.g., a plasmid or a viral vector.

5 A still further aspect of the invention relates to a host cell comprising a nucleic acid molecule as defined above, wherein said host cell is capable of expressing an anti-CD3 antibody encoded by said nucleic acid molecule.

In some embodiments, an antibody provided herein has a dissociation constant (Kd) of ≤ 1 μM , ≤ 100 nM, ≤ 10 nM, ≤ 1 nM, ≤ 0.1 nM, ≤ 0.01 nM, or ≤ 0.001 nM (e.g., 10^{-6}M or less, e.g., from 10^{-8}M to 10^{-13}M , e.g., from 10^{-9}M to 10^{-13}M). In preferred aspects, this affinity range is the
10 “optimal affinity range”, which retains anti-tumor activity but has reduced toxicity due to reduced cytokine release. In certain preferred aspects, the anti-CD3 antibody has an affinity in the range of 30 nM to 40 nM. In preferred aspects, the anti-CD3 antibody has an affinity in the range of 30-40 nM as measured by alanine scanning of the HC CDR3 of the antibody.

In some embodiments, Kd is measured by a radiolabeled antigen binding assay (RIA). In
15 some embodiments, an RIA is performed with the Fab version of an antibody of interest and its antigen. For example, solution binding affinity of Fabs for antigen is measured by equilibrating Fab with a minimal concentration of (125I)-labeled antigen in the presence of a titration series of unlabeled antigen, then capturing bound antigen with an anti-Fab antibody-coated plate (see, e.g., Chen et al., J. Mol. Biol. 293:865-881(1999)).

20 In some embodiments, Kd is measured using a BIACORE® surface plasmon resonance assay. For example, an assay using a BIACORE®-2000 or a BIACORE®-3000 (BIAcore, Inc., Piscataway, N.J.) is performed at 25° C. with immobilized antigen CM5 chips at ~10 response units (RU). In one embodiment, carboxymethylated dextran biosensor chips (CM5, BIACORE, Inc.) are activated with N-ethyl-N'-(3-dimethylaminopropyl)-carbodiimide hydrochloride (EDC) and
25 N-hydroxysuccinimide (NHS) according to the supplier's instructions. Antigen is diluted with 10 mM sodium acetate, pH 4.8, to 5 g/ml (0.2 M) before injection at a flow rate of 5 $\mu\text{l}/\text{minute}$ to achieve approximately 10 response units (RU) of coupled protein. Following the injection of antigen, 1 M ethanolamine is injected to block unreacted groups. For kinetics measurements, two-fold serial dilutions of Fab (0.78 nM to 500 nM) are injected in PBS with 0.05% polysorbate 20
30 (TWEEN-20™) surfactant (PBST) at 25° C. at a flow rate of approximately 25 $\mu\text{l}/\text{min}$. Association rates (kon) and dissociation rates (KO) are calculated using a simple one-to-one Langmuir binding

model (BIAcore® Evaluation Software version 3.2) by simultaneously fitting the association and dissociation sensorgrams. The equilibrium dissociation constant (K_d) is calculated as the ratio k_{on}/k_{off} . See, for example, Chen et al., *J. Mol. Biol.* 293:865-881 (1999). If the on-rate exceeds $10^6 \text{ M}^{-1}\text{s}^{-1}$ by the surface plasmon resonance assay above, then the on-rate can be determined
5 by using a fluorescent quenching technique that measures the increase or decrease in fluorescence emission intensity (excitation=295 nm; emission=340 nm, 16 nm band-pass) at 25° C. of a 20 nM anti-antigen antibody (Fab form) in PBS, pH 7.2, in the presence of increasing concentrations of antigen as measured in a spectrometer, such as a stop-flow equipped spectrophotometer (Aviv Instruments) or a 8000-series SLM-AMINCO™ spectrophotometer (ThermoSpectronic) with a
10 stirred cuvette.

In preferred aspects, the modality to estimate affinity is production of a monovalent anti-CD3 antibody followed by titration on live CD3 expressing cells and determination of MFI by flow cytometry to determine an EC_{50} value. Advantageously, this may represent an exacting context (monovalent and CD3 on cells) in which a therapeutic based on the anti-CD3 antibodies
15 of the invention is used.

In some embodiments, an antibody provided herein is an antibody fragment. Antibody fragments include, but are not limited to, Fab, Fab', Fab'-SH, F(ab')₂, Fv, and scFv fragments, and other fragments described below. For a review of certain antibody fragments, see Hudson et al. *Nat. Med.* 9:129-134 (2003). For a review of scFv fragments, see, e.g., Pluckthun, in *The Pharmacology of Monoclonal Antibodies*, vol. 113, Rosenberg and Moore eds., (Springer-Verlag, New York), pp. 269-315 (1994); see also WO 93/16185; and U.S. Pat. Nos. 5,571,894 and 5,587,458. For discussion of Fab and F(ab')₂ fragments comprising salvage receptor binding epitope residues and having increased in vivo half-life, see U.S. Pat. No. 5,869,046.
20

Diabodies are antibody fragments with two antigen-binding sites that may be bivalent or bispecific. See, e.g., EP 404,097; WO 1993/01161; Hudson et al. *Nat. Med.* 9:129-134 (2003); and Hollinger et al. *Proc. Natl. Acad. Sci. USA* 90: 6444-6448 (1993). Triabodies and tetrabodies are also described in Hudson et al. *Nat. Med.* 9:129-134 (2003).
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Single-domain antibodies are antibody fragments comprising all or a portion of the heavy chain variable domain or all or a portion of the light chain variable domain of an antibody. In certain embodiments, a single-domain antibody is a human single-domain antibody (Domantis, Inc., Waltham, Mass.; see, e.g., U.S. Pat. No. 6,248,516 B1).
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Antibody fragments can be made by various techniques, including but not limited to proteolytic digestion of an intact antibody as well as production by recombinant host cells (e.g., *E. coli* or phage).

In some embodiments, an antibody provided herein is a chimeric antibody. Certain
5 chimeric antibodies are described, e.g., in U.S. Pat. No. 4,816,567; and Morrison et al. Proc. Natl. Acad. Sci. USA, 81:6851-6855 (1984)). In one example, a chimeric antibody comprises a non-human variable region (e.g., a variable region derived from a mouse, rat, hamster, rabbit, or non-human primate, such as a monkey) and a human constant region. In a further example, a chimeric antibody is a “class switched” antibody in which the class or subclass has been changed from that
10 of the parent antibody. Chimeric antibodies include antigen-binding fragments thereof.

In preferred embodiments, an antibody provided herein is a human antibody. Human antibodies can be produced using various techniques known in the art. Human antibodies are described generally in van Dijk and van de Winkel, *Curr. Opin. Pharmacol.* 5: 368-74 (2001) and Lonberg, *Curr. Opin. Immunol.* 20:450-459 (2008).

The human antibodies may be prepared by administering an immunogen to a transgenic
15 animal that has been modified to produce intact human antibodies or intact antibodies with human variable regions in response to antigenic challenge. Such animals typically contain all or a portion of the human immunoglobulin loci, which replace the endogenous immunoglobulin loci, or which are present extrachromosomally or integrated randomly into the animal's chromosomes. In such
20 transgenic mice, the endogenous immunoglobulin loci have generally been inactivated. For a review of methods for obtaining human antibodies from transgenic animals, see Lonberg, *Nat. Biotech.* 23:1117-1125 (2005). See also, e.g., U.S. Pat. Nos. 6,075,181 and 6,150,584 describing XENOMOUSE™ technology; U.S. Pat. No. 5,770,429 describing HUMAB® technology; U.S. Pat. No. 7,041,870 describing K-M MOUSE® technology, and U.S. Patent Application
25 Publication No. US 2007/0061900, describing VELOCIMOUSE® technology). Human variable regions from intact antibodies generated by such animals may be further modified, e.g., by combining with a different human constant region.

Human antibodies can also be made by hybridoma-based methods. Human myeloma and mouse-human heteromyeloma cell lines for the production of human monoclonal antibodies have
30 been described. (See, e.g., Kozbor *J. Immunol.*, 133: 3001 (1984); Brodeur et al., *Monoclonal Antibody Production Techniques and Applications*, pp. 51-63 (Marcel Dekker, Inc., New York,

1987); and Boerner et al., J. Immunol., 147: 86 (1991).) Human antibodies generated via human B-cell hybridoma technology are also described in Li et al., Proc. Natl. Acad. Sci. USA, 103: 3557-3562 (2006). Additional methods include those described, for example, in U.S. Pat. No. 7,189,826 (describing production of monoclonal human IgM antibodies from hybridoma cell lines) and Ni, Xiandai Mianyixue, 26(4):265-268 (2006) (describing human-human hybridomas). Human hybridoma technology (Trioma technology) is also described in Vollmers and Brandlein, Histology and Histopathology, 20(3):927-937 (2005) and Vollmers and Brandlein, Methods and Findings in Experimental and Clinical Pharmacology, 27(3):185-91 (2005).

For example, in preferred aspects, technologies such as AlivaMab Mouse technology are used to produce the anti-CD3 antibodies described herein. AlivaMab Mouse technology is used to generate panels of monoclonal antibodies (mAbs) against CD3.

The AlivaMab Mouse is a transgenic mouse that produces chimeric human-mouse monoclonal antibodies comprising fully human Fab and upper hinge regions and mouse middle hinge and Fc regions. Optimized constant domains facilitate the generation and identification of antibodies that retain structure-function characteristics. Antibodies of the invention produced using AlivaMab Mouse technology possess biophysical properties, which are predictive and comparable to that of fully human antibody counterparts.

Antibodies of the invention may be produced by AlivaMab Kappa Mice, and may include a chimeric immunoglobulin heavy (IgH) chain and a human immunoglobulin kappa (IgK) light chain. The antibodies of the invention produced by AlivaMab Lambda Mice may include a chimeric IgH chain and a human immunoglobulin lambda (IgK) light chain. The chimeric IgH chain of the AlivaMab Mouse anti-CD3 antibodies may include a human variable region comprising a human variable heavy (VH) domain, a human diversity heavy (DH) domain, and a human joining heavy (JH) domain, a human constant heavy 1 (CH1) domain, a human upper hinge region (except for O_μ, which is naturally missing an upper hinge region), a mouse middle hinge region, a mouse CH2 domain, and a mouse CH3 domain.

When an anti-CD3 antibody is discovered, the human heavy chain variable region is readily appended to a fully human constant region while maintaining the antigen-binding characteristics of the parent chimeric antibody that were developed *in vivo* in the AlivaMab Mouse. In one embodiment, the human heavy chain variable region, CH1 and, optionally, upper hinge region of

the chimeric antibody are appended to human hinge, a human CH2 domain and a human CH3 domain in order to produce a fully human anti-CD3 antibody as disclosed herein.

Portions of variable regions from the antibodies produced from AlivaMab Mouse technology may include all or a combination of the complementarity determining regions (CDRs) of the VH and/or VL. The variable regions may be formatted with constant regions, either native or modified for various desired effector functions, in a standard antibody structure (two heavy chains with two light chains). The variable regions may also be formatted as multi-specific antibodies, e.g., bispecific antibodies binding to two different epitopes or to two different antigens. The variable regions may also be formatted as antibody fragments, e.g., single-domain antibodies comprising a single VH or VL, Fabs or Fab'2. The antibodies may also be used as antibody-drug conjugates, or carry other additions such as small molecule toxins, biologic toxins, cytokines, oligopeptides, or RNAs to increase therapeutic modality and/or increase safety.

Methods for producing the anti-CDR3 antibodies of the invention using AlivaMab mouse technology may include immunizing AlivaMab Kappa Mice and AlivaMab Lambda Mice with an antigen. Generally, within two weeks, the mice are sacrificed and terminal materials collected. Splens and lymph nodes may be prepared and fused with myeloma cells (such as CRL-2016 cells) using a PEG based method as generally described in "Antibodies: A Laboratory Manual" (Harlow and Lane 1988 CSH Press) to establish hybridomas.

Hybridomas may be grown in 384-well tissue culture plates and supernatants from individual wells were screened by ELISA for production of antibodies recognizing the antigen of interest. Positive wells are then transferred to 48-well plates, expanded, and supernatants were collected for antigen binding confirmation by ELISA. Positive supernatants may also be counter-screened against a non-related histidine-tagged protein. Hybridoma lines each from AlivaMab Kappa Mice and AlivaMab Lambda Mice are confirmed to bind to the antigen specifically by ELISA and are picked at random and single-cell cloned into 96-well plates. They are grown into colonies and the supernatant from these individual colonies is screened by ELISA to re-confirm monoclonal antibody binding to the antigen of interest. These supernatants are then screened by FACS to confirm binding to the native antigen expressed on cells.

AlivaMab Mouse technology and methods for producing antibodies using such technologies can be found in WO 2010/039900 and WO 2011/123708, which are incorporated herein in their entirety.

In certain methods for designing and/or producing the anti-CD3 antibodies of the invention, by elucidating the amino acid and/or nucleotide sequences of the CDR3 variable regions and/or one of CDR3 heavy chain (HC) and light chain (LC or λ) variable regions of the anti-CD3 antibodies produced, for example, using Alivama Mouse technology, a series of “clusters” or “motifs” are identified within the sequences. These clusters represent convergent somatic hypermutations (SHM) in the variable region sequences. Clustering may provide insight into the functionally related sequences and the diversity of the total population of antibodies and their variable region sequences. Sequences that are descendants from the same parent B cell or convergently evolved the sequences in the same cluster should be functionally more related than sequences belonging to other clusters. Convergent SHMs are likely functionally related mutations, e.g., they share a specific affinity for CD3. These SHM may inform the development of recombinant anti-CD3 antibodies with improved properties, such as specific binding for CD3 fragments.

Specific SHMs in the anti-CD3 antibody variable regions of the antibodies disclosed herein are found in Tables 1-4, with the amino acids representing an SHM indicated in bold.

Human antibodies may also be generated by isolating Fv clone variable domain sequences selected from human-derived phage display libraries. Such variable domain sequences may then be combined with a desired human constant domain.

Antibodies of the invention may be isolated by screening combinatorial libraries for antibodies with the desired activity or activities. For example, a variety of methods is known in the art for generating phage display libraries and screening such libraries for antibodies possessing the desired binding characteristics. Such methods are reviewed, e.g., in Hoogenboom et al. in *Methods in Molecular Biology* 178:1-37 (O'Brien et al., ed., Human Press, Totowa, N.J., 2001) and further described, e.g., in the McCafferty et al., *Nature* 348:552-554; Clackson et al., *Nature* 352: 624-628 (1991); Marks et al., *J. Mol. Biol.* 222: 581-597 (1992); Marks and Bradbury, in *Methods in Molecular Biology* 248:161-175 (Lo, ed., Human Press, Totowa, N.J., 2003); Sidhu et al., *J. Mol. Biol.* 338(2): 299-310 (2004); Lee et al., *J. Mol. Biol.* 340(5): 1073-1093 (2004); Fellouse, *Proc. Natl. Acad. Sci. USA* 101(34): 12467-12472 (2004); and Lee et al., *J. Immunol. Methods* 284(1-2): 119-132(2004).

In certain phage display methods, repertoires of VH and VL genes are separately cloned by polymerase chain reaction (PCR) and recombined randomly in phage libraries, which can then

be screened for antigen-binding phage as described in Winter et al., *Ann. Rev. Immunol.*, 12: 433-455 (1994). Phage typically display antibody fragments, either as single-chain Fv (scFv) fragments or as Fab fragments. Libraries from immunized sources provide high-affinity antibodies to the immunogen without the requirement of constructing hybridomas. Alternatively, the naive repertoire can be cloned (e.g., from human) to provide a single source of antibodies to a wide range of non-self and also self-antigens without any immunization as described by Griffiths et al., *EMBO J.*, 12: 725-734 (1993). Finally, naive libraries can also be made synthetically by cloning unarranged V-gene segments from stem cells, and using PCR primers containing random sequence to encode the highly variable CDR3 regions and to accomplish rearrangement in vitro, as described by Hoogenboom and Winter, *J. Mol. Biol.*, 227: 381-388 (1992). Patent publications describing human antibody phage libraries include, for example: U.S. Pat. No. 5,750,373, and US Patent Publication Nos. 2005/0079574, 2005/0119455, 2005/0266000, 2007/0117126, 2007/0160598, 2007/0237764, 2007/0292936, and 2009/0002360.

