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Methods of isolating T cell receptors having antigenic specificity for a cancer specific mutation

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ABSTRACT OF THE DISCLOSURE

Disclosed are methods of isolating a TCR having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising: identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence; inducing autologous APCs of the patient to present the mutated amino acid sequence; co-culturing autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence; selecting the autologous T cells; and isolating a nucleotide sequence that encodes the TCR from the selected autologous T cells, wherein the TCR has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation. Also disclosed are related methods of preparing a population of cells, populations of cells, TCRs, pharmaceutical compositions, and methods of treating or preventing cancer.

METHODS OF ISOLATING T CELL RECEPTORS HAVING ANTIGENIC SPECIFICITY
FOR A CANCER-SPECIFIC MUTATION

INCORPORATION-BY-REFERENCE OF MATERIAL SUBMITTED
ELECTRONICALLY

[0001] This application is a divisional application of Australian Patent Application No.2021200388, filed on 21 January 2021, which is a divisional of Australian Patent Application No. 2014407539 filed on 2 October 2014, and is related to International Patent Application No. PCT/US2014/058796, filed on 2 October 2014, each of which is incorporated herein by reference in its entirety. Incorporated by reference in its entirety herein is a computer-readable nucleotide/amino acid sequence listing submitted concurrently herewith and identified as follows: One 29,577 Byte ASCII (Text) file named "718291ST25.TXT," dated 15 September 2014.

BACKGROUND OF THE INVENTION

[0002] Adoptive cell therapy (ACT) using cells that have been genetically engineered to express an anti-cancer antigen T cell receptor (TCR) can produce positive clinical responses in some cancer patients. Nevertheless, obstacles to the successful use of TCR-engineered cells for the widespread treatment of cancer and other diseases remain. For example, TCRs that specifically recognize cancer antigens may be difficult to identify and/or isolate from a patient. Accordingly, there is a need for improved methods of obtaining cancer-reactive TCRs.

BRIEF SUMMARY OF THE INVENTION

[0003] An embodiment of the invention provides a method of isolating a TCR, or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising: identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence; inducing autologous antigen presenting cells (APCs) of the patient to present the mutated amino acid sequence; co-culturing autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence; selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a major histocompatibility complex (MHC) molecule expressed by the patient; and isolating a nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, from the

selected autologous T cells, wherein the TCR, or the antigen-binding portion thereof, has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation.

[0004] Another embodiment of the invention provides a method of preparing a population of cells that express a TCR, or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising: identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence; inducing autologous APCs of the patient to present the mutated amino acid sequence; co-culturing autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence; selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a MHC molecule expressed by the patient; isolating a nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, from the selected autologous T cells, wherein the TCR, or the antigen-binding portion thereof, has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation; and introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into peripheral blood mononuclear cells (PBMC) to obtain cells that express the TCR, or the antigen-binding portion thereof.

[0005] Additional embodiments of the invention provide related populations of cells, TCRs or an antigen-binding portion thereof, pharmaceutical compositions, and methods of treating or preventing cancer.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWING(S)

[0006] Figure 1A is a graph showing the number of spots per 1×10^3 ($1e3$) cells measured by interferon (IFN)- γ enzyme-linked immunosorbent spot (ELISPOT) assay after a 20 hour co-culture of 3737-TIL with OKT3 or dendritic cells (DCs) transfected with green fluorescent protein (GFP) RNA, or the indicated tandem mini-gene (TMG) construct. “>” denotes greater than 500 spots per 1×10^3 cells. Mock-transfected cells were treated with transfection reagent only without addition of nucleic acid.

[0007] Figure 1B is a graph showing the percentage of CD4+ 3737-TIL that were OX40+ following co-culture with OKT3 or DCs transfected with GFP RNA, TMG-1, or the indicated

wild type (wt) gene *ALK*, *CD93*, *ERBB2IP*, *FCER1A*, *GRXCRI*, *KIF9*, *NAGS*, *NLRP2*, or *RAC3*. Mock-transfected cells were treated with transfection reagent only without addition of nucleic acid.