Alternatively, in certain aspects, the present invention includes humanized variants of the antibodies described herein or humanized antibodies comprising a one or more of SEQ ID NOS: 2-233, or a fragment thereof. Methods for humanizing antibodies are well known in the art.

In certain aspects, the anti-CD3 antibodies of the invention are, or form a part of, a multispecific antibody.

Multispecific antibodies are monoclonal antibodies that have binding specificities for at least two different sites. In some embodiments, bispecific antibodies may bind to two different epitopes of CD3 (e.g., CD3 ϵ or CD3 γ). In some embodiments, one of the binding specificities is for CD3 (e.g., CD3 ϵ or CD3 γ) and the other is for any other antigen (e.g., a second biological molecule, e.g., a cell surface antigen, e.g., a tumor antigen). Accordingly, a bispecific anti-CD3 antibody may have binding specificities for CD3 and a second biological molecule, such as a second biological molecule (e.g., a tumor antigen).

Techniques for making multispecific antibodies include, but are not limited to, recombinant co-expression of two immunoglobulin heavy chain-light chain pairs having different specificities (see Milstein and Cuello, *Nature* 305: 537 (1983)), WO 93/08829, and Traunecker et al., *EMBO J.* 10: 3655 (1991)), and “knob-in-hole” engineering (see, e.g., U.S. Pat. No. 5,731,168). “Knob-in-hole” engineering of multispecific antibodies may be utilized to generate a first arm containing a knob and a second arm containing the hole into which the knob of the first arm may bind. The

knob of the multispecific antibodies of the invention may be an anti-CD3 arm in one embodiment. Alternatively, the knob of the multispecific antibodies of the invention may be an anti-target/antigen arm in one embodiment. The hole of the multispecific antibodies of the invention may be an anti-CD3 arm in one embodiment. Alternatively, the hole of the multispecific antibodies
5 of the invention may be an anti-target/antigen arm in one embodiment.

There other ways of making multispecific antibodies. For example, multispecific antibodies may be engineered using immunoglobulin crossover (also known as Fab domain exchange or CrossMab format) technology (see e.g., WO2009/080253; Schaefer et al., Proc. Natl. Acad. Sci. USA, 108:11187-11192 (2011)). Multi-specific antibodies may also be made by
10 engineering electrostatic steering effects for making antibody Fc-heterodimeric molecules (WO 2009/089004A1); cross-linking two or more antibodies or fragments (see, e.g., U.S. Pat. No. 4,676,980, and Brennan et al., Science, 229: 81 (1985)); using leucine zippers to produce bi-specific antibodies (see, e.g., Kostelny et al., J. Immunol., 148(5):1547-1553 (1992)); using “diabody” technology for making bispecific antibody fragments (see, e.g., Hollinger et al., Proc.
15 Natl. Acad. Sci. USA, 90:6444-6448 (1993)); and using single-chain Fv (sFv) dimers (see, e.g. Gruber et al., J. Immunol., 152:5368 (1994)); and preparing trispecific antibodies as described, e.g., in Tutt et al. J. Immunol 147: 60 (1991).

In certain embodiments, amino acid sequence variants of the anti-CD3 antibodies of the invention (e.g., bispecific anti-CD3 antibodies of the invention that bind to CD3 and a second
20 biological molecule, e.g., a cell surface antigen, e.g., a tumor antigen, such as certain members and/or fragments of the epidermal growth factor receptor (HER/EGFR/ERBB) family, and/or different CD3 epitopes) are contemplated. For example, it may be desirable to improve the binding affinity and/or other biological properties of the antibody. Amino acid sequence variants of an antibody may be prepared by introducing appropriate modifications into the nucleotide sequence
25 encoding the antibody, or by peptide synthesis. Such modifications include, for example, deletions from, and/or insertions into and/or substitutions of residues within the amino acid sequences of the antibody. Any combination of deletion, insertion, and substitution can be made to arrive at the final construct, provided that the final construct possesses the desired characteristics, for example, antigen-binding.

In certain aspects, the anti-CD3 antibodies of the invention are multi-specific antibodies.
30 In certain aspects, the multi-specific antibodies are bispecific antibodies, trispecific antibodies,

and/or of greater multi-specificity that exhibit specificity to CD3 and another molecule and/or another epitope of CD3. For example, such antibodies can bind to both CD3 and to an antigen that is important for targeting the antibody to a particular cell type or tissue (for example, to an antigen associated with a cancer antigen of a tumor being treated). In some embodiments, multi-specific antibodies of the invention bind to molecules (receptors or ligands) involved in immunomodulatory pathways, such as CTLA4, TIM3, TIM4, OX40, CD40, GITR, 4-1-BB, CD27/CD70, ICOS, B7-H4, LIGHT, PD-1 or LAG3, which may provide control or modulation of the multi-specific antibodies' therapeutic effects. Furthermore, a multispecific antibody may bind to effector molecules such as cytokines (e.g., IL-7, IL-15, IL-12, IL-4 TGF-beta, IL-10, IL-17, IFN γ , Flt3, BLys) and/or chemokines (e.g., CCL21). Methods are known in the art for producing bispecific antibodies.

In certain embodiments, antibody variants having one or more amino acid substitutions are provided. Sites of interest for substitutional mutagenesis include the HVRs and FRs. Conservative substitutions are shown in Table A under the heading of "preferred substitutions." More substantial changes are provided in Table A under the heading of "exemplary substitutions," and as further described below in reference to amino acid side chain classes. Amino acid substitutions may be introduced into an antibody of interest and the products screened for a desired activity, for example, retained/improved antigen binding, decreased immunogenicity, or improved ADCC or CDC.

20

TABLE A

Exemplary and Preferred Amino Acid Substitutions		
Original Residue	Exemplary Substitutions	Preferred Substitutions
Ala (A)	Val; Leu; Ile	Val
Arg (R)	Lys; Gln; Asn	Lys
Asn (N)	Gln; His; Asp; Lys; Arg	Gln
Asp (D)	Glu; Asn	Glu
Cys (C)	Ser; Ala	Ser
Gln (Q)	Asn; Glu	Asn

Glu (E)	Asp; Gln	Asp
Gly (G)	Ala	Ala
His (H)	Asn; Gln; Lys; Arg	Arg
Ile (I)	Leu; Val; Met; Ala; Phe ; Norleucine	Leu
Leu (L)	Norleucine; Ile; Val; Met; Ala, Phe	Ile
Lys (K)	Arg; Gln; Asn	Arg
Met (M)	Leu; Phe; Ile	Leu
Phe (F)	Trp; Leu; Val; Ile; Ala; Tyr	Tyr
Pro (P)	Ala	Ala
Ser (S)	Thr	Thr
Thr (T)	Val; Ser	Ser
Trp (W)	Tyr; Phe	Tyr
Tyr (Y)	Trp; Phe; Thr; Ser	Phe
Val (V)	Ile; Leu; Met; Phe; Ala; Norleucine	Leu

Amino acids may be grouped according to common side-chain properties:

- (1) hydrophobic: Met, Ala, Val, Leu, Ile;
- (2) neutral hydrophilic: Cys, Ser, Thr, Asn, Gln;
- 5 (3) acidic: Asp, Glu;
- (4) basic: His, Lys, Arg;
- (5) residues that influence chain orientation: Gly, Pro;
- (6) aromatic: Trp, Tyr, Phe.

10 Non-conservative substitutions will entail exchanging a member of one of these classes for another class.

One type of substitutional variant involves substituting one or more hypervariable region residues of a parent antibody (e.g. a humanized or human antibody). Generally, the resulting variant(s) selected for further study will have modifications (e.g., improvements) in certain biological properties (e.g., increased affinity, reduced immunogenicity) relative to the parent antibody and/or will have substantially retained certain biological properties of the parent antibody. An exemplary substitutional variant is an affinity matured antibody, which may be
15 conveniently generated, e.g., using phage display-based affinity maturation techniques. Briefly,

one or more HVR residues are mutated and the variant antibodies displayed on phage and screened for a particular biological activity (e.g. binding affinity).

Alterations (e.g., substitutions) may be made in HVRs, e.g., to improve antibody affinity. Such alterations may be made in HVR “hotspots,” i.e., residues encoded by codons that undergo mutation at high frequency during the somatic maturation process (see, e.g., Chowdhury, *Methods Mol. Biol.* 207:179-196 (2008)), and/or residues that contact antigen, with the resulting variant VH or VL being tested for binding affinity. Affinity maturation by constructing and reselecting from secondary libraries has been described, e.g., in Hoogenboom et al. in *Methods in Molecular Biology* 178:1-37 (O'Brien et al., ed., Human Press, Totowa, N.J., (2001).) In some embodiments of affinity maturation, diversity is introduced into the variable genes chosen for maturation by any of a variety of methods (e.g., error-prone PCR, chain shuffling, or oligonucleotide-directed mutagenesis). A secondary library is then created. The library is then screened to identify any antibody variants with the desired affinity. Another method to introduce diversity involves HVR-directed approaches, in which several HVR residues (e.g., 4-6 residues at a time) are randomized. HVR residues involved in antigen binding may be specifically identified, e.g., using alanine scanning mutagenesis or modeling. CDR-H3 and CDR-L3 in particular are often targeted.

In certain embodiments, substitutions, insertions, or deletions may occur within one or more HVRs so long as such alterations do not substantially reduce the ability of the antibody to bind antigen. For example, conservative alterations (e.g., conservative substitutions as provided herein) that do not substantially reduce binding affinity may be made in HVRs. Such alterations may, for example, be outside of antigen contacting residues in the HVRs. In certain embodiments of the variant VH and VL sequences provided above, each HVR either is unaltered, or contains no more than one, two, or three amino acid substitutions.

A useful method for identification of residues or regions of an antibody that may be targeted for mutagenesis is called “alanine scanning mutagenesis” as described by Cunningham and Wells (1989) *Science*, 244:1081-1085. In this method, a residue or group of target residues (e.g., charged residues such as arg, asp, his, lys, and glu) are identified and replaced by a neutral or negatively charged amino acid (e.g., alanine or polyalanine) to determine whether the interaction of the antibody with antigen is affected. Further substitutions may be introduced at the amino acid locations demonstrating functional sensitivity to the initial substitutions. Alternatively, or additionally, a crystal structure of an antigen-antibody complex to identify contact points

between the antibody and antigen may be used. Such contact residues and neighboring residues may be targeted or eliminated as candidates for substitution. Variants may be screened to determine whether they contain the desired properties.

Amino acid sequence insertions include amino- and/or carboxyl-terminal fusions ranging in length from one residue to polypeptides containing a hundred or more residues, as well as intra-sequence insertions of single or multiple amino acid residues. Examples of terminal insertions include an antibody with an N-terminal methionyl residue. Other insertional variants of the antibody molecule include the fusion to the N- or C-terminus of the antibody to an enzyme or a polypeptide which increases the serum half-life of the antibody.

In certain embodiments, anti-CD3 antibodies of the invention can be altered to increase or decrease the extent to which the antibody is glycosylated. Addition or deletion of glycosylation sites to anti-CD3 antibody of the invention may be conveniently accomplished by altering the amino acid sequence such that one or more glycosylation sites is created or removed.

Where the antibody comprises an Fc region, the carbohydrate attached thereto may be altered. Native antibodies produced by mammalian cells typically comprise a branched, biantennary oligosaccharide that is generally attached by an N-linkage to Asn297 of the CH2 domain of the Fc region. See, e.g., Wright et al. TIBTECH 15:26-32 (1997). The oligosaccharide may include various carbohydrates, e.g., mannose, N-acetyl glucosamine (GlcNAc), galactose, and sialic acid, as well as a fucose attached to a GlcNAc in the "stem" of the biantennary oligosaccharide structure. In some embodiments, modifications of the oligosaccharide in an antibody of the invention may be made in order to create antibody variants with certain improved properties.

In one embodiment, anti-CD3 antibody variants are provided having a carbohydrate structure that lacks fucose attached (directly or indirectly) to an Fc region. For example, the amount of fucose in such antibody may be from 1% to 80%, from 1% to 65%, from 5% to 65%, or from 20% to 40%. The amount of fucose is determined by calculating the average amount of fucose within the sugar chain at Asn297, relative to the sum of all glycostructures attached to Asn 297 (e.g. complex, hybrid and high mannose structures) as measured by MALDI-TOF mass spectrometry, as described in WO 2008/077546, for example. Asn297 refers to the asparagine residue located at about position 297 in the Fc region (EU numbering of Fc region residues); however, Asn297 may also be located about 3 amino acids upstream or downstream of position

297, i.e., between positions 294 and 300, due to minor sequence variations in antibodies. Such fucosylation variants may have improved ADCC function. See, e.g., US Patent Publication Nos. US 2003/0157108 (Presta, L.); US 2004/0093621 (Kyowa Hakko Kogyo Co., Ltd). Examples of publications related to “defucosylated” or “fucose-deficient” antibody variants include: US 5 2003/0157108; WO 2000/61739; WO 2001/29246; US 2003/0115614; US 2002/0164328; US 2004/0093621; US 2004/0132140; US 2004/0110704; US 2004/0110282; US 2004/0109865; WO 2003/085119; WO 2003/084570; WO 2005/035586; WO 2005/035778; WO2005/053742; WO2002/031140; Okazaki et al. *J. Mol. Biol.* 336:1239-1249 (2004); Yamane-Ohnuki et al. *Biotech. Bioeng.* 87: 614 (2004). Examples of cell lines capable of producing defucosylated 10 antibodies include Lec13 CHO cells deficient in protein fucosylation (Ripka et al. *Arch. Biochem. Biophys.* 249:533-545 (1986); US Pat Appl No US 2003/0157108 A1, Presta, L; and WO 2004/056312 A1, Adams et al., especially at Example 11), and knockout cell lines, such as alpha-1,6-fucosyltransferase gene, FUT8, knockout CHO cells (see, e.g., Yamane-Ohnuki et al. *Biotech. Bioeng.* 87: 614 (2004); Kanda, Y. et al., *Biotechnol. Bioeng.*, 94(4):680-688 (2006); and 15 WO2003/085107).

Anti-CD3 antibodies variants are further provided with bisected oligosaccharides, for example, in which a biantennary oligosaccharide attached to the Fc region of the antibody is bisected by GlcNAc. Such antibody variants may have reduced fucosylation and/or improved ADCC function. Examples of such antibody variants are described, e.g., in WO 2003/011878 20 (Jean-Mairet et al.); U.S. Pat. No. 6,602,684 (Umana et al.); and US 2005/0123546 (Umana et al.). Antibody variants with at least one galactose residue in the oligosaccharide attached to the Fc region are also provided. Such antibody variants may have improved CDC function. Such antibody variants are described, e.g., in WO 1997/30087 (Patel et al.); WO 1998/58964 (Raju, S.); and WO 1999/22764 (Raju, S.).