[0008] Figures 2A-2C are graphs showing the number of spots per 1×10^3 ($1e3$) cells measured by IFN- γ ELISPOT assay at 20 hours for 3737-TIL (A), DMF5 T cells (B), or T4 T cells (C) that were co-cultured with DCs transfected with TMG-1 (A) or 624-CIITA cells (B) and (C) that had been pre-incubated with nothing, or the indicated HLA-blocking antibodies (against MHC-I, MHC-II, HLA-DP, HLA-DQ, or HLA-DR) (A-C).

[0009] Figure 2D is a graph showing the number of spots per 1×10^3 ($1e3$) cells measured by IFN- γ ELISPOT assay at 20 hours for 3737-TIL co-cultured with autologous DQ-0301/-0601 B cells (grey bars) or allogeneic EBV-B cells partially matched at the HLA-DQ 05/0601 locus (black bars) or the HLA-DQ-0201/0301 locus (unshaded bars) that had been pulsed overnight with DMSO, mutated (mut) ALK or mut ERBB2IP 25-AA long peptides. ETGHLENGNKYPNLE (SEQ ID NO: 53);

[0010] Figure 2E is a graph showing the number of spots per 1×10^3 ($1e3$) cells measured by IFN- γ ELISPOT assay at 20 hours for 3737-TIL co-cultured with autologous B cells that had been pulsed overnight with the mut ERBB2IP 25-AA peptide TSFLSINSKEETGHLENGNKYPNLE (SEQ ID NO: 73), or the indicated truncated mut ERBB2IP peptides FLSINSKEETGHLENGNKYPNLE (SEQ ID NO: 30), SINSKEETGHLENGNKYPNLE (SEQ ID NO: 31), NSKEETGHLENGNKYPNLE (SEQ ID NO: 32), KEETGHLENGNKYPNLE (SEQ ID NO: 33), ETGHLENGNKYPNLE (SEQ ID NO: 53), TSFLSINSKEETGHL (SEQ ID NO: 34), TSFLSINSKEETGHLEN (SEQ ID NO: 35), TSFLSINSKEETGHLENGN (SEQ ID NO: 36), TSFLSINSKEETGHLENGNKY (SEQ ID NO: 37), or TSFLSINSKEETGHLENGNKYPN (SEQ ID NO: 38).

[0011] Figure 3A is a graph showing the percentage of various TCR V β clonotypes in 3737-TIL, measured by flow cytometry gated on live CD4+ (shaded) or CD8+ (unshaded) T cells.

[0012] Figure 3B is a graph showing the IFN- γ levels (pg/ml) detected in patient 3737 serum samples measured at the indicated number of days pre- and post-adoptive cell transfer of 3737-TIL on Day 0 (indicated by arrow). Error bars are standard error of the mean (SEM).

[0013] Figure 3C is a graph showing the total tumor burden (circles) (measured as % of pre-treatment baseline) or tumor burden in the lung (triangles) or liver (squares) at the indicated number of months relative to cell transfer on day 0 (indicated by arrow).

[0014] Figure 3D is a graph showing the percentage of various TCR V β clonotypes in CD4⁺ V β 22⁻ OX40⁺ 3737-TIL, as measured by flow cytometry.

[0015] Figures 4A and 4B are graphs showing the frequency of the two ERBB2IP-mutation-specific TCR β -CDR3 clonotypes V β 22⁺ (A) and V β 5.2⁺ (B) in the blood (circles) of patient 3737 at various times pre- and post-adoptive cell transfer with 3737-TIL, a tumor before cell transfer (diamonds), and various tumors after cell transfer (Tu-1-Post (squares), Tu-2-Post (\blacktriangle), and Tu-3-Post (\blacktriangledown)). Shaded bars indicate the frequency of the two ERBB2IP-mutation-specific TCR β -CDR3 clonotypes V β 22⁺ (A) and V β 5.2⁺ (B) in the transferred cells (3737-TIL). “X” indicates “Not detected.”

[0016] Figure 4C is a graph showing *ERBB2IP* expression relative to *ACTB* in 3737-TIL (T cells) and various tumors pre (Tu-Pre) and post (Tu-1-post, Tu-2-post, and Tu-3-post) adoptive cell transfer.

[0017] Figure 4D is a graph showing the total tumor burden (circles) (measured as % of pre-treatment baseline) or tumor burden in the lung (triangles) or liver (squares) at the indicated number of months relative to cell transfer (indicated by arrows).

[0018] Figure 5A is a schematic of an example of tandem minigene (TMG) construct, which encoded polypeptides containing 6 identified mutated amino acid residues flanked on their N- and C- termini, 12 amino acids on both sides. The mutated KIF2C sequence is DSSLQARLFPGLTIKIQRSNGLIHS (SEQ ID NO: 57).

[0019] Figure 5B is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2359 T cells co-cultured overnight with autologous melanocytes or COS-7 cells co-transfected with HLA-A*0205 and TMG construct RJ-1 (structure shown in Fig. 9A), RJ-2, RJ-3, RJ-4, RJ-5, RJ-6, RJ-7, RJ-8, RJ-9, RJ-10, RJ-11, RJ-12, or an empty vector.

[0020] Figure 5C is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2359 co-cultured with COS-7 cells transfected with HLA-A*0205 and an RJ-1 variant in which the gene indicated “wt” in the table was converted back to the WT sequence. The *KIF2C* WT sequence is DSSLQARLFPGLAIKIQRSNGLIHS (SEQ ID NO: 65).

[0021] Figure 5D is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2359 co-cultured with COS-7 cells transfected with an empty vector, *KIF2C* WT, or mutated *KIF2C* cDNA construct, together with HLA cDNA construct (identifying each shaded bar from left to right): HLA-A*0101 (unshaded bars), HLA-A*0201 (grey bars), or HLA-A*0205 (black bars).

[0022] Figure 5E is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2359 T cells co-cultured overnight with HEK293 cells stably expressing HLA-A*0205 that were pulsed with various concentrations (μ M) of KIF2C₁₀₋₁₉ WT (RLFPGLAIKI; SEQ ID NO: 58) (bottom line in graph) or mutated KIF2C₁₀₋₁₉ (RLFPGLTIKI; SEQ ID NO: 59) (top line in graph).

[0023] Figure 6A is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2591 T cells co-cultured with autologous melanocytes or HEK293 cells stably expressing HLA-C*0701 transfected with an empty vector or a TMG construct selected from the group consisting of DW-1 to DW-37.

[0024] Figure 6B is a schematic showing the structure of TMG construct DW-6. The mutated POLA2 sequence is TIEGTRSSGSHFVFPVPSLRDVHHE (SEQ ID NO: 64).

[0025] Figure 6C is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2591 co-cultured with COS-7 cells transfected with HLA-C*0701 and a DW-6 variant in which the gene indicated "wt" in the table was converted back to the WT sequence. The *POLA2* WT sequence is TIEGTRSSGSHLVFPVPSLRDVHHE (SEQ ID NO: 66).

[0026] Figure 6D is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2591 co-cultured with COS-7 cells transfected with an empty vector, *POLA2* WT, or mutated *POLA2* cDNA construct, together with HLA cDNA construct (identifying each bar from left to right): HLA-C*0401 (unshaded bars), HLA-C*0701 (grey bars), or HLA-C*0702 (black bars).

[0027] Figure 6E is a graph showing the level of IFN- γ (pg/mL) secreted by TIL 2591 T cells co-cultured overnight with HEK293 cells stably expressing HLA-C*0701 that were pulsed with various concentrations (μ M) of POLA2₄₁₃₋₄₂₂ WT (TRSSGSHLVF; SEQ ID NO: 67) (bottom line in graph) or mutated POLA2₄₁₃₋₄₂₂ (TRSSGSHFVF; SEQ ID NO: 68) (top line in graph).

[0028] Figures 7A-7F are computerized tomography (CT) scans of the lungs of Patient 3737 taken prior to (A-C) and six months after (D-F) the second administration of mutation-reactive cells. The arrows point to cancerous lesions.