25 In certain embodiments, one or more amino acid modifications may be introduced into the Fc region of an anti-CD3 antibody of the invention thereby generating an Fc region variant (see e.g., US 2012/0251531). The Fc region variant may comprise a human Fc region sequence (e.g., a human IgG1, IgG2, IgG3 or IgG4 Fc region) comprising an amino acid modification (e.g., a substitution) at one or more amino acid positions. In certain embodiments, the invention 30 contemplates an anti-CD3 antibody variant that possesses some but not all effector functions, which make it a desirable candidate for applications in which the half life of the antibody in vivo

is important yet certain effector functions (such as complement and ADCC) are unnecessary or deleterious. In vitro and/or in vivo cytotoxicity assays can be conducted to confirm the reduction/depletion of CDC and/or ADCC activities. For example, Fc receptor (FcR) binding assays can be conducted to ensure that the antibody lacks Fc γ R binding (hence likely lacking ADCC activity), but retains FcRn binding ability. The primary cells for mediating ADCC, NK cells, express Fc γ RIII only, whereas monocytes express Fc γ RI, Fc γ RII and Fc γ RIII. FcR expression on hematopoietic cells is summarized in Table 3 on page 464 of Ravetch and Kinetic, *Annu. Rev. Immunol.* 9:457-492 (1991). Non-limiting examples of in vitro assays to assess ADCC activity of a molecule of interest is described in U.S. Pat. No. 5,500,362 (see, e.g. Hellstrom, I. et al. *Proc. Nat'l Acad. Sci. USA* 83:7059-7063 (1986)) and Hellstrom, I et al., *Proc. Nat'l Acad. Sci. USA* 82:1499-1502 (1985); U.S. Pat. No. 5,821,337 (see Bruggemann, M. et al., *J. Exp. Med.* 166:1351-1361 (1987)). Alternatively, non-radioactive assays methods may be employed (see, for example, ACTI™ non-radioactive cytotoxicity assay for flow cytometry (CellTechnology, Inc. Mountain View, Calif.; and CytoTox 96 non-radioactive cytotoxicity assay (Promega, Madison, Wis.). Useful effector cells for such assays include peripheral blood mononuclear cells (PBMC) and Natural Killer (NK) cells.

Alternatively, or additionally, ADCC activity of the molecule of interest may be assessed in vivo, e.g., in an animal model such as that disclosed in Clynes et al. *Proc. Nat'l Acad. Sci. USA* 95:652-656 (1998). C1q binding assays may also be carried out to confirm that the antibody is unable to bind C1q and hence lacks CDC activity. See, e.g., C1q and C3c binding ELISA in WO 2006/029879 and WO 2005/100402. To assess complement activation, a CDC assay may be performed (see, for example, Gazzano-Santoro et al. *J. Immunol. Methods* 202:163 (1996); Cragg, M. S. et al. *Blood*. 101:1045-1052 (2003); and Cragg, M. S. and M. J. Glennie *Blood*. 103:2738-2743 (2004)). FcRn binding and in vivo clearance/half life determinations can also be performed using methods known in the art (see, e.g., Petkova, S. B. et al. *Int'l. Immunol.* 18(12):1759-1769 (2006)).

Antibodies with reduced effector function include those with substitution of one or more of Fc region residues 238, 265, 269, 270, 297, 327 and 329 (U.S. Pat. Nos. 6,737,056 and 8,219,149). Such Fc mutants include Fc mutants with substitutions at two or more of amino acid positions 265, 269, 270, 297 and 327, including the so-called "DANA" Fc mutant with substitution of residues 265 and 297 to alanine (U.S. Pat. Nos. 7,332,581 and 8,219,149).

In certain embodiments, the proline at position 329 of a wild-type human Fc region in the antibody is substituted with glycine or arginine or an amino acid residue large enough to destroy the proline sandwich within the Fc/Fc γ receptor interface that is formed between the proline 329 of the Fc and tryptophan residues Trp 87 and Trp 110 of Fc γ RIII (Sondermann et al.: Nature 406, 267-273 (20 Jul. 2000)). In certain embodiments, the antibody comprises at least one further amino acid substitution. In one embodiment, the further amino acid substitution is S228P, E233P, L234A, L235A, L235E, N297A, N297D, or P331S, and still in another embodiment the at least one further amino acid substitution is L234A and L235A of the human IgG1 Fc region or S228P and L235E of the human IgG4 Fc region (see e.g., US 2012/0251531), and still in another embodiment the at least one further amino acid substitution is L234A and L235A and P329G of the human IgG1 Fc region.

Certain antibody variants with improved or diminished binding to FcRs are described. (See, e.g., U.S. Pat. No. 6,737,056; WO 2004/056312, and Shields et al., J. Biol. Chem. 9(2): 6591-6604 (2001).)

In certain embodiments, an antibody variant comprises an Fc region with one or more amino acid substitutions which improve ADCC, e.g., substitutions at positions 298, 333, and/or 334 of the Fc region (EU numbering of residues).

In some embodiments, alterations are made in the Fc region that result in altered (i.e., either improved or diminished) C1q binding and/or Complement Dependent Cytotoxicity (CDC), e.g., as described in U.S. Pat. No. 6,194,551, WO 99/51642, and Idusogie et al. J. Immunol. 164: 4178-4184 (2000).

Antibodies with increased half-lives and improved binding to the neonatal Fc receptor (FcRn), which is responsible for the transfer of maternal IgGs to the fetus (Guyer et al., J. Immunol. 117:587 (1976) and Kim et al., J. Immunol. 24:249 (1994)), are described in US2005/0014934A1 (Hinton et al.). Those antibodies comprise an Fc region with one or more substitutions therein which improve binding of the Fc region to FcRn. Such Fc variants include those with substitutions at one or more of Fc region residues: 238, 256, 265, 272, 286, 303, 305, 307, 311, 312, 317, 340, 356, 360, 362, 376, 378, 380, 382, 413, 424 or 434, e.g., substitution of Fc region residue 434 (U.S. Pat. No. 7,371,826). See also Duncan & Winter, Nature 322:738-40 (1988); U.S. Pat. Nos. 5,648,260; 5,624,821; and WO 94/29351 concerning other examples of Fc region variants.

In some aspects the bispecific antibody comprises an Fc region comprising an N297G mutation. In some embodiments, the bispecific antibody comprising the N297G mutation comprises one or more heavy chain constant domains, wherein the one or more heavy chain constant domains are selected from a first CH1 domain, a first CH2 domain, a first CH3 domain, a second CH1 domain, second CH2 domain, and a second CH3 domain.

In certain embodiments, it may be desirable to create cysteine engineered antibodies in which one or more residues of an antibody are substituted with cysteine residues. In some embodiments, the substituted residues occur at accessible sites of the antibody. By substituting those residues with cysteine, reactive thiol groups are thereby positioned at accessible sites of the antibody and may be used to conjugate the antibody to other moieties, such as drug moieties or linker-drug moieties, to create an immunoconjugate, as described further herein. In certain embodiments, any one or more of the following residues may be substituted with cysteine: V205 (Kabat numbering) of the light chain; A118 (EU numbering) of the heavy chain; and S400 (EU numbering) of the heavy chain Fc region. Cysteine engineered antibodies may be generated as described, for example, in WO 2016/040856, which is incorporated by reference in its entirety herein, including any drawings.

In certain embodiments, the bispecific antibody provided herein may be further modified to contain additional non-proteinaceous moieties that are known in the art and readily available. The moieties suitable for derivatization of the antibody include but are not limited to water soluble polymers. Non-limiting examples of water soluble polymers include, but are not limited to, polyethylene glycol (PEG), copolymers of ethylene glycol/propylene glycol, carboxymethylcellulose, dextran, polyvinyl alcohol, polyvinyl pyrrolidone, poly-1,3-dioxolane, poly-1,3,6-trioxane, ethylene/maleic anhydride copolymer, polyaminoacids (either homopolymers or random copolymers), and dextran or poly(n-vinyl pyrrolidone)polyethylene glycol, propylene glycol homopolymers, polypropylene oxide/ethylene oxide co-polymers, polyoxyethylated polyols (e.g., glycerol), polyvinyl alcohol, and mixtures thereof. Polyethylene glycol propionaldehyde may have advantages in manufacturing due to its stability in water. The polymer may be of any molecular weight, and may be branched or unbranched. The number of polymers attached to the antibody may vary, and if more than one polymer are attached, they can be the same or different molecules. In general, the number and/or type of polymers used for derivatization can be determined based on considerations including, but not limited to, the

particular properties or functions of the antibody to be improved, whether the antibody derivative will be used in a therapy under defined conditions, etc.

In another embodiment, conjugates of an antibody and nonproteinaceous moiety that may be selectively heated by exposure to radiation are provided. In one embodiment, the
5 nonproteinaceous moiety is a carbon nanotube (Kam et al., Proc. Natl. Acad. Sci. USA 102: 11600-11605 (2005)). The radiation may be of any wavelength, and includes, but is not limited to, wavelengths that do not harm ordinary cells, but which heat the nonproteinaceous moiety to a temperature at which cells proximal to the antibody-nonproteinaceous moiety are killed.

The bispecific antibodies of the invention may be produced using recombinant methods
10 and compositions, for example, as described in U.S. Pat. No. 4,816,567. In one embodiment, an isolated nucleic acid encoding an anti-CD3 antibody described herein is provided. Such nucleic acid may encode an amino acid sequence comprising the VL and/or an amino acid sequence comprising the VH of the antibody (e.g., the light and/or heavy chains of the antibody). In a further embodiment, one or more vectors (e.g., expression vectors) comprising such nucleic acid are
15 provided. In a further embodiment, a host cell comprising such nucleic acid is provided. In one such embodiment, a host cell comprises (e.g., has been transformed with): (1) a vector comprising a nucleic acid that encodes an amino acid sequence comprising the VL of the antibody and an amino acid sequence comprising the VH of the antibody, or (2) a first vector comprising a nucleic acid that encodes an amino acid sequence comprising the VL of the antibody and a second vector
20 comprising a nucleic acid that encodes an amino acid sequence comprising the VH of the antibody. In one embodiment, the host cell is eukaryotic, e.g. a Chinese Hamster Ovary (CHO) cell or lymphoid cell (e.g., Y0, NS0, Sp20 cell). In one embodiment, a method of making a bispecific antibody is provided, wherein the method comprises culturing a host cell comprising a nucleic acid encoding the antibody under conditions suitable for expression of the antibody, and optionally
25 recovering the antibody from the host cell (or host cell culture medium).

For recombinant production of an antibody, a nucleic acid encoding an antibody is isolated and inserted into one or more vectors for further cloning and/or expression in a host cell. Such nucleic acid may be readily isolated and sequenced using conventional procedures (e.g., by using
30 oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of the antibody).

Suitable host cells for cloning or expression of antibody-encoding vectors include prokaryotic or eukaryotic cells described herein. For example, antibodies may be produced in bacteria, in particular when glycosylation and Fc effector function are not needed. For expression of antibody fragments and polypeptides in bacteria, see, e.g., U.S. Pat. Nos. 5,648,237, 5,789,199, and 5,840,523. (See also Charlton, *Methods in Molecular Biology*, Vol. 248 (B. K. C. Lo, ed., Humana Press, Totowa, N.J., 2003), pp. 245-254, describing expression of antibody fragments in *E. coli*.) After expression, the antibody may be isolated from the bacterial cell paste in a soluble fraction and can be further purified.

In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable cloning or expression hosts for antibody-encoding vectors, including fungi and yeast strains whose glycosylation pathways have been “humanized,” resulting in the production of an antibody with a partially or fully human glycosylation pattern. See Gerngross, *Nat. Biotech.* 22:1409-1414 (2004), and Li et al., *Nat. Biotech.* 24:210-215 (2006).

Suitable host cells for the expression of glycosylated antibody are also derived from multicellular organisms (invertebrates and vertebrates). Examples of invertebrate cells include plant and insect cells. Numerous baculoviral strains have been identified which may be used in conjunction with insect cells, particularly for transfection of *Spodoptera frugiperda* cells.

Plant cell cultures can also be utilized as hosts. See, e.g., U.S. Pat. Nos. 5,959,177, 6,040,498, 6,420,548, 7,125,978, and 6,417,429 (describing PLANTIBODIES™ technology for producing antibodies in transgenic plants).

Vertebrate cells may also be used as hosts. For example, mammalian cell lines that are adapted to grow in suspension may be useful. Other examples of useful mammalian host cell lines are monkey kidney CV1 line transformed by SV40 (COS-7); human embryonic kidney line (293 or 293 cells as described, e.g., in Graham et al., *J. Gen Virol.* 36:59 (1977)); baby hamster kidney cells (BHK); mouse sertoli cells (TM4 cells as described, e.g., in Mather, *Biol. Reprod.* 23:243-251 (1980)); monkey kidney cells (CV1); African green monkey kidney cells (VERO-76); human cervical carcinoma cells (HELA); canine kidney cells (MDCK; buffalo rat liver cells (BRL 3A); human lung cells (W138); human liver cells (Hep G2); mouse mammary tumor (MMT 060562); TRI cells, as described, e.g., in Mather et al., *Annals N.Y. Acad. Sci.* 383:44-68 (1982); MRC 5 cells; and FS4 cells. Other useful mammalian host cell lines include Chinese hamster ovary (CHO) cells, including DHFR-CHO cells (Urlaub et al., *Proc. Natl. Acad. Sci. USA* 77:4216 (1980)), and

myeloma cell lines such as Y0, NS0 and Sp2/0. For a review of certain mammalian host cell lines suitable for antibody production, see, e.g., Yazaki and Wu, *Methods in Molecular Biology*, Vol. 248 (B. K. C. Lo, ed., Humana Press, Totowa, N.J.), pp. 255-268 (2003).

5 The antibodies of the invention may be identified, screened for, or characterized for their physical/chemical properties and/or biological activities by various assays known in the art.

In one aspect, the antibody of the invention is tested for its antigen binding activity, for example, by known methods such as ELISA, Western blot, etc. In another aspect, competition assays may be used to identify an antibody that competes with an anti-CD3 antibody of the invention for binding to CD3. In an exemplary competition assay, immobilized CD3 is incubated
10 in a solution comprising a first labeled antibody that binds to CD3 and a second unlabeled antibody that is being tested for its ability to compete with the first antibody for binding to CD3. The second antibody may be present in a hybridoma supernatant. As a control, immobilized CD3 is incubated in a solution comprising the first labeled antibody but not the second unlabeled antibody. After incubation under conditions permissive for binding of the first antibody to CD3, excess unbound
15 antibody is removed, and the amount of label associated with immobilized CD3 is measured. If the amount of label associated with immobilized CD3 is substantially reduced in the test sample relative to the control sample, then that indicates that the second antibody is competing with the first antibody for binding to CD3. See, e.g., Harlow and Lane (1988) *Antibodies: A Laboratory Manual*. Ch. 14 (Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.).

20 In one aspect, assays are provided for identifying antibodies having biological activity. Biological activity may include, for example, binding to CD3 (e.g., CD3 on the surface of a T cell), or a peptide fragment thereof, either *in vivo*, *in vitro*, or *ex vivo*. In the case of a bispecific antibody of the invention, biological activity may also include, for example, effector cell activation (e.g., T cell (e.g., CD8+ and/or CD4+ T cell) activation), effector cell population expansion (i.e.,
25 an increase in T cell count), target cell population reduction (i.e., a decrease in the population of cells expressing the second biological molecule on their cell surfaces), and/or target cell killing. In some embodiments, the activity comprises ability to support B cell killing and/or the activation of the cytotoxic T cells.

In certain embodiments, any of the antibodies of the invention may be used to detect the
30 presence of CD3 in a biological sample. The term “detecting” as used herein encompasses quantitative or qualitative detection. In certain embodiments, a biological sample comprises a cell

or tissue. In certain embodiments, the method comprises contacting the biological sample with an anti-CD3 antibody as described herein under conditions permissive for binding of the bispecific antibody to CD3 and another antigen, and detecting whether a complex is formed between the bispecific antibody and CD3. Such method may be an in vitro or in vivo method.

5 In certain embodiments, labeled antibodies are provided. Labels include, but are not limited to, labels or moieties that are detected directly (such as fluorescent, chromophoric, electron-dense, chemiluminescent, and radioactive labels), as well as moieties, such as enzymes or ligands, that are detected indirectly, e.g., through an enzymatic reaction or molecular interaction. Exemplary labels include, but are not limited to, the radioisotopes ^{32}P , ^{14}C , ^{125}I , ^3H , and ^{131}I , fluorophores
10 such as rare earth chelates or fluorescein and its derivatives, rhodamine and its derivatives, dansyl, umbelliferone, luciferases, e.g., firefly luciferase, and bacterial luciferase (see for example, U.S. Pat. No. 4,737,456, which is incorporated by reference in its entirety herein, including any drawings), luciferin, 2,3-dihydrophthalazinediones, horseradish peroxidase (HRP), alkaline phosphatase, β -galactosidase, glucoamylase, lysozyme, saccharide oxidases, e.g., glucose oxidase,
15 galactose oxidase, and glucose-6-phosphate dehydrogenase, heterocyclic oxidases such as uricase and xanthine oxidase, coupled with an enzyme that employs hydrogen peroxide to oxidize a dye precursor such as HRP, lactoperoxidase, or microperoxidase, biotin/avidin, spin labels, bacteriophage labels, stable free radicals, and the like.