DETAILED DESCRIPTION OF THE INVENTION

[0029] An embodiment of the invention provides a method of isolating a TCR, or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation. The invention provides many advantages.

For example, the inventive methods may rapidly assess a large number of mutations restricted by all of the patient's MHC molecules at one time, which may identify the full repertoire of the patient's mutation-reactive T cells. Additionally, by distinguishing immunogenic cancer mutations from (a) silent cancer-specific mutations (which do not encode a mutated amino acid sequence) and (b) cancer-specific mutations that encode a non-immunogenic amino acid sequence, the inventive methods may identify one or more cancer-specific, mutated amino acid sequences that may be targeted by a TCR, or an antigen-binding portion thereof. In addition, the invention may provide TCRs, and antigen-binding portions thereof, having antigenic specificity for mutated amino acid sequences encoded by cancer-specific mutations that are unique to the patient, thereby providing "personalized" TCRs, and antigen-binding portions thereof, that may be useful for treating or preventing the patient's cancer. The inventive methods may also avoid the technical biases inherent in traditional methods of identifying cancer antigens such as, for example, those using cDNA libraries, and may also be less time-consuming and laborious than those methods. For example, the inventive methods may select mutation-reactive T cells without co-culturing the T cells with tumor cell lines, which may be difficult to generate, particularly for e.g., epithelial cancers. Without being bound to a particular theory or mechanism, it is believed that the inventive methods may identify and isolate TCRs, or antigen-binding portions thereof, that target the destruction of cancer cells while minimizing or eliminating the destruction of normal, non-cancerous cells, thereby reducing or eliminating toxicity. Accordingly, the invention may also provide TCRs, or antigen-binding portions thereof, that successfully treat or prevent cancer such as, for example, cancers that do not respond to other types of treatment such as, for example, chemotherapy alone, surgery, or radiation.

[0030] The method may comprise identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence. The cancer cell may be obtained from any bodily sample derived from a patient which contains or is expected to contain tumor or cancer cells. The bodily sample may be any tissue sample such as blood, a tissue sample obtained from the primary tumor or from tumor metastases, or any other sample containing tumor or cancer cells. The nucleic acid of the cancer cell may be DNA or RNA.

[0031] In order to identify cancer-specific mutations, the method may further comprise sequencing nucleic acid such as DNA or RNA of normal, noncancerous cells and comparing

the sequence of the cancer cell with the sequence of the normal, noncancerous cell. The normal, noncancerous cell may be obtained from the patient or a different individual.

[0032] The cancer-specific mutation may be any mutation in any gene which encodes a mutated amino acid sequence (also referred to as a “non-silent mutation”) and which is expressed in a cancer cell but not in a normal, noncancerous cell. Non-limiting examples of cancer-specific mutations that may be identified in the inventive methods include missense, nonsense, insertion, deletion, duplication, frameshift, and repeat expansion mutations. In an embodiment of the invention, the method comprises identifying at least one gene containing a cancer-specific mutation which encodes a mutated amino acid sequence. However, the number of genes containing such a cancer-specific mutation that may be identified using the inventive methods is not limited and may include more than one gene (for example, about 2, about 3, about 4, about 5, about 10, about 11, about 12, about 13, about 14, about 15, about 20, about 25, about 30, about 40, about 50, about 60, about 70, about 80, about 90, about 100, about 150, about 200, about 400, about 600, about 800, about 1000, about 1500, about 2000 or more, or a range defined by any two of the foregoing values). Likewise, in an embodiment of the invention, the method comprises identifying at least one cancer-specific mutation which encodes a mutated amino acid sequence. However, the number of such cancer-specific mutations that may be identified using the inventive methods is not limited and may include more than one cancer-specific mutation (for example, about 2, about 3, about 4, about 5, about 10, about 11, about 12, about 13, about 14, about 15, about 20, about 25, about 30, about 40, about 50, about 60, about 70, about 80, about 90, about 100, about 150, about 200, about 400, about 600, about 800, about 1000, about 1500, about 2000 or more, or a range defined by any two of the foregoing values). In an embodiment in which more than one cancer-specific mutation is identified, the cancer-specific mutations may be located in the same gene or in different genes.

[0033] In an embodiment, identifying one or more genes in the nucleic acid of a cancer cell comprises sequencing the whole exome, the whole genome, or the whole transcriptome of the cancer cell. Sequencing may be carried out in any suitable manner known in the art. Examples of sequencing techniques that may be useful in the inventive methods include Next Generation Sequencing (NGS) (also referred to as “massively parallel sequencing technology”) or Third Generation Sequencing. NGS refers to non-Sanger-based high-throughput DNA sequencing technologies. With NGS, millions or billions of DNA strands may be sequenced in parallel, yielding substantially more throughput and minimizing the

need for the fragment-cloning methods that are often used in Sanger sequencing of genomes. In NGS, nucleic acid templates may be randomly read in parallel along the entire genome by breaking the entire genome into small pieces. NGS may, advantageously, provide nucleic acid sequence information of a whole genome, exome, or transcriptome in very short time periods, e.g., within about 1 to about 2 weeks, preferably within about 1 to about 7 days, or most preferably, within less than about 24 hours. Multiple NGS platforms which are commercially available or which are described in the literature can be used in the context of the inventive methods, e.g., those described in Zhang et al., *J. Genet. Genomics*, 38(3): 95-109 (2011) and Voelkerding et al., *Clinical Chemistry*, 55: 641-658 (2009).

[0034] Non-limiting examples of NGS technologies and platforms include sequencing-by-synthesis (also known as “pyrosequencing”) (as implemented, e.g., using the GS-FLX 454 Genome Sequencer, 454 Life Sciences (Branford, CT), ILLUMINA SOLEXA Genome Analyzer (Illumina Inc., San Diego, CA), or the ILLUMINA HISEQ 2000 Genome Analyzer (Illumina), or as described in, e.g., Ronaghi et al., *Science*, 281(5375): 363-365 (1998)), sequencing-by-ligation (as implemented, e.g., using the SOLID platform (Life Technologies Corporation, Carlsbad, CA) or the POLONATOR G.007 platform (Dover Systems, Salem, NH)), single-molecule sequencing (as implemented, e.g., using the PACBIO RS system (Pacific Biosciences (Menlo Park, CA) or the HELISCOPE platform (Helicos Biosciences (Cambridge, MA)), nano-technology for single-molecule sequencing (as implemented, e.g., using the GRIDON platform of Oxford Nanopore Technologies (Oxford, UK), the hybridization-assisted nano-pore sequencing (HANS) platforms developed by Nabsys (Providence, RI), and the ligase-based DNA sequencing platform with DNA nanoball (DNB) technology referred to as probe-anchor ligation (cPAL)), electron microscopy-based technology for single-molecule sequencing, and ion semiconductor sequencing.

[0035] The method may comprise inducing autologous antigen presenting cells (APCs) of the patient to present the mutated amino acid sequence. The APCs may include any cells which present peptide fragments of proteins in association with major histocompatibility complex (MHC) molecules on their cell surface. The APCs may include, for example, any one or more of macrophages, DCs, langerhans cells, B-lymphocytes, and T-cells. Preferably, the APCs are DCs. By using autologous APCs from the patient, the inventive methods may, advantageously, identify TCRs, and antigen-binding portions thereof, that have antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation that is presented in the context of an MHC molecule expressed by the patient. The MHC molecule

can be any MHC molecule expressed by the patient including, but not limited to, MHC Class I, MHC Class II, HLA-A, HLA-B, HLA-C, HLA-DM, HLA-DO, HLA-DP, HLA-DQ, and HLA-DR molecules. The inventive methods may, advantageously, identify mutated amino acid sequences presented in the context of any MHC molecule expressed by the patient without using, for example, epitope prediction algorithms to identify MHC molecules or mutated amino acid sequences, which may be useful only for a select few MHC class I alleles and may be constrained by the limited availability of reagents to select mutation-reactive T cells (e.g., an incomplete set of MHC tetramers). Accordingly, in an embodiment of the invention, the inventive methods advantageously identify mutated amino acid sequences presented in the context of any MHC molecule expressed by the patient and are not limited to any particular MHC molecule. Preferably, the autologous APCs are antigen-negative autologous APCs.