20 Production of Anti-CD3 Antibodies and Antibody Compositions

An additional aspect of the invention relates to methods for producing an anti-CD3 antibodies and compositions of the invention. One embodiment of this aspect of the invention relates to a method for producing an anti-CD3 antibody as defined herein, comprising providing a host cell capable of expressing an anti-CD3 antibody, cultivating said host cell under conditions
25 suitable for expression of the antibody, and isolating the resulting antibody.

An antibody or antibody composition of the present invention may be produced by methods generally known in the art for production of recombinant monoclonal or polyclonal antibodies. Thus, in the case of production of a single antibody of the invention, any method known in the art for production of recombinant monoclonal antibodies may be used. For production of an antibody
30 composition comprising two or more anti-CD3 antibodies of the invention, the individual antibodies may be produced separately, i.e., each antibody being produced in a separate bioreactor,

or the individual antibodies may be produced together in single bioreactor. When the number of different antibodies in a composition is more than e.g., two or three, it will generally be preferably for reasons of cost efficiency to produce the antibodies together in a single bioreactor. On the other hand, when the composition only contains a small number of different antibodies, e.g., two, three
5 or possibly four different antibodies, a decision to produce them separately in different bioreactors or together in a single bioreactor will have to be made based on the individual circumstances. If the antibody composition is produced in more than one bioreactor, the purified anti-CD3 antibody composition can be obtained by pooling the antibodies obtained from individually purified supernatants from each bioreactor. Various approaches are known in the art for production of a
10 polyclonal antibody composition in multiple bioreactors, where the cell lines or antibody preparations are combined at a later point upstream or prior to or during downstream processing.

In the case of production of two or more individual antibodies in a single bioreactor, this may be performed, for example, based on site-specific integration of the antibody coding sequence into the genome of the individual host cells, ensuring that the V_H and V_L protein chains are
15 maintained in their original pairing during production. Furthermore, the site-specific integration minimizes position effects, and therefore the growth and expression properties of the individual cells in the polyclonal cell line are expected to be very similar. Generally, the method involves the following: i) a host cell with one or more recombinase recognition sites; ii) an expression vector with at least one recombinase recognition site compatible with that of the host cell; iii) generation
20 of a collection of expression vectors by transferring the selected V_H and V_L coding pairs from the screening vector to an expression vector such that a full-length antibody or antibody fragment can be expressed from the vector (such a transfer may not be necessary if the screening vector is identical to the expression vector); iv) transfection of the host cell with the collection of expression vectors and a vector coding for a recombinase capable of combining the recombinase recognition
25 sites in the genome of the host cell with that in the vector; v) obtaining/generating a polyclonal cell line from the transfected host cell and vi) expressing and collecting the antibody composition from the polyclonal cell line.

An alternative approach is to produce two or more different antibodies in a single bioreactor. This method involves generation of a polyclonal cell line capable of expressing a
30 polyclonal antibody or other polyclonal protein comprising two or more distinct members by a) providing a set of expression vectors, wherein each of said vectors comprises at least one copy of

a distinct nucleic acid encoding a distinct member of the polyclonal protein, separately transfecting host cells with each of the expression vectors under conditions avoiding site-specific integration of the expression vectors into the genome of the cells, thereby obtaining two or more compositions of cells, each composition expressing one distinct member of the polyclonal protein, and c) mixing
5 the at least two compositions of cells to obtain a polyclonal cell line.

The antibodies of the invention may be produced in various types of cells, including mammalian cells as well as non-mammalian eukaryotic or prokaryotic cells, such as plant cells, insect cells, yeast cells, fungi, *E. coli* etc. However, the antibodies are preferably produced in mammalian cells, for example CHO cells, COS cells, BHK cells, myeloma cells (e.g., Sp2/0 or
10 NS0 cells), fibroblasts such as NIH 3T3, or immortalized human cells such as HeLa cells or HEK 293 cells.

Methods for transfecting a nucleic acid sequence into a host cell are well-known in the art (see, e.g., Sambrook et al., *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratory Press, 3rd Edition, 2001). For site-specific integration a suitable host cell will comprise
15 one or more recombinase recognition sites in its genome. In this case, a suitable expression vector comprises a recombination recognition site matching the recombinase recognition site(s) of the host cell.

One embodiment of the present invention is thus a polyclonal cell line capable of expressing two or more anti-CD3 antibodies of the present invention. A further embodiment is a
20 polyclonal cell line wherein each individual cell is capable of expressing a single V_H and V_L pair, and the polyclonal cell line as a whole is capable of expressing a collection of V_H and V_L pairs, where each V_H and V_L pair encodes an anti-CD3 antibody.

Therapeutic Compositions

Another aspect of the invention is a pharmaceutical composition comprising as an active
25 ingredient at least one anti-CD3 antibody of the invention, or an anti-CD3 Fab or another anti-CD3 antibody fragment composition. Such compositions are intended for amelioration, prevention and/or treatment of cancer. The pharmaceutical composition may be administered to a human or to a domestic animal.

In addition to at least one antibody of the invention or fragment thereof, the pharmaceutical
30 composition will further comprise at least one pharmaceutically acceptable diluent, carrier or

excipient. These may for example include preservatives, stabilizers, surfactants/wetting agents, emulsifying agents, solubilizers, salts for regulating the osmotic pressure and/or buffers. Solutions or suspensions may further comprise viscosity-increasing substances, such as sodium carboxymethylcellulose, carboxymethylcellulose, dextran, polyvinylpyrrolidone or gelatin. A suitable pH value for the pharmaceutical composition will generally be in the range of about 5.5 to 8.5, such as about 6 to 8, e.g., about 7, maintained where appropriate by use of a buffer.

Conventional pharmaceutical practice may be employed to provide suitable formulations or compositions to administer to e.g., cancer patients. The administration will typically be therapeutic, meaning that it is administered after a cancer condition has been diagnosed. Any appropriate route of administration may be employed, for example parenteral, intravenous, intra-arterial, subcutaneous, intramuscular, intraperitoneal, intranasal, aerosol, suppository or oral administration. Pharmaceutical compositions of the invention will typically be administered in the form of liquid solutions or suspensions, more typically aqueous solutions or suspensions, in particular isotonic aqueous solutions or suspensions.

As an alternative to a liquid formulation, the compositions of the invention may be prepared in lyophilized form comprising the at least one antibody alone or together with a carrier, for example mannitol, in which case the composition is reconstituted with a liquid such as sterile water prior to use.

The pharmaceutical compositions comprise from approximately 1% to approximately 95%, preferably from approximately 20% to approximately 90%, active ingredient. Pharmaceutical compositions according to the invention may e.g., be produced in unit dose form, such as in the form of ampoules, vials, suppositories, tablets or capsules. The formulations can be administered to human individuals in therapeutically or prophylactically effective amounts (e.g., amounts which prevent, eliminate, or reduce a pathological condition) to provide therapy for a cancerous disease or other condition. The preferred dosage of therapeutic agent to be administered is likely to depend on such variables as the severity of the cancer, the overall health status of the particular patient, the formulation of the compound excipients, and its route of administration.

Therapeutic Uses of Antibodies and Compositions According to the Invention

The anti-CD3 antibodies and pharmaceutical compositions according to the present invention may be used for the treatment or amelioration of a disease, in a mammal, in particular treatment of cancer in humans.

Some embodiments provide a method of treating or delaying the progression of a cell proliferative disorder or an autoimmune disorder in a subject in need thereof, the method comprising administering to the subject an effective amount any one of the antibodies described herein (in a monospecific, bi-specific, or multi-specific format). In another aspect, the invention features a method of enhancing or decreasing immune function in a subject having a cell proliferative disorder or an autoimmune disorder, the method comprising administering to the subject any one of the antibodies described herein (in a monospecific, bi-specific, or multi-specific format).

In any of the uses or methods set forth herein, the cell proliferative disorder can be cancer. In some embodiments, the cancer is selected from the group consisting of breast cancer, colorectal cancer, non-small cell lung cancer, non-Hodgkin's lymphoma (NHL), B cell lymphoma, B cell leukemia, multiple myeloma, renal cancer, prostate cancer, liver cancer, head and neck cancer, melanoma, ovarian cancer, mesothelioma, glioblastoma, germinal-center B-cell-like (GCB) DLBCL, activated B-cell-like (ABC) DLBCL, follicular lymphoma (FL), mantle cell lymphoma (MCL), acute myeloid leukemia (AML), chronic lymphoid leukemia (CLL), marginal zone lymphoma (MZL), small lymphocytic leukemia (SLL), lymphoplasmacytic lymphoma (LL), Waldenstrom macroglobulinemia (WM), central nervous system lymphoma (CNSL), Burkitt's lymphoma (BL), B-cell prolymphocytic leukemia, Splenic marginal zone lymphoma, Hairy cell leukemia, Splenic lymphoma/leukemia, unclassifiable, Splenic diffuse red pulp small B-cell lymphoma, Hairy cell leukemia variant, Waldenstrom macroglobulinemia, Heavy chain diseases, a Heavy chain disease, γ Heavy chain disease, Heavy chain disease, Plasma cell myeloma, Solitary plasmacytoma of bone, Extrasosseous plasmacytoma, Extranodal marginal zone lymphoma of mucosa-associated lymphoid tissue (MALT lymphoma), Nodal marginal zone lymphoma, Pediatric nodal marginal zone lymphoma, Pediatric follicular lymphoma, Primary cutaneous follicle centre lymphoma, T-cell/histiocyte rich large B-cell lymphoma, Primary DLBCL of the CNS, Primary cutaneous DLBCL, leg type, EBV-positive DLBCL of the elderly, DLBCL associated with chronic inflammation, Lymphomatoid granulomatosis, Primary mediastinal (thymic) large B-cell lymphoma, Intravascular large B-cell lymphoma, ALK-positive large B-cell

lymphoma, Plasmablastic lymphoma, Large B-cell lymphoma arising in HHV8-associated multicentric Castleman disease, Primary effusion lymphoma: B-cell lymphoma, unclassifiable, with features intermediate between diffuse large B-cell lymphoma and Burkitt lymphoma, and B-cell lymphoma, unclassifiable, with features intermediate between diffuse large B-cell lymphoma and classical Hodgkin lymphoma. In some embodiments, the preferred cancer is germinal-center B-cell-like (GCB) DLBCL, activated B-cell-like (ABC) DLBCL, follicular lymphoma (FL), mantle cell lymphoma (MCL), acute myeloid leukemia (AML), chronic lymphoid leukemia (CLL), marginal zone lymphoma (MZL), small lymphocytic leukemia (SLL), lymphoplasmacytic lymphoma (LL), Waldenstrom macroglobulinemia (WM), central nervous system lymphoma (CNSL), or Burkitt's lymphoma (BL).

In some embodiments, the autoimmune disorder is selected from the group consisting of rheumatoid arthritis, juvenile rheumatoid arthritis, systemic lupus erythematosus (SLE), Wegener's disease, inflammatory bowel disease, idiopathic thrombocytopenic purpura (ITP), thrombotic thrombocytopenic purpura (TTP), autoimmune thrombocytopenia, multiple sclerosis, psoriasis, IgA nephropathy, IgM polyneuropathies, myasthenia gravis, vasculitis, diabetes mellitus, Reynaud's syndrome, Sjorgen's syndrome, glomerulonephritis, Neuromyelitis Optica (NMO), and IgG neuropathy.

In some embodiments, the antibody is in a kit comprising: (a) a composition comprising any one of the antibodies described herein (in a monospecific, bi-specific, or multi-specific format) and (b) a package insert comprising instructions for administering the composition to a subject to treat or delay progression of a cell proliferative disorder. In some embodiments, the antibody within the kit is lyophilized.

In any of the preceding uses or methods, the subject can be a human.

Dose and Route of Administration

The antibodies and compositions of the invention will be administered in an effective amount for treatment of the condition in question, i.e., at dosages and for periods of time necessary to achieve a desired result. A therapeutically effective amount may vary according to factors such as the particular condition being treated, the age, sex and weight of the patient, and whether the anti-CD3 antibodies are being administered as a stand-alone treatment or in combination with one or more additional anti-cancer treatments.

An effective amount for tumor therapy may be measured by its ability to stabilize disease progression and/or ameliorate symptoms in a patient, and preferably to reverse disease progression, e.g., by reducing tumor size. The ability of an antibody or composition of the invention to inhibit cancer may be evaluated by in vitro assays, e.g., as described in the examples, as well as in suitable animal models that are predictive of the efficacy in human tumors. Suitable dosage regimens will be selected in order to provide an optimum therapeutic response in each particular situation, for example, administered as a single bolus or as a continuous infusion, and with possible adjustment of the dosage as indicated by the exigencies of each case.

In some embodiments, the antibody is administered to the subject in a dosage of about 0.01 mg/kg to about 10 mg/kg. In some embodiments, the antibody is administered to the subject in a dosage of about 0.1 mg/kg to about 10 mg/kg. In some embodiments, the antibody is administered to the subject in a dosage of about 1 mg/kg. In some embodiments, the antibody is administered subcutaneously, intravenously, intramuscularly, topically, orally, transdermally, intraperitoneally, intraorbitally, by implantation, by inhalation, intrathecally, intraventricularly, or intranasally. In some embodiments, the antibody is administered subcutaneously. In some embodiments, the bispecific antibody is administered intravenously.

Pharmaceutical Formulations

Pharmaceutical formulations of the antibodies of the invention may be prepared by mixing such antibody having the desired degree of purity with one or more optional pharmaceutically acceptable carriers (Remington's Pharmaceutical Sciences 16th edition, Osol, A. Ed. (1980)), in the form of lyophilized formulations or aqueous solutions. Pharmaceutically acceptable carriers are generally nontoxic to recipients at the dosages and concentrations employed, and include, but are not limited to: buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid and methionine; preservatives (such as octadecyldimethylbenzyl ammonium chloride; hexamethonium chloride; benzalkonium chloride; benzethonium chloride; phenol, butyl or benzyl alcohol; alkyl parabens such as methyl or propyl paraben; catechol; resorcinol; cyclohexanol; 3-pentanol; and m-cresol); low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, histidine, arginine, or lysine; monosaccharides, disaccharides, and other carbohydrates including

glucose, mannose, or dextrans; chelating agents such as EDTA; sugars such as sucrose, mannitol, trehalose or sorbitol; salt-forming counter-ions such as sodium; metal complexes (e.g. Zn-protein complexes); and/or non-ionic surfactants such as polyethylene glycol (PEG). Exemplary pharmaceutically acceptable carriers herein further include interstitial drug dispersion agents such as soluble neutral-active hyaluronidase glycoproteins (sHASEGP), for example, human soluble PH-20 hyaluronidase glycoproteins, such as rHuPH20 (HYLENEX®, Baxter International, Inc.). Certain exemplary sHASEGPs and methods of use, including rHuPH20, are described in US Patent Publication Nos. 2005/0260186 and 2006/0104968. In one aspect, a sHASEGP is combined with one or more additional glycosaminoglycanases such as chondroitinases.

Exemplary lyophilized antibody formulations are described in U.S. Pat. No. 6,267,958. Aqueous antibody formulations include those described in U.S. Pat. No. 6,171,586 and WO2006/044908, the latter formulations including a histidine-acetate buffer.

The formulation herein may also contain more than one active ingredient as necessary for the particular indication being treated, preferably those with complementary activities that do not adversely affect each other. For example, it may be desirable to further provide an additional therapeutic agent (e.g., a chemotherapeutic agent, a cytotoxic agent, a growth inhibitory agent, and/or an anti-hormonal agent, such as those recited herein above). Such active ingredients are suitably present in combination in amounts that are effective for the purpose intended. Active ingredients may be entrapped in microcapsules prepared, for example, by coacervation techniques or by interfacial polymerization, for example, hydroxymethylcellulose or gelatin-microcapsules and poly-(methylmethacrylate) microcapsules, respectively, in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nano-particles and nanocapsules) or in macroemulsions. Such techniques are disclosed in Remington's Pharmaceutical Sciences 16th edition, Osol, A. Ed. (1980).

Sustained-release preparations may be prepared. Suitable examples of sustained-release preparations include semipermeable matrices of solid hydrophobic polymers containing the bispecific antibody, which matrices are in the form of shaped articles, for example, films, or microcapsules. The formulations to be used for in vivo administration are generally sterile. Sterility may be readily accomplished, e.g., by filtration through sterile filtration membranes.

EXAMPLES

Example 1

The present example provides the results of a hybridoma-based method used to generate multiple, different anti-CD3 antibodies of the invention.

5 Briefly, the multiple, different anti-CD3 antibodies were produced using hybridoma technology. Although the present inventors have used this methodology to produce a large number of anti-CD3 antibodies, this Example provides 58 CD3-specific antibodies discovered by the present inventors. By elucidating the amino acid and/or nucleotide sequences of these antibodies' CDR3 heavy chain (HC) and light chain (LC or λ) variable regions, the present inventors identified
10 a series of "clusters" or "motifs" within the sequences. These clusters represent convergent somatic hypermutations (SHM) in the variable region sequences. Clustering may provide insight into the functionally related sequences and the diversity of the total population of antibodies and their variable region sequences. Sequences that are descendants from the same parent B cell or convergently evolved the sequences in the same cluster should be functionally more related than
15 sequences belonging to other clusters.

In the present Example, the variable chain region sequences were provided by different antibodies. Thus, any clustering can most likely be attributed to convergent SHM, which are likely functionally related mutations, e.g., they share a specific affinity for CD3. These SHM may inform the development of recombinant anti-CD3 antibodies with improved properties, such as specific
20 binding for CD3 fragments.

Hybridomas

Hybridomas were produced that each expressed a different anti-CD3 antibody. Methods known in the art to generate antibodies and/or hybridoma cells are known in the art.

25 For example, different anti-CD3 antibodies are obtained from different populations of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical and/or bind the same epitope except for possible variants that arise during production of the monoclonal antibody, such variants generally being present in minor amounts.

By way of example, in certain hybridoma methods, a mouse or other appropriate host
30 animal, such as a hamster, is immunized with a CD3 antigen to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the protein used for immunization.

Alternatively, lymphocytes may be immunized in vitro. Lymphocytes then are fused with myeloma cells using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell.

5 The hybridoma cells thus prepared are seeded and grown in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells. For example, if the parental myeloma cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substances prevent the growth of HGPRT-deficient cells.

10 In some embodiments, the myeloma cells are those that fuse efficiently, support stable high-level production of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. Among these, in some embodiments, the myeloma cell lines are murine myeloma lines, such as those derived from mouse tumors. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal
15 antibodies.

Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against CD3. In some embodiments, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent
20 assay (ELISA).

After hybridoma cells are identified that produce antibodies of the desired specificity, affinity, and/or activity, the clones may be subcloned by limiting dilution procedures and grown by standard methods. Suitable culture media for this purpose include, for example, D-MEM or RPMI-1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in
25 an animal.

The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, chromatography, gel electrophoresis, dialysis, or affinity chromatography.

30 DNA encoding the monoclonal antibodies is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding

specifically to genes encoding the heavy and light chains of murine antibodies). In some embodiments, the hybridoma cells serve as a source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as E. coli cells, simian COS cells, Chinese Hamster Ovary (CHO) cells, or myeloma cells that do not otherwise
5 produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells.

In a further embodiment, antibodies or antibody fragments can be isolated from antibody phage libraries generated using the techniques described in McCafferty et al., Nature, 348:552-554 (1990). Clackson et al., Nature, 352:624-628 (1991) and Marks et al., J. Mol. Biol., 222:581-
10 597 (1991) describe the isolation of murine and human antibodies, respectively, using phage libraries. Subsequent publications describe the production of high affinity (nM range) human antibodies by chain shuffling (Marks et al., Bio/Technology, 10:779-783 (1992)), as well as combinatorial infection and in vivo recombination as a strategy for constructing very large phage libraries (Waterhouse et al., Nuc. Acids. Res., 21:2265-2266 (1993)). Thus, these techniques are
15 viable alternatives to traditional monoclonal antibody hybridoma techniques for isolation of monoclonal antibodies.

The DNA also may be modified, for example, by substituting the coding sequence for human heavy- and light chain constant domains in place of the homologous murine sequences, or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence
20 for a non-immunoglobulin polypeptide.

Sequencing

Briefly, a cDNA library was created from anti-CD3 antibody mRNA obtained from a
25 hybridoma culture of cells that express different anti-CD3 antibodies.

The sequences were aligned and all the unique anti-CD3 antibody sequences identified. Alignments the sequences revealed the uniqueness of each particular sequence and their corresponding antibodies. The antibody variable sequences of each different antibody were analyzed using a custom AlivaAlign sequencing software.

The identified CDR3 HC and VL sequences are provided in Table 1 as SEQ ID NOS: 2-
30 59 and 60-117. The corresponding heavy chain variable region and light chain variable region

sequences are provided in Table 2 as SEQ ID NOS: 118-175 and 176-233 respectively. As shown in Table 2, the heavy and light chain variable sequences were paired and identified to a particular antibody.

5 *Clustering*

The HC and VL chain variable region sequences were clustered using an algorithm that assigned sequences to a cluster when they shared identical hV, hJ, IV, and IJ genes, had identical HCDR3 lengths and were at least 90% identical (Hamming distance) for their HCDR3s within the cluster.

10 Table 3 provides the identified SMH for each heavy and light chain variable region sequence. In Table 3, the respective HC and VL variable region sequences are identifiable and pairable by the identity of their associated CD3 antibody.

15 Table 4 provides data that informs the listed clusters, i.e., 1-8, of the CD3 antibodies. The table also shows gene usage for heavy and light chains as well as difference to germline sequences as a percentage. The corresponding and paired HC and VL CDR3 sequences are also provided. Within each cluster are related sequences or “subclusters”, e.g., 1.1 and 1.2. Surprisingly, some antibodies provided identical heavy and light chain nucleotide sequences between the antibodies.

TABLE 1		
CDR3 Sequences		
CDR3 VH Amino Acid Sequence	Type	SEQ ID NO:
ARMRYNYFDY	VH	SEQ ID NO: 2
ARVKWDLDDY	VH	SEQ ID NO: 3
AREQWEPLYFDY	VH	SEQ ID NO: 4
AHLAHWGPYFDY	VH	SEQ ID NO: 5
ARFSNWGSGGYFDY	VH	SEQ ID NO: 6
ARERWALLFFEY	VH	SEQ ID NO: 7
AREKWELLYFDY	VH	SEQ ID NO: 8

ARERWELLFFDY	VH	SEQ ID NO: 9
AREKWVQLYFDF	VH	SEQ ID NO: 10
ARFSNWGSGGYFDY	VH	SEQ ID NO: 11
ARMRYNYFDY	VH	SEQ ID NO: 12
ARVVRDYGMDV	VH	SEQ ID NO: 13
ARERWELLFFDY	VH	SEQ ID NO: 14
AREKWELLYFDY	VH	SEQ ID NO: 15
AREKWVQLYFDF	VH	SEQ ID NO: 16
AREQWELLYFDY	VH	SEQ ID NO: 17
ARGGVGRNYYYYGMDV	VH	SEQ ID NO: 18
ARDGRWELLAFDI	VH	SEQ ID NO: 19
AKMRYNYFDY	VH	SEQ ID NO: 20
AKMRYNYFDY	VH	SEQ ID NO: 21
AREQWELLYFDY	VH	SEQ ID NO: 22
AREQWELLYFDY	VH	SEQ ID NO: 23
ARMRYNYFDY	VH	SEQ ID NO: 24
ARERYNYFDY	VH	SEQ ID NO: 25
ARERYNYFDY	VH	SEQ ID NO: 26
ARERYNYFDY	VH	SEQ ID NO: 27
ARERYNYFDF	VH	SEQ ID NO: 28
AREPYNYFDY	VH	SEQ ID NO: 29
ARERYNYFDY	VH	SEQ ID NO: 30
ARERYNYFDY	VH	SEQ ID NO: 31
ARERYNYFDY	VH	SEQ ID NO: 32
ARERYNYFDS	VH	SEQ ID NO: 33

ARVRRNYGMDV	VH	SEQ ID NO: 34
ARVRRNYGMDV	VH	SEQ ID NO: 35
ARVRRNYGMDV	VH	SEQ ID NO: 36
ARERWELLYFDY	VH	SEQ ID NO: 37
ARVRRNYGMDV	VH	SEQ ID NO: 38
ARVRRNYGMDV	VH	SEQ ID NO: 39
ARVRRNYGMDV	VH	SEQ ID NO: 40
ARVRRNYDMDV	VH	SEQ ID NO: 41
ARVRRNYGMDV	VH	SEQ ID NO: 42
AREQWVLLYFDY	VH	SEQ ID NO: 43
ARVRRNYGMDV	VH	SEQ ID NO: 44
ARVRRNYGMDV	VH	SEQ ID NO: 45
ARVRRNYGMDV	VH	SEQ ID NO: 46
ARERWELLYFDY	VH	SEQ ID NO: 47
ARERKYPLQFDY	VH	SEQ ID NO: 48
VRERKYPLYFDY	VH	SEQ ID NO: 49
ARVRRNYGMDV	VH	SEQ ID NO: 50
ARVRRNYGMDV	VH	SEQ ID NO: 51
ARERKYPLQFDY	VH	SEQ ID NO: 52
ARERKYLLYFDY	VH	SEQ ID NO: 53
ARVRRNYGMDV	VH	SEQ ID NO: 54
ARVRRNYGMDV	VH	SEQ ID NO: 55
ARVRRNYDMDV	VH	SEQ ID NO: 56
ARERKYLLYFDY	VH	SEQ ID NO: 57
SRERWNLLFFDY	VH	SEQ ID NO: 58

ARELWELLYFDY	VH	SEQ ID NO: 59
CDR3 λ chain sequences	Type	SEQ ID NO
SYDSSNVVF	λ	SEQ ID NO: 60
QAWDSSTHVV	λ	SEQ ID NO: 61
SYDSSNRVF	λ	SEQ ID NO: 62
CSYAGSSTLI	λ	SEQ ID NO: 63
QVWDSSSDHPV	λ	SEQ ID NO: 64
SFDRSNRVF	λ	SEQ ID NO: 65
SYDSSNRVF	λ	SEQ ID NO: 66
SYDSSNRVF	λ	SEQ ID NO: 67
SYDSSNRVF	λ	SEQ ID NO: 68
QVWDSTSDHPV	λ	SEQ ID NO: 69
SYDSSNVVF	λ	SEQ ID NO: 70
SYDSSNHVVF	λ	SEQ ID NO: 71
SYDSSNRVF	λ	SEQ ID NO: 72
SYDSSNRVF	λ	SEQ ID NO: 73
SYDSSNRVF	λ	SEQ ID NO: 74
SYDSSNRVF	λ	SEQ ID NO: 75
GTWDSSLSAVV	λ	SEQ ID NO: 76
QSADTSGTYRV	λ	SEQ ID NO: 77
SYDGSNVVF	λ	SEQ ID NO: 78
SYDGSNVVF	λ	SEQ ID NO: 79
SYDSSNRVF	λ	SEQ ID NO: 80

SYDRSNRVF	λ	SEQ ID NO: 81
SYDSSNVVF	λ	SEQ ID NO: 82
SYDSSNRVF	λ	SEQ ID NO: 83
SYNGNNRVF	λ	SEQ ID NO: 84
SYDNSNRVF	λ	SEQ ID NO: 85
SYDGSNRVF	λ	SEQ ID NO: 86
SYDNSNRVF	λ	SEQ ID NO: 87
SYDSSNRVF	λ	SEQ ID NO: 88
SYDNNNRVF	λ	SEQ ID NO: 89
SYDNSNRVF	λ	SEQ ID NO: 90
SYDGNNRVF	λ	SEQ ID NO: 91
SYDSSNHVVF	λ	SEQ ID NO: 92
SYDSSNHVVF	λ	SEQ ID NO: 93
SYDSSNWVF	λ	SEQ ID NO: 94
SYDSSNRVF	λ	SEQ ID NO: 95
SYDSNNHVVF	λ	SEQ ID NO: 96
SYDSSNHVVF	λ	SEQ ID NO: 97
SYDSSDHVVF	λ	SEQ ID NO: 98
SYDSGNWVF	λ	SEQ ID NO: 99
SYDSNNWVF	λ	SEQ ID NO: 100
SYDSSNRVF	λ	SEQ ID NO: 101
YDSSNHVVF	λ	SEQ ID NO: 102
SYDSSNHVVF	λ	SEQ ID NO: 103
SYDSSNHVVF	λ	SEQ ID NO: 104

SYDSSNRVF	λ	SEQ ID NO: 105
SYDSSNRVF	λ	SEQ ID NO: 106
SYDSSVRVF	λ	SEQ ID NO: 107
SYDSSNWVF	λ	SEQ ID NO: 108
SYDSSNHWVF	λ	SEQ ID NO: 109
SFDSSNRVF	λ	SEQ ID NO: 110
SYDRSNRVF	λ	SEQ ID NO: 111
SYDSSNHWVF	λ	SEQ ID NO: 112
SYDSSNHWVF	λ	SEQ ID NO: 113
SYDSSNHWVF	λ	SEQ ID NO: 114
SFDNSNRVF	λ	SEQ ID NO: 115
SYDSSNRVF	λ	SEQ ID NO: 116
SYDSNRVF	λ	SEQ ID NO: 117

Table 2 -- anti-CD3 antibody Variable Region Sequences

Antibody	HC Sequence	SEQ ID NO: 118	LC Sequence	SEQ ID NO: 176
70701_05 C15A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSMNW VRQTPGKGLEWVSSISNSCNYYADSVKGRFTISRDS AKNSLYLQMNSLRAEDTAVYFCARMRYNYFDYWG QGTLVTVSS	SEQ ID NO: 118	NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVVYEDNQRPSPGVPDFRFS GSIASSNSASLTISGLKTEDEADYYCQSYDSS NVVFGGGTKLTVL	SEQ ID NO: 176
70701_05 D06A	EVQLVESGGGLVQPGGSLRLSCVASGFTFSNYWMN WVRQAPGKGLEWVANIKKDGSEKYYVDSVKGRFTIS RDNAKNSLYLQMNSLRAEDTAVYFCARVKWDLDDY WGQGTILVTVSS	SEQ ID NO: 119	SYELTQPPSVSVSPGQTASITCSGDKLGHKYV CWYQQKPGQSPVLVIYQDNKRKPSGIPERFSGS NSGNTATLTISGTQAMDEADYYCQAWDSSTH VVFVGGGTKLTVL	SEQ ID NO: 177
70701_05 L17A	EVQLVESGGGLVKPGGSLRLSCAASGFNLSTYSMNW VRQAPGKGLEWVSSISRSSYYIYADAVKGRFTISR NAQNSLYLQMNSLRAEDTAVYFCAREQWEPLYFDY WGQGTILVTVSS	SEQ ID NO: 120	NVMLIQPHSVSESPGKTVTISCTRSSGSIASKY VQWYQQRPGSSPTTVVYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFVGGGTKLTVL	SEQ ID NO: 178
70701_05 M03A	QITLKESGPTLVKPTQTLITCSFSGFSTTSVGVGW IRQPPGKALECLALYWNDDKRYSPSLKNRLTIKDT KNQVLTMTNMDPVDTATYFCAHLAHWGPYFDYW GQGTILVTVSS	SEQ ID NO: 121	QSALTQSASVSGSPGQSITISCTGTSSDVGSYN LVSWYQQHPGKAPKLMIEGSKRPSGVSNRF SGSKSGNTASLTISGLQAEDEADYYCCSYAGS STLIFGGGTKLTVL	SEQ ID NO: 179
70701_06 A03A	EVLLVESGGGLVKPGGSLRLSCEASGFTFSRYSMNW VRQAPGKGLEWVSSISRSSRYIYADSVKGRFTISR NAKNSLYLQMISLRAEDTAVFYCARFSNWGSGGYFD YWGQGTILVTVSS	SEQ ID NO: 122	SYVLTQPPSVSVAPGQTASITCGGNIGSKSVH WYQQKPGQAPVLVYVDDTDRPSGIPERFSGS NSGNTATLTISRVEAGDEADYYCQVWDDSSD HPVFGGGTKLTVL	SEQ ID NO: 180
70701_06 A16A	EVQLVESGGGLVKPGGSLRLSCAASGFTVNNYSMNW VRQAPGKGLEWVSSISRSSNYIYADSVKGRFTISR	SEQ ID NO: 123	NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDYQRPSPGVPDFRFSG	SEQ ID NO: 181