[0036] Inducing autologous APCs of the patient to present the mutated amino acid sequence may be carried out using any suitable method known in the art. In an embodiment of the invention, inducing autologous APCs of the patient to present the mutated amino acid sequence comprises pulsing the autologous APCs with peptides comprising the mutated amino acid sequence or a pool of peptides, each peptide in the pool comprising a different mutated amino acid sequence. Each of the mutated amino acid sequences in the pool may be encoded by a gene containing a cancer specific mutation. In this regard, the autologous APCs may be cultured with a peptide or a pool of peptides comprising the mutated amino acid sequence in a manner such that the APCs internalize the peptide(s) and display the mutated amino acid sequence(s), bound to an MHC molecule, on the cell membrane. In an embodiment in which more than one gene is identified, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence, the method may comprise pulsing the autologous APCs with a pool of peptides, each peptide in the pool comprising a different mutated amino acid sequence. Methods of pulsing APCs are known in the art and are described in, e.g., Solheim (Ed.), *Antigen Processing and Presentation Protocols (Methods in Molecular Biology)*, Human Press, (2010). The peptide(s) used to pulse the APCs may include the mutated amino acid(s) encoded by the cancer-specific mutation. The peptide(s) may further comprise any suitable number of contiguous amino acids from the endogenous protein encoded by the identified gene on each of the carboxyl side and the amino side of the mutated amino acid(s). The number of contiguous amino acids from the endogenous protein flanking each side of the mutation is not limited and may be, for

example, about 4, about 5, about 6, about 7, about 8, about 9, about 10, about 11, about 12, about 13, about 14, about 15, about 16, about 17, about 18, about 19, about 20, or a range defined by any two of the foregoing values. Preferably, the peptide(s) comprise(s) about 12 contiguous amino acids from the endogenous protein on each side of the mutated amino acid(s).

[0037] In an embodiment of the invention, inducing autologous APCs of the patient to present the mutated amino acid sequence comprises introducing a nucleotide sequence encoding the mutated amino acid sequence into the APCs. The nucleotide sequence is introduced into the APCs so that the APCs express and display the mutated amino acid sequence, bound to an MHC molecule, on the cell membrane. The nucleotide sequence encoding the mutated amino acid may be RNA or DNA. Introducing a nucleotide sequence into APCs may be carried out in any of a variety of different ways known in the art as described in, e.g., Solheim et al. *supra*. Non-limiting examples of techniques that are useful for introducing a nucleotide sequence into APCs include transformation, transduction, transfection, and electroporation. In an embodiment in which more than one gene is identified, the method may comprise preparing more than one nucleotide sequence, each encoding a mutated amino acid sequence encoded by a different gene, and introducing each nucleotide sequence into a different population of autologous APCs. In this regard, multiple populations of autologous APCs, each population expressing and displaying a different mutated amino acid sequence, may be obtained.

[0038] In an embodiment in which more than one gene is identified, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence, the method may comprise introducing a nucleotide sequence encoding the more than one gene. In this regard, in an embodiment of the invention, the nucleotide sequence introduced into the autologous APCs is a TMG construct, each minigene comprising a different gene, each gene including a cancer-specific mutation that encodes a mutated amino acid sequence. Each minigene may encode one mutation identified by the inventive methods flanked on each side of the mutation by any suitable number of contiguous amino acids from the endogenous protein encoded by the identified gene, as described herein with respect to other aspects of the invention. The number of minigenes in the construct is not limited and may include for example, about 5, about 10, about 11, about 12, about 13, about 14, about 15, about 20, about 25, or more, or a range defined by any two of the foregoing values. The APCs express the mutated amino acid sequences encoded by the TMG construct and display the mutated amino acid sequences,

bound to an MHC molecule, on the cell membranes. In an embodiment, the method may comprise preparing more than one TMG construct, each construct encoding a different set of mutated amino acid sequences encoded by different genes, and introducing each TMG construct into a different population of autologous APCs. In this regard, multiple populations of autologous APCs, each population expressing and displaying mutated amino acid sequences encoded by different TMG constructs, may be obtained.

[0039] The method may comprise culturing autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence. The T cells can be obtained from numerous sources in the patient, including but not limited to tumor, blood, bone marrow, lymph node, the thymus, or other tissues or fluids. The T cells can include any type of T cell and can be of any developmental stage, including but not limited to, CD4+/CD8+ double positive T cells, CD4+ helper T cells, e.g., Th1 and Th2 cells, CD8+ T cells (e.g., cytotoxic T cells), tumor infiltrating cells (e.g., tumor infiltrating lymphocytes (TIL)), peripheral blood T cells, memory T cells, naïve T cells, and the like. The T cells may be CD8+ T cells, CD4+ T cells, or both CD4+ and CD8+ T cells. The method may comprise co-culturing the autologous T cells and autologous APCs so that the T cells encounter the mutated amino acid sequence presented by the APCs in such a manner that the autologous T cells specifically bind to and immunologically recognize a mutated amino acid sequence presented by the APCs. In an embodiment of the invention, the autologous T cells are co-cultured in direct contact with the autologous APCs.

[0040] The method may comprise selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a MHC molecule expressed by the patient. The phrase “antigenic specificity,” as used herein, means that a TCR, or the antigen-binding portion thereof, expressed by the autologous T cells can specifically bind to and immunologically recognize the mutated amino acid sequence encoded by the cancer-specific mutation. The selecting may comprise identifying the T cells that have antigenic specificity for the mutated amino acid sequence and separating them from T cells that do not have antigenic specificity for the mutated amino acid sequence. Selecting the autologous T cells having antigenic specificity for the mutated amino acid sequence may be carried out in any suitable manner. In an embodiment of the invention, the method comprises expanding the numbers of autologous T cells, e.g., by co-culturing with a T cell growth factor, such as interleukin (IL)-2 or IL-15, or as described herein with respect to other

aspects of the invention, prior to selecting the autologous T cells. In an embodiment of the invention, the method does not comprise expanding the numbers of autologous T cells with a T cell growth factor, such as IL-2 or IL-15 prior to selecting the autologous T cells.

[0041] For example, upon co-culture of the autologous T cells with the APCs that present the mutated amino acid sequence, T cells having antigenic specificity for the mutated amino acid sequence may express any one or more of a variety of T cell activation markers which may be used to identify those T cells having antigenic specificity for the mutated amino acid sequence. Such T cell activation markers may include, but are not limited to, programmed cell death 1 (PD-1), lymphocyte-activation gene 3 (LAG-3), T cell immunoglobulin and mucin domain 3 (TIM-3), 4-1BB, OX40, and CD107a. Accordingly, in an embodiment of the invention, selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selecting the T cells that express any one or more of PD-1, LAG-3, TIM-3, 4-1BB, OX40, and CD107a. Cells expressing one or more T cell activation markers may be sorted on the basis of expression of the marker using any of a variety of techniques known in the art such as, for example, fluorescence-activated cell sorting (FACS) or magnetic-activated cell sorting (MACS) as described in, e.g., Turcotte et al., *Clin. Cancer Res.*, 20(2): 331-43 (2013) and Gros et al., *J. Clin. Invest.*, 124(5): 2246-59 (2014).

[0042] In another embodiment of the invention, selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selecting the T cells (i) that secrete a greater amount of one or more cytokines upon co-culture with APCs that present the mutated amino acid sequence as compared to the amount of the one or more cytokines secreted by a negative control or (ii) in which at least twice as many of the numbers of T cells secrete one or more cytokines upon co-culture with APCs that present the mutated amino acid sequence as compared to the numbers of negative control T cells that secrete the one or more cytokines. The one or more cytokines may comprise any cytokine the secretion of which by a T cell is characteristic of T cell activation (e.g., a TCR expressed by the T cells specifically binding to and immunologically recognizing the mutated amino acid sequence). Non-limiting examples of cytokines, the secretion of which is characteristic of T cell activation, include IFN- γ , IL-2, and tumor necrosis factor alpha (TNF- α), granulocyte/monocyte colony stimulating factor (GM-CSF), IL-4, IL-5, IL-9, IL-10, IL-17, and IL-22.