	NAKNSLYQMNSLRAEDTAVYYCARERWALLFFEY WGQGLVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSFDRSN RVFGGTKLTVL	
70701_06 B15A	EVQLVESGGGLVKPGGSLRLSCAASGENLSTYSMNW VRQAPGKGLEWVSSISRSSSIYYADSVKGRFTISRDN AQNLSLYQMNSLRAEDTAVYYCAREKWELLYFDYW GQGLTVTVSS	SEQ ID NO: 124		NVMLIQPHSVSESPGKTVTISCTRSSGSIASKY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 182
70701_06 M17A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSSYSMNW VRQAPGKGLEWVASMSRSSSIYYAHSVKGRFTISRDN NAKNSLYQMNSLRAEDTAVYYCARERWELFFDY WGQGLVTVSS	SEQ ID NO: 125		NFMLTQPHSVSESPGKTVTISCTRSSGNIARNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 183
70701_07 N16A	EVQLVESGGGLVKPGGSLRLSCAASGFTLSSYNMDW VRQAPGKGLEWVSSISRSSSIYYADSVKGRFTISRDN NAENSLYQMNSLRAEDTAVYYCAREKWVQLYDFD WGQGLVTVSS	SEQ ID NO: 126		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNF VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 184
70701_08 J03A	EVQLVESGGGLVKPGGSLRLSCAASGFTFRYSMNW VRQAPGKGLEWVSSISRSSSIYYADSVKGRFTISRDN AKNSLYQMNSLRAEDTAVYYCAREFSNWGSGGYFD YWGQGLVTVSS	SEQ ID NO: 127		SYVLTQPPSVSVAPQGTARITCGGNNIGKSKSV HWYQQKFGQAPVLVYDDSDRPSGIPERFSG SNSENTATLTSRVEAGDEADYYCQVVDST DHPVFGGGTKLTVL	SEQ ID NO: 185
70701_09 C02A	EVQLVESGGGLVKPGGSLRLSCAASGFTFRYSMNW VRQTPGKGLEWVSSISSNVIYYADSVKGRFTISRDN AKNSLYQMNSLRAEDTAVYYCARMRYNYFDYWG QGTLVTVSS	SEQ ID NO: 128		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFS GSIASSNSASLTISGLKTEDEADYYCQSYDSS NVVFGGGTKLTVL	SEQ ID NO: 186
70701_09 K17A	EVQLVESGGGLGQPGGSLRLSCAASGFTFSSYSMNW VRQAPGKGLEWVYSISRSSSIYYADSVKGRFHSRDK AKNSLYQMNSLRDEDTAVYYCARVVRDYGMDVW GQGTTVTVSS	SEQ ID NO: 129		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 187
70701_10 D02A	EVQLVESGGGLVKPGGSLRLSCAASGFTLSSYSMNW VRQAPGKGLEWVSTMRSRSSSIYYADSVKGRFTISRDN	SEQ ID NO: 130		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG	SEQ ID NO: 188

	NAKNSLYQMNSLRAEDTAVYYCARERWELLFHDY WGQGTLVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	
70701_10 F19A	EVQLVESGGGLVKPGGSLRLSCAASGFTISRYSMNW VRQAPGKGLEWVSSISRSSSHIYYADSVKGRFTISRDN AKNSMYLQMNSLRAEDTAVYYCAREKWELLYFDY WGQGTLVTVSS	SEQ ID NO: 131		NVMLTQHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 189
70701_10 J09A	EVQLVESGGGLVKPGGSLRLSCAASGFTLSSYSMDW VRQAPGKGLEWVSSISRSSNYIYYADSVKGRFTISRDRD NAENSLYLQMNSLRAEDTAVYYCAREKWVQLYFDF WGQGTLVTVSS	SEQ ID NO: 132		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNF VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 190
70701_10 N20A	EVQLVEFGGGLVRPGGSLRLSCVASGFTSFIFTMNVV RQAPGKGLEWVSTSSRSTSYIYYADSVKGRFTISRDN AKNSLYLQMNSLRAEDTAVYYCAREQWELLYFDYW GQGTLVTVSS	SEQ ID NO: 133		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 191
70701_11 B10A	QVQLVQSGAEVKKPGSSVKVSKCASGGTFSSYAISW VRQAPGQGLEWMGGHPIFATANYAQKFKGRVITAD KSTSTAYMELSSLRSEDTAVYYCARGGVGRNYYYYG MDVWGGQGTITVTVSS	SEQ ID NO: 134		QSVLTQPPSVSAAPGQKVTISCSGSSNIGNNY VSWYQQLPGTAPKLLIYDNYKRPSGIPDRFSG SKSGTSA TLGHITGLQGTGDEADYYCGTWDSSLS AVVFGGGTKLTVL	SEQ ID NO: 192
70701_11 K23A	EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYGMNW VRQAPGKGLEWVYSISRSTRYYADSVKGRFTISRDRD NAKNSLYLQMNSLRDEDTSVYYCARDGRWELLAFDI WGQGTMTVTVSS	SEQ ID NO: 135		SYELTQPPSVSVSPGQTARITCSGDALPKQYA YWYQKPGQAPLVLYKDSERPSGIPERFSGS SSGTTVTLTISGVQAEDEADYYCQASADTSPTY RVFGGGTKLTVL	SEQ ID NO: 193
70701_11 L01A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRY SINWV RQAPGKGLEWVSSISSSSSYIYYADSLKGRFTISRDN RNSLYLQMNSLRAEDTAVYYCAKMRNYFDYWGQ GALVTVSS	SEQ ID NO: 136		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN VVFGGGTKLTVL	SEQ ID NO: 194
70701_11 M11A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRY SINWV RQAPGKGLEWVSSISSSSSYIYYADSLKGRFTISRDN	SEQ ID NO: 137		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG	SEQ ID NO: 195

	RNSLYLQMNLSRAEDTAVYYCAKMRNYFDYWGG GALVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDGSN VVFGGTKLTVL	
70701_12 F16A	EVQLVEFGGGLVRPGGSLRLSCVASGFTFSFTMNWV RQAPGKGLEWVSTSSRSTSYIYADSVKGRFTISRDN AKNSLYLQMNLSRAEDTAVYYCAREQWELLYFDYW GGTLVTVSS	SEQ ID NO: 138		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 196
70701_12 J12A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYINWV RQAPGKGLEWVSSSSRSSYIYADSVKGRFTISRDN AKNSLYLQMNLSRAEDTAVYYCAREQWELLYFDYW GGTLVTVSS	SEQ ID NO: 139		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDRSN RVFGGTKLTVL	SEQ ID NO: 197
70701_12 O16A	EVQLVESGGGLVKPGGSLRLSCGASGFTFSYSMNW VRQAPGKGLEWVSSISSRVIYADSLRGRFTISRDN AKNSLYLQMNLSRAEDTAVYYCARMRYNYFDYW GGTLVTVSS	SEQ ID NO: 140		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN VVFGGTKLTVL	SEQ ID NO: 198
70701_13 C01A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSMNW VRQAPGKGLEWVSSISGSSRYIYADSVKGRFTVSRD NAKNSLYLQMSLSPEDTAVYYCARERYNYFDYW GGTLVTVSS	SEQ ID NO: 141		NFMLTQPHSVSESPGKTVTISCTRSSGSIANNF VQWFQRPFGSPTIDVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 199
70701_13 G21A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSIYSMNWV RQAPGKGLEWVSSISKSTFIYADSVKGRFTISRDN KNSLYLQMNLSAEDTAVYYCARERYNYFDYWGGQ TLVTVSS	SEQ ID NO: 142		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYSCQSYNGNN RVFGGTKLTVL	SEQ ID NO: 200
70701_13 H15A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSLNW VRQAPGKGLEWVSSISSRVIYADSVQGRFTISRDN AKNSLYLQMNLSRAEDTAVYYCARERYNYFDYWGGQ GTLVTVSS	SEQ ID NO: 143		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYFCQSYDSSN RVFGGTKLTVL	SEQ ID NO: 201
70701_14 F08A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSMNW VRQAPGKGLEWVSSISSRVIYADSVKGRFTISRDN	SEQ ID NO: 144		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTAVIYEDNQRPSPGVPPDRFSG	SEQ ID NO: 202

	AKNSLYLQMNSLRAEDTAVVYCARERYNYFDYWGQ GTLVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	
70701_14 F21A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSMNW VRQAPGKGLEWVSSISSSRYIFYADSVKGRFTISRDN NAKNSLYLQMNSLRAEDTAVVYCARERYNYFDYWGQ QGTLVSVSS	SEQ ID NO: 145		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	SEQ ID NO: 203
70701_14 H05A	EVHLVESGGGLVKPGGSLRLSCAASGFTFSRYSMNW VRQAPGKGLEWVSSISSSRYIFYADSVKGRFTISRDN AKNSLYLQMNSLRAEDTAVVYCARERYNYFDYWGQ GTLVTVSS	SEQ ID NO: 146		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	SEQ ID NO: 204
70701_14 K01A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSRYSLNW VRQAPGKGLEWVSSISSSRYIFYADSVKGRFTISRDN AKNSLYLQMNSLRAEDTAVVYCARERYNYFDYWGQ GTLVTVSS	SEQ ID NO: 147		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	SEQ ID NO: 205
70701_14 K02A	EVQLVESGGGLVKRGGSLRLSCATSGFTLSRYSMNW VRQAPGKGLEWVSSISSSRYIFYADSVKGRFTISRDN AKNSLYLQMNSLRAEDTAVVYCARERYNYFDYWGQ GTLVTVSS	SEQ ID NO: 148		DFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTAVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	SEQ ID NO: 206
70701_14 M09A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSIYSMNWV RQAPGKGLEWVSSISKSSRYIFYADSVKGRFTISRDN KNSLYLQMNSLRAEDTAVVYCARERYNYFDSWGQ TLVTVSS	SEQ ID NO: 149		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN RVFGGGTKLTVL	SEQ ID NO: 207
70701_16 D18A	EVQLVESGGGSVQPGGSLRLTCAASGFTFSIYSMNW VRQAPGKGLEWVSYISRSSSTIYADSVKGRFTISRDN AKNSLYLQMNSLRDEDTAVVYCARVRRNYGMDVW QGTTVTVSS	SEQ ID NO: 150		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPPDRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDGSN HWVFGGGTKLTVL	SEQ ID NO: 208
70701_16 E13A	EVQLVESGGGLVQPGGSLRLSCAASGFTFSIYSMNW VRQAPGKGLEWVSYISRSSTIYADSVKGRFTISRDN	SEQ ID NO: 151		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPPDRFSG	SEQ ID NO: 209

	NAKNSLYQMNSLRDEDTA VYYCARVRRNYGMDV WGQGTITVSS			SIDSSNSASLTISGLKTEDEAEAYCQSYDSSN HWVFGGGTKLTVL	
70701_16 L13A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSMNW VRQAPGKGLEWVSYISRSSSTIYYADSVKGRFTISRDN AKNSLYQMNSLRDEDTA VYYCARVRRNYGMDVW GQGTITVTVSS	SEQ ID NO: 152		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISRLKTEDEADY CQSYDSSN WVFGGGTKLTVL	SEQ ID NO: 210
70701_16 P13A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSMNW VRQAPGKGLEWVSSSSRSSTIYYADSVKGRFTITRD NAKNSLYQMNSLRAEDTA VYYCARERWELLYFDY WGQGTITVTVSS	SEQ ID NO: 153		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADY CQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 211
70701_17 A23A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSMNW VRQAPGKGLEWVSYISRSSNTIYYADSVKGRFTISRDN NARNLSLYQMNSLRDEDAA VYYCARVRRNYGMDV WGQGTITVTVSS	SEQ ID NO: 154		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADY CQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 212
70701_17 G08A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSMNW VRQAPGKGLEWVSYISSSGSTIYYADSVKGRFTISRDN NAKNSLYQMNSLRDEDTA VYYCARVRRNYGMDV WGQGTITVTVSS	SEQ ID NO: 155		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADY CQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 213
70701_17 H17A	EVQLVESGGGLVQPGGSLRLSCVVSFGFTSSYSMNW VRQAPGKGLEWVSYISSHSSTIYYADSVKGRLTISRDN NAKNSLYQMNSLRDEDTA VYYCARVRRNYGMDV WGQGTITVTVSS	SEQ ID NO: 156		KLMLTQPHSVSESPGKTVTISCTRSSGSIARNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLRTEDEADY CQSYDSSD HWVFGGGTKLTVL	SEQ ID NO: 214
70701_18 A16A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSMNW VRQAPGKGLEWVSYISSSNTIYYADSVKGRFTISRDN AKNSLYQMNSLRDEDTA VYYCARVRRNYDMDVW GQGTITVTVSS	SEQ ID NO: 157		NFMLTQPHSVSESPGKTVTISCTRSSGNIARNY VQWYQQRPGSSPTTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADY CQSYDSGN WVFGGGAKLTVL	SEQ ID NO: 215
70701_18 E19A	EVQLVESGGGLVQPGGSLRLSCAASGFTSSYSVNW VRQAPGKGLEWVSYISRSSSIYYVDSVKGRFTISRDN	SEQ ID NO: 158		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTTVIYDDNQRPSPGVPDFRFSG	SEQ ID NO: 216

	AKNSLYLQMNSLRDEDTAVYYCARVRRNYGMDVW GQGTTVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDSSN WVFGGGTKLTVL	
70701_18 M14A	EVQLVESGGGLVKPGGSLRLSCAASGFIILNNYSMNW VRQAPGKGLWVSSISRSSSTIYYADSVKGRFTISRDN AKNSLYLQMNSLRDEDTAVYYCAREQWVLLYFDY WGQGTLLTVTVSS	SEQ ID NO: 159		NFMLTQPHSVSESPGKTTITICTRRRNGNIASNY VQWYQQRPGSPPTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 217
70701_19 A10A	EAQLVESGGGLVQPGGSLRLSCAASGFTFSSYSMNW VRQAPGKGLWVSYISRSSSTIYYADSAKGRFTISRDN AKNSLYLQMNSLRDEDTAVYYCARVRRNYGMDVW GQGTTVTVSS	SEQ ID NO: 160		HFMLTQPHSVSESPGKTVTICTRTRSSGSIASNY VQWYQQRPGSPPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 218
70701_19 C22A	EVQLVESGGGLVQPGESLRLSCAASGFTFSSYSMNW VRQAPGKGLWVSYISRSSSTIYYADSVKGRFTISRDN NARNSLYLQMNSLRDEDTAVYYCARVRRNYGMDV WGQGTTVTVTVSS	SEQ ID NO: 161		NFMLTQPHSVSESPGKTVTICTRTRSSGSIASNY VQWYQQRPGSPPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 219
70701_19 D03A	EVQLVESGGGLVQPGGSLRLSCAVSGFTFSSYSMNW VRQAPGKGLWVSYSSSSSTIYYADSVKGRFTISRDN NAKNSVYLQMNSLRDEDTAVYYCARVRRNYGMDV WGQGTTVTVTVSS	SEQ ID NO: 162		NFMLTQPHSVSESPGKTVTICTRTRSSGSIATNY VQWYQQRPGSPPTLIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 220
70701_19 E13A	EVHLVESGGGLVKPGGSLRLSCAASGIFTSRYSMNW VRQAPGKGLWVSSISRSSNYIYYADSVKGRFTISRDN NAKNSLYLQMNSLRAEDTAVYYCARERWELLYFDY WGQGTLLTVTVSS	SEQ ID NO: 163		NFMLTQPHSVSESPGKTVTICTRTRSSGSIASNY VQWYQQRPGSPPTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKLTVL	SEQ ID NO: 221
70701_19 I16A	QVQLQESGPGLVKPSQTLSTCTVSGGSITSGASYWS WIRQHPGKGLWIGIYIYSGSTYHSPSLKSRVTISVDI SKNRFSLMLSSVTAAADTAVYYCAREKYPQLQFDYWG QGTLLTVTVSS	SEQ ID NO: 164		NFMLTQPHSVSESPGKTVTICTRTRSSGSIASNY VQWYQQRPGSPPTVIYEDSQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFGGGTKVTVL	SEQ ID NO: 222
70701_19 J10A	QVQLQESGPGLMKPSQTLSTCTVSGGSISGVYYWN WIRQHPGKGLWIGIYIYSGSTIYNSPLKSRVTISVD	SEQ ID NO: 165		NFMLTQPHSVSESPGKTVTICTRTRSSGSIASNY VQWVQQRPGSPPTVIFEDNQRPSPGVPDFRFSG	SEQ ID NO: 223

	TSKNQFSLKLSVTAADTAVYYCVRRERKYPYFYDYW GQGLTVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDSSV RVFGGGTKLTVL	
70701_19 O15A	EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYSMNW VRQAPGKGLWEVSYISSSSTIYYADSVKGRFTISRDN AKNSLFLQMNLSLRDEDTAVYYCARVRRRNYGMDVW GQGTITVTVSS	SEQ ID NO: 166		NFVLPQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLTIEDEADYYCQSYDSSN WVFGGGTKLTVL	SEQ ID NO: 224
70701_20 O18A	EVQLVESGGGLVQPGGSLRLSCAASGLTFSRYSMNW VRQAPGKGLYWVSYISTSTIYYADSVKGRFTISR NAKNSLSLQMNLSLRDEDTAVYYCARVRRRNYGMDV WGQGTITVTVSS	SEQ ID NO: 167		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGEGTKLTVL	SEQ ID NO: 225
70701_21 A06A	QVQLQESGPGLVPEPSQTLSLICTVSGGSISSGASYWSW IRQHPGKGLWIGIYYSGTTFYENPSLKSRTISLDT KNQFSLKLSVTAADTAVYYCARERKYPLOFDYWG QGTITVTVSS	SEQ ID NO: 168		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQLRPGSSPTVIYEDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSFDSN RVFGGGTKLTVL	SEQ ID NO: 226
70701_21 J18A	QVQLQESGPGLVKPSQTLSLCTVSGGSISSGAYCWS WIRQHPGKGLWIGIYYSGTTFYENPSLKSRTISVDT SKKQFSLKLSVTAADTAVYYCARERKYLLEYFDYWG QGTITVTVSS	SEQ ID NO: 169		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYEDYQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDRSN RVFGGGTKLTVL	SEQ ID NO: 227
70701_22 D03A	EMQLVESGGGLVQRGGSLRLSCAASKFTFSGYAMN WVRQAPGKGLWEVSYISSTIYYADSVKGRFTISR DNAKNSLYLQMNLSLRDEDTAVYYCARVRRRNYGMD VWGQGTITVTVSS	SEQ ID NO: 170		YFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 228
70701_22 O09A	EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYSINWV RQAPGKGLWEVSYISSTIYYADSVKGRFTISRDN AKNSLYLQMNLSLRDEDTAVYYCARVRRRNYGMDVW GQGTITVTVSS	SEQ ID NO: 171		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYDDNQRPSPGVPDFRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGGTKLTVL	SEQ ID NO: 229
70701_23 F03A	EVQLVSGGGGLVQPGGSLRLSCAASGFTFSSYSMNW VRQAPGKGLWEVSYISSSSTIYYADSVKGRFTISRDN	SEQ ID NO: 172		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSSPTVIYEDNQRPSPGVPDFRFSG	SEQ ID NO: 230

	AKNSLYQMNSLRDEDTAVYYCARVRRNYDMDVW GQGTIVTVSS			SIDSSNSASLTISGLKTEDEADYYCQSYDSSN HWVFGGKLTIVL	
70701_23 N16A	QVQLRESGPGLVKPSQTLSTLCTVSGGSINSGGYCWS WIRQHPGKGLEWIGSIYSGTYNPSLKRVTISVDT SKNQFSLKLSVTAADTAVYYCARERKYLIFDYWG QGTLVTVSS	SEQ ID NO: 173		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSPPTVIYEDNQRPSPGVPRFSG SIDSSNSASLTISGLKTEDEADYYCQSFDNSN RVFAGGKLTIVL	SEQ ID NO: 231
70701_24 E09A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSYCMNW VRQAPGKGLEWVSSISRSSSIYIYADSVKGRFTISRDN AKNSLYQMNSLRAEDTAVYYCSRERWNLFFDYW GQGTIVTVSS	SEQ ID NO: 174		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSPPTVIYEDNQRPSPGVPRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFAGGKLTIVL	SEQ ID NO: 232
70701_24 N03A	EVQLVESGGGLVKPGGSLRLSCAASGFTFSYINWV RQAPGKGLEWVSSISRSSSIYIYADSVKGRFTISRDN KNSLYQMNSLRAEDTAVYYCARELWELLYFDYWG QGTLVTVSS	SEQ ID NO: 175		NFMLTQPHSVSESPGKTVTISCTRSSGSIASNY VQWYQQRPGSPPTVIYEDNQRPSPGVPRFSG SIDSSNSASLTISGLKTEDEADYYCQSYDSSN RVFAGGKLTIVL	SEQ ID NO: 233

Table 3 – identified somatic hypermutations in the CD3 antibodies

Heavy Chain									
Antibody	SHM	N- Glycosylation	Deamidation	Isomerisation (DG/DS)	Isomerisation (DT/DD/DH)	Aberrant Cysteines	Missing Cysteines	Integrin Binding	CDR Met and Trp
70701_05C15A	S31R A40T S53N S55C S56N N74S		NS NS NS	DS DS	DT	c54			CDR3
70701_05D06A	A23V S3IN S35N Q53K		NS NS	DG DS	DT				CDR1 CDR3
70701_05L17A	T28N F29L S31T S53R S63A K76Q	NL	NS NS		DT				CDR3

7070L_11M11A	S31R M34H V64L K76R	NS NS	DS	DT			CDR3
7070L_12F16A	S7F K13R A23V S31I Y32F S33T S50T I51S S53R S55T	NS NS	DS	DT			CDR3
7070L_12J12A	S31R M34H I51S S53R	NS NS	DS	DT			CDR3
7070L_12O16A	A23G S56R V64L K65R	NS NS	DS	DT			CDR3
7070L_13C01A	S31R S53G S56R I70V N84S R87S A88P	NS	DS	DT			
7070L_13G21A	S31I S53K S54T S56T Y57F Y59F R87S	NS NS	DS	DT			
7070L_13H15A	S31R M34L S56R Y59F K65Q Y95F	NS NS	DS	DT			
7070L_14F08A	S31R S56R Y59F	NS NS	DS	DT			
7070L_14F21A	S31R S56R Y59F I70M S85N	NS	DS	DT			
7070L_14H05A	Q3H S31R S56R	NS NS	DS	DT			
7070L_14K01A	S31R M34L S56R	NS NS	DS	DT			
7070L_14K02A	P14R A24T F29L S31R V93L	NS NS	DS	DT			
7070L_14M09A	S31I S53K S56R Y59F R87S	NS NS	DS DS	DT			
7070L_16D18A	L11S S21T S31T S53R	NS NS	DS	DT			CDR3
7070L_16E13A	S53R S55T	NS NS	DS	DT			CDR3
7070L_16L13A	S53R	NS NS	DS	DT			CDR3
7070L_16P13A	I51S S53R S71T	NS NS	DS	DT			CDR3
7070L_17A23A	S53R S56N K76R N84S T91A	NS NT	DS				CDR3

70701_17G08A	S55GN77S		NS	DS	DT				CDR3
70701_17H17A	A23V A24V S55H F68L		NS NS	DS	DT				CDR3
70701_18A16A	S55I S56N		NS NS NT	DS	DT				CDR3
70701_18E19A	M34V S53R T57I A61V		NS NS	DS	DT				CDR3
70701_18M14A	T28I F29L S30N S31N S53R E89D	NY	NS NS	DS	DT DD				CDR3
70701_19A10A	V2A S53R V64A		NS NS	DS	DT				CDR3
70701_19C22A	G16E S53R S56N K76R		NS NS NT	DS	DT				CDR3
70701_19D03A	A24V I51S L79V		NS NS	DS	DT				CDR3
70701_19E13A	Q3H F27I S31R S53R S56N		NS NS	DS	DT				CDR3
70701_19I16A	S30T G33A Y34S Y61H N62S Q79R K83M				DT DT				
70701_19J10A	V12M G33V S37N Y52D A98V				DT DT				
70701_19O15A	Y80F		NS NS	DS	DT				CDR3
70701_20O18A	F27L S31R E46Y S53T S56T Y80S		NS NS	DS	DT				CDR3
70701_21A06A	K13E T21I G33A Y34S S58T Y61F V73L				DT DT				
70701_21J18A	G33A Y35C Y52S N78K				DT DT	c34			
70701_22D03A	V2M P14R G26K S31G S33A S55T I58T		NS NS	DS	DT				CDR3

70701_22O09A	S31R M34I S54N S55T	NT	NS NS NT	DS	DT							CDR3
70701_23F03A	E6G		NS NS	DS	DT							CDR3
70701_23N16A	Q5R S30N Y35C Y52S		NS		DT DT	c34						
70701_24B09A	S33C S53R A97S		NS NS	DS	DT	c32						CDR3
70701_24N03A	M34I S53R		NS NS	DS	DT							CDR3
Corresponding Light Chain Sequences												
Antibody	SHM	N- Glycosylation	Deamidation	Isomerisation (DG/DS)	Isomerisation (DT/DD/DH)	Aberrant Cysteines	Missing Cysteines	Integrin Binding	CDR Met and Trp			
70701_05C15A	149V D68A		NS	DS								
70701_05D06A	D29H A32V S51N		NS NT	DS		c33						CDR3
70701_05L17A	F2V T51N32K		NS	DS DS								
70701_05M03A	P7S		NT			c91						
70701_06A03A	R19S S51T		NS NT	DS	DT DD DH							CDR3
70701_06A16A	T47P N53Y Y94F S96R		NS	DS								
70701_06B15A	F2V T51N32K		NS	DS DS								
70701_06M17A	S28N S31R		NS	DS DS								
70701_07N16A	Y33F N72K			DS DS								
70701_08J03A	G67E S93T		NS NT	DS DS	DD DH							CDR3
70701_09C02A	149V D68A		NS	DS								

70701_09K17A	I20F E51V			NS	DS DS				CDR3
70701_10D02A				NS	DS DS				
70701_10F19A	F2V P7T			NS	DS DS				
70701_10I09A	Y33F			NS	DS DS				
70701_10N20A				NS	DS DS				
70701_11B10A	N53Y				DS				CDR3
70701_11K23A	S92T				DS	DT			
70701_11L01A	S96G			NS	DG DS				
70701_11M11A	S96G			NS	DG DS				
70701_12F16A				NS	DS DS				
70701_12J12A	S96R			NS	DS				
70701_12O16A	S28N			NS	DS DS				
70701_13C01A	S31N Y33F Y37F T47D			NS	DS DS				
70701_13G21A	Y90S D95N S96G S97N			NG NS	DS				
70701_13H15A	T47P Y90F S96N			NS NS	DS				
70701_14F08A	S28N T47A S96G			NS	DG DS				
70701_14F21A	S96N			NS NS	DS				
70701_14H05A				NS	DS DS				
70701_14K01A	N53Y D61A S96N S97N			NS	DS				
70701_14K02A	T46A S95N			NS NS	DS				

70701_14M09A	N53Y Y90S S96G S97N		NS	DG DS					
70701_16D18A	E51D		NS	DS DS	DD				CDR3
70701_16E13A	E51D D88E		NS	DS DS	DD				CDR3
70701_16L13A	Y37F E51D G80R		NS	DS DS	DD				CDR3
70701_16P13A			NS	DS DS					
70701_17A23A	E51D S97N		NS	DS DS	DD				CDR3
70701_17G08A	E51D N72S			DS DS	DD				CDR3
70701_17H17A	NIK F2L S31R E51D K82R N98D		NS	DS DS	DD DH				CDR3
70701_18A16A	S28N S31R S97G		NS	DS DS					CDR3
70701_18E19A	E51D S97N		NS	DS DS	DD				CDR3
70701_18M14A	V18I S25R S26N S28N S43R		NG NS	DS DS					
70701_19A10A	E50D		NS	DS DS	DD				CDR3
70701_19C22A	E51D		NS	DS DS	DD				CDR3
70701_19D03A	S31T V48L E51D		NS	DS DS	DD				CDR3
70701_19E13A			NS	DS DS					
70701_19I16A	N53S		NS	DS DS DS					
70701_19J10A	Y37F Y50F		NS	DS DS					
70701_19O15A	M3V T5P T47S E51D N53S K82T		NS	DS DS DS	DD				CDR3
70701_20O18A	E51D		NS	DS DS	DD				CDR3
70701_21A06A	Q39L Y94F		NS	DS DS					

70701_21I18A	I29V N53Y S96R		NS	DS									
70701_22D03A	E50D		NS	DS DS	DD								CDR3
70701_22O09A	E51D		NS	DS DS	DD								CDR3
70701_23F03A			NS	DS DS									CDR3
70701_23N16A	Y94F S96N		NS NS	DS									
70701_24B09A			NS	DS DS									
70701_24N03A	S43C S97N		NS	DS DS						c42			

Table 4 -- variable region sequence clusters

Cluster	Identicals	Antibody	IsoType	V Gene	V % Match	D Gene	D % Match	J Gene	J % Match	CDR3	IsoType	V Gene	V % Match	J Gene	J % Match	CDR3
Clu st 1.1		70701_06A 03A	IGHG 1	IGHV 3-21	95.2 %	IGHD 7-27	100.0 %	IGH J4	97.8 %	ARFSNWGSGGYFD Y	λ	IGLV 3-21	99.0 %	IGL J2	100.0 %	QVWDSSSD HPV
Clu st 1.2		70701_08J0 3A	IGHG 2a	IGHV 3-21	99.0 %	IGHD 7-27	100.0 %	IGH J4	97.8 %	ARFSNWGSGGYFD Y	λ	IGLV 3-21	99.3 %	IGL J2	100.0 %	QVWDSTSD HPV
Clu st 2.1		70701_05C1 5A	IGHG 1	IGHV 3-21	96.6 %	IGHD 2-2	100.0 %	IGH J4	100.0 %	ARMRYNYFDY	λ	IGLV 6-57	99.0 %	IGL J2	100.0 %	SYDSSNVVVF
Clu st 2.2		70701_09C0 2A	IGHG 1	IGHV 3-21	98.0 %	IGHD 2-2	100.0 %	IGH J4	100.0 %	ARMRYNYFDY	λ	IGLV 6-57	99.0 %	IGL J2	100.0 %	SYDSSNVVVF
Clu st 2.3	identical _2	70701_11L0 1A	IGHG 1	IGHV 3-21	97.6 %	IGHD 2-2	100.0 %	IGH J4	97.9 %	AKMRYNYFDY	λ	IGLV 6-57	99.0 %	IGL J2	100.0 %	SYDGSNVV F