[0043] For example, a TCR, or an antigen-binding portion thereof, or a T cell expressing the TCR, or the antigen-binding portion thereof, may be considered to have “antigenic

specificity” for the mutated amino acid sequence if the T cells, or T cells expressing the TCR, or the antigen-binding portion thereof, secrete at least twice as much IFN- γ upon co-culture with (a) antigen-negative APCs pulsed with a concentration of a peptide comprising the mutated amino acid sequence (e.g., about 0.05 ng/mL to about 10 μ g/mL, e.g., 0.05 ng/mL, 0.1 ng/mL, 0.5 ng/mL, 1 ng/mL, 5 ng/mL, 100 ng/mL, 1 μ g/mL, 5 μ g/mL, or 10 μ g/mL) or (b) APCs into which a nucleotide sequence encoding the mutated amino acid sequence has been introduced as compared to the amount of IFN- γ secreted by a negative control. The negative control may be, for example, (i) T cells expressing the TCR, or the antigen-binding portion thereof, co-cultured with (a) antigen-negative APCs pulsed with the same concentration of an irrelevant peptide (e.g., the wild-type amino acid sequence, or some other peptide with a different sequence from the mutated amino acid sequence) or (b) APCs into which a nucleotide sequence encoding an irrelevant peptide sequence has been introduced, or (ii) untransduced T cells (e.g., derived from PBMC, which do not express the TCR, or antigen binding portion thereof) co-cultured with (a) antigen-negative APCs pulsed with the same concentration of a peptide comprising the mutated amino acid sequence or (b) APCs into which a nucleotide sequence encoding the mutated amino acid sequence has been introduced. A TCR, or an antigen-binding portion thereof, or a T cell expressing the TCR, or the antigen-binding portion thereof, may also have “antigenic specificity” for the mutated amino acid sequence if T cells, or T cells expressing the TCR, or the antigen-binding portion thereof, secrete a greater amount of IFN- γ upon co-culture with antigen-negative APCs pulsed with higher concentrations of a peptide comprising the mutated amino acid sequence as compared to a negative control, for example, any of the negative controls described above. IFN- γ secretion may be measured by methods known in the art such as, for example, enzyme-linked immunosorbent assay (ELISA).

[0044] Alternatively or additionally, a TCR, or an antigen-binding portion thereof, or a T cell expressing the TCR, or the antigen-binding portion thereof, may be considered to have “antigenic specificity” for the mutated amino acid sequence if at least twice as many of the numbers of T cells, or T cells expressing the TCR, or the antigen-binding portion thereof, secrete IFN- γ upon co-culture with (a) antigen-negative APCs pulsed with a concentration of a peptide comprising the mutated amino acid sequence or (b) APCs into which a nucleotide sequence encoding the mutated amino acid sequence has been introduced as compared to the numbers of negative control T cells that secrete IFN- γ . The concentration of peptide and the negative control may be as described herein with respect to other aspects of the invention.

The numbers of cells secreting IFN- γ may be measured by methods known in the art such as, for example, ELISPOT.

[0045] While T cells having antigenic specificity for the mutated amino acid sequence may both (1) express any one or more T cells activation markers described herein and (2) secrete a greater amount of one or more cytokines as described herein, in an embodiment of the invention, T cells having antigenic specificity for the mutated amino acid sequence may express any one or more T cell activation markers without secreting a greater amount of one or more cytokines or may secrete a greater amount of one or more cytokines without expressing any one or more T cell activation markers.

[0046] In another embodiment of the invention, selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selectively growing the autologous T cells that have antigenic specificity for the mutated amino acid sequence. In this regard, the method may comprise co-culturing the autologous