Clu st 2.3	identical _2	70701_11M 11A	IGHG 1	IGHV 3-21	97.6 %	IGHD 2-2	100.0 %	IGH J4	97.9 %	AKMRYNYFDY	λ	IGLV 6-57	99.0 %	IGL J2	100.0 %	SYDGSNVV F
Clu st 2.4		70701_12O 16A	IGHG 1	IGHV 3-21	98.3 %	IGHD 5-24	100.0 %	IGH J4	100.0 %	ARMRYNYFDY	λ	IGLV 6-57	99.3 %	IGL J2	97.3 %	SYDSSNVVF
Clu st 3.1		70701_05L1 7A	IGHG 1	IGHV 3-21	97.6 %	IGHD 1-14	100.0 %	IGH J4	97.8 %	AREQWEPLYFDY	λ	IGLV 6-57	97.3 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.10		70701_19E1 3A	IGHG 2a	IGHV 3-21	98.3 %	IGHD 1-26	100.0 %	IGH J4	100.0 %	ARERWELLYFDY	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.11		70701_24N 03A	IGHG 1	IGHV 3-21	98.6 %	IGHD 1-26	91.7 %	IGH J4	100.0 %	ARELWELLYFDY	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSNNRVF
Clu st 3.2		70701_06B1 5A	IGHG 1	IGHV 3-21	97.6 %	IGHD 1-26	92.3 %	IGH J4	100.0 %	AREKWELLYFDY	λ	IGLV 6-57	97.3 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.3		70701_06M 17A	IGHG 2b	IGHV 3-21	98.3 %	IGHD 1-26	100.0 %	IGH J4	100.0 %	ARERWELFFDY	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.4		70701_10D 02A	IGHG 2a	IGHV 3-21	98.0 %	IGHD 1-26	100.0 %	IGH J4	100.0 %	ARERWELFFDY	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.5		70701_10F1 9A	IGHG 1	IGHV 3-21	98.3 %	IGHD 1-26	100.0 %	IGH J4	100.0 %	AREKWELLYFDY	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.6	identical _1	70701_10N 20A	IGHG 1	IGHV 3-21	94.9 %	IGHD 1-26	92.3 %	IGH J4	97.9 %	AREQWELLYFDY	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNRVF
Clu st 3.6	identical _1	70701_12F1 6A	IGHG 1	IGHV 3-21	95.3 %	IGHD 1-26	92.3 %	IGH J4	97.9 %	AREQWELLYFDY	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNRVF

Chu st 3.7	70701_12JI 2A	IGHG 1	IGHV 3-21	98.6 %	IGHD 1-26	92.3 %	IGH J4	100.0 %	AREQWELLYFDY	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDRSNRVF
Chu st 3.8	70701_16PI 3A	IGHG 1	IGHV 3-21	99.0 %	IGHD 1-26	100.0 %	IGH J4	100.0 %	ARERWELLYFDY	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNRVVF
Chu st 3.9	70701_18M 14A	IGHG 2a	IGHV 3-21	96.3 %			IGH J4	100.0 %	AREQWVLLYFDY	λ	IGLV 6-57	97.6 %	IGL J3	100.0 %	SYDSSNRVVF
Chu st 4.1	70701_07N 16A	IGHG 2a	IGHV 3-21	97.3 %	IGHD 1-1	100.0 %	IGH J4	97.8 %	AREKWVQLYFDF	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSSNRVVF
Chu st 4.2	70701_10J0 9A	IGHG 2a	IGHV 3-21	97.6 %	IGHD 1-1	100.0 %	IGH J4	97.8 %	AREKWVQLYFDF	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNRVVF
Chu st 5.1	70701_13C0 1A	IGHG 2a	IGHV 3-21	97.0 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	97.6 %	IGL J3	100.0 %	SYDSSNRVVF
Chu st 5.2	70701_13G 21A	IGHG 1	IGHV 3-21	96.6 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	98.0 %	IGL J3	100.0 %	SYNGNNRV F
Chu st 5.3	70701_13H 15A	IGHG 1	IGHV 3-21	97.0 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDNSNRVVF
Chu st 5.4	70701_14F0 8A	IGHG 2b	IGHV 3-21	98.0 %	IGHD 1-14	100.0 %	IGH J4	97.9 %	ARERYNYFDF	λ	IGLV 6-57	98.3 %	IGL J3	100.0 %	SYDGSNRV F
Chu st 5.5	70701_14F2 1A	IGHG 1	IGHV 3-21	98.3 %	IGHD 2-2	100.0 %	IGH J4	97.9 %	AREPYNYFDY	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDNSNRVVF
Chu st 5.6	70701_14H 05A	IGHG 2b	IGHV 3-21	99.0 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNRVVF

Chu st 5.7	70701_14K 01A	IGHG 1	IGHV 3-21	98.6 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	98.3 %	IGL J3	97.1 %	SYDNNNRV F
Chu st 5.8	70701_14K 02A	IGHG 1	IGHV 3-21	98.0 %	IGHD 1-14	100.0 %	IGH J4	100.0 %	ARERYNYFDY	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDNSNRVVF
Chu st 5.9	70701_14M 09A	IGHG 1	IGHV 3-21	97.3 %	IGHD 3-16	100.0 %	IGH J4	97.8 %	ARERYNYFDS	λ	IGLV 6-57	98.0 %	IGL J3	100.0 %	SYDGNNRV F
Chu st 6.1	70701_09K 17A	IGHG 1	IGHV 3-48	97.6 %	IGHD 6-13	100.0 %	IGH J6	100.0 %	ARVVRDYGMDV	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSSNHW VF
Chu st 6.10	70701_19A 10A	IGHG 1	IGHV 3-48	98.6 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	YDSSNHW FG
Chu st 6.11	70701_19C2 2A	IGHG 1	IGHV 3-48	98.3 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNHW VF
Chu st 6.12	70701_19D 03A	IGHG 1	IGHV 3-48	98.6 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	98.3 %	IGL J3	100.0 %	SYDSSNHW VF
Chu st 6.13	70701_19O 15A	IGHG 1	IGHV 3-48	99.3 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	97.6 %	IGL J3	100.0 %	SYDSSNHW F
Chu st 6.14	70701_20O 18A	IGHG 1	IGHV 3-48	96.6 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.3 %	IGL J3	97.4 %	SYDSSNHW VF
Chu st 6.15	70701_22D 03A	IGHG 1	IGHV 3-48	95.6 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNHW VF
Chu st 6.16	70701_22O 09A	IGHG 1	IGHV 3-48	98.3 %			IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.3 %	IGL J3	100.0 %	SYDSSNHW VF

Clu st 6.17	70701_23F0 3A	IGHG 2a	IGHV 3-48	99.7 %				IGH J6	98.0 %	ARVRRNYDMDV	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNHW VF
Clu st 6.2	70701_16D 18A	IGHG 2a	IGHV 3-48	98.3 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDSSNHW VF
Clu st 6.3	70701_16E1 3A	IGHG 1	IGHV 3-48	99.0 %				IGH J6	98.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSSNHW VF
Clu st 6.4	70701_16L1 3A	IGHG 1	IGHV 3-48	99.0 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDSSNHW F
Clu st 6.5	70701_17A 23A	IGHG 1	IGHV 3-48	98.5 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSSNHW VF
Clu st 6.6	70701_17G 08A	IGHG 2a	IGHV 3-48	99.3 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	99.0 %	IGL J3	100.0 %	SYDSSNHW VF
Clu st 6.7	70701_17H 17A	IGHG 1	IGHV 3-48	96.9 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	97.3 %	IGL J3	100.0 %	SYDSSDHW VF
Clu st 6.8	70701_18A 16A	IGHG 1	IGHV 3-48	99.3 %				IGH J6	98.0 %	ARVRRNYDMDV	λ	IGLV 6-57	98.6 %	IGL J3	97.3 %	SYDSSGNWV F
Clu st 6.9	70701_18E1 9A	IGHG 1	IGHV 3-48	96.9 %				IGH J6	100.0 %	ARVRRNYGMDV	λ	IGLV 6-57	98.6 %	IGL J3	97.3 %	SYDSSNHW F
Clu st 7.1	70701_19H 6A	IGHG 1	IGHV 4-31	97.7 %				IGH J4	100.0 %	ARERKYPLQFDY	λ	IGLV 6-57	99.0 %	IGL J3	97.2 %	SYDSSNRVF
Clu st 7.2	70701_21A 06A	IGHG 1	IGHV 4-31	97.0 %				IGH J4	100.0 %	ARERKYPLQFDY	λ	IGLV 6-57	98.3 %	IGL J3	100.0 %	SFDSSNRVF

Chu st 8.1	70701_21J1 8A	IGHG 1	IGHV 4-31	98.3 %			IGH J4	100.0 %	ARERKYLlyFDY	λ	IGLV 6-57	98.6 %	IGL J3	100.0 %	SYDRSNRVF
Chu st 8.2	70701_23N 16A	IGHG 1	IGHV 4-31	98.0 %			IGH J4	100.0 %	ARERKYLlyFDY	λ	IGLV 6-57	99.0 %	IGL J3	97.2 %	SFDNSNRVf
	70701_11B1 0A	IGHG 2a	IGHV 1-69	99.7 %	IGHD 3-10	100.0 %	IGH J6	100.0 %	ARGGVGRNlyYY GMDV	λ	IGLV 1-51	99.3 %	IGL J2	100.0 %	GTWDSLS AVV
	70701_05M 03A	IGHG 2b	IGHV 2-5	98.0 %	IGHD 7-27	100.0 %	IGH J4	100.0 %	AHLAHWGPYFDY	λ	IGLV 2-23	99.7 %	IGL J2	100.0 %	CSYAGSSTL I
	70701_06A 16A	IGHG 2a	IGHV 3-21	97.3 %			IGH J4	97.7 %	ARERWALLFFEY	λ	IGLV 6-57	98.0 %	IGL J3	100.0 %	SFDRSNRVF
	70701_24B0 9A	IGHG 1	IGHV 3-21	98.0 %	IGHD 1-26	83.3 %	IGH J4	100.0 %	SRERWNLFFDY	λ	IGLV 6-57	99.7 %	IGL J3	100.0 %	SYDSSNRVf
	70701_11K 23A	IGHG 1	IGHV 3-48	97.6 %	IGHD 1-26	100.0 %	IGH J3	100.0 %	ARDGRWELLAFDI	λ	IGLV 3-25	99.3 %	IGL J3	100.0 %	QSADTSGT YRV
	70701_05D 06A	IGHG 2a	IGHV 3-7	98.0 %	IGHD 1-26	91.7 %	IGH J4	100.0 %	ARVKWDLlyD	λ	IGLV 3-1	98.2 %	IGL J2	100.0 %	QAWDSSSTH VV
	70701_19J1 0A	IGHG 1	IGHV 4-31	96.3 %			IGH J4	100.0 %	VRERKYPLlyFDY	λ	IGLV 6-57	98.3 %	IGL J3	100.0 %	SYDSSVRVf

CLAIMS

What is claimed is:

1. An anti-CD3 antibody or antibody fragment comprising a heavy chain (VH) CDR3 sequence comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59.
2. The anti-CD3 antibody or antibody fragment of claim 1, wherein the antibody or fragment comprises a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.
3. The anti-CD3 antibody or antibody fragment of claim 1, wherein the antibody or fragment comprises a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175.
4. The anti-CD3 antibody or antibody fragment of claim 1, wherein the antibody or fragment comprises a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.
5. The anti-CD3 antibody or antibody fragment of claim 1, wherein the antibody or fragment comprises:
 - a) a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least

85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175; and

b) a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

6. An anti-CD3 antibody or antibody fragment comprising a heavy chain variable region and/or a light chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

7. The anti-CD3 antibody or antibody fragment of claim 6, wherein the antibody or antibody fragment comprises a heavy chain variable region comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

8. The anti-CD3 antibody or antibody fragment of claim 6, wherein the antibody or antibody fragment comprises a light chain variable region comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11;

Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

9. The anti-CD3 antibody or antibody fragment of claim 6, wherein the antibody or antibody fragment comprises a heavy chain CDR3 sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

10. The anti-CD3 antibody or antibody fragment of claim 6, wherein the antibody or antibody fragment comprises a light chain CDR3 sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

11. A therapeutic composition comprising an anti-CD3 antibody or antibody fragment comprising a heavy chain (VH) CDR3 sequence comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59.

12. The therapeutic composition of claim 11, wherein the antibody or fragment comprises a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least

85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

13. The therapeutic composition of claim 11, wherein the antibody or fragment comprises a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175.

14. The therapeutic composition of claim 11, wherein the antibody or fragment comprises a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

15. The therapeutic composition of claim 11, wherein the therapeutic composition comprises a plurality of different CD3 antibodies, wherein each different CD3 antibody comprises a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175 and/or a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

16. The therapeutic composition of claim 11, wherein the antibody or fragment comprises a heavy chain variable region and/or a light chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust

6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

17. A method of treating a cancer patient with a therapeutic composition comprising at least one anti-CD3 antibody or fragment thereof, wherein said anti-CD3 antibody or fragment comprises:

a heavy chain (VH) CDR3 sequence comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59;

a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117;

a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175; and/or

a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

18. The method of claim 17, wherein the least one anti-CD3 antibody or fragment thereof comprises a heavy chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

19. The method of claim 17, wherein the least one anti-CD3 antibody or fragment thereof comprises a light chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

20. The method of claim 17, wherein the least one anti-CD3 antibody or fragment thereof comprises a light chain variable region and heavy chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

21. A bispecific antibody or multi-specific antibody that binds to CD3 and another antigen, wherein a targeting arm of the bispecific antibody or multi-specific antibody comprises an anti-CD3 antibody or antibody fragment comprising a heavy chain (VH) CDR3 sequence comprising an amino acid sequence selected from one of SEQ ID NOS: 2-59, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 2-59.

22. The bispecific antibody or multi-specific antibody of claim 21, wherein the anti-CD3 antibody or fragment comprises a light chain CDR3 sequence (VL or λ) comprising an amino acid sequence selected from one of SEQ ID NOS: 60-117, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 60-117.

23. The bispecific antibody or multi-specific antibody of claim 21, wherein the anti-CD3 antibody or fragment comprises a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%, at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175.

24. The bispecific antibody or multi-specific antibody claim 21, wherein the anti-CD3 antibody or fragment comprises a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

25. The bispecific antibody or multi-specific antibody of claim 21, wherein the anti-CD3 antibody or fragment comprises:

a) a heavy chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 118-175, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 118-175; and

b) a light chain variable region comprising an amino acid sequence selected from one of SEQ ID NOS: 176-233, and/or a sequence that comprises an amino acid sequence having at least 85%, at least 90% at least 95%; at least 96%, at least 97%, at least 98% or at least 99% sequence identity with any of SEQ ID NOS: 176-233.

26. A bispecific antibody or multi-specific antibody that binds to CD3 and another antigen, wherein a targeting arm of the bispecific antibody or multi-specific antibody comprises an anti-CD3 antibody or antibody fragment that comprises a heavy chain variable region and/or a light chain variable region sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust

6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

27. The bispecific antibody or multi-specific antibody of claim 26, wherein the anti-CD3 antibody or antibody fragment comprises a heavy chain variable region comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

28. The bispecific antibody or multi-specific antibody of claim 26, wherein the anti-CD3 antibody or antibody fragment comprises a light chain variable region comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

29. The bispecific antibody or multi-specific antibody of claim 26, wherein the anti-CD3 antibody or antibody fragment comprises a heavy chain CDR3 sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.

30. The bispecific antibody or multi-specific antibody of claim 26, wherein the anti-CD3 antibody or antibody fragment comprises a light chain CDR3 sequence comprising the somatic hypermutations (SMH) of clusters: Clust 1.1; Clust 1.2; Clust 2.1; Clust 2.2; Clust 2.3; Clust 2.3; Clust 2.4; Clust 3.1; Clust 3.10; Clust 3.11; Clust 3.2; Clust 3.3; Clust 3.4; Clust 3.5; Clust 3.6; Clust 3.6; Clust 3.7; Clust 3.8; Clust 3.9; Clust 4.1; Clust 4.2; Clust 5.1; Clust 5.2; Clust 5.3; Clust 5.4; Clust 5.5; Clust 5.6; Clust 5.7; Clust 5.8; Clust 5.9; Clust 6.1; Clust 6.10; Clust 6.11; Clust 6.12; Clust 6.13; Clust 6.14; Clust 6.15; Clust 6.16; Clust 6.17; Clust 6.2; Clust 6.3; Clust 6.4; Clust 6.5; Clust 6.6; Clust 6.7; Clust 6.8; Clust 6.9; Clust 7.1; Clust 7.2; Clust 8.11 and/or Clust 8.2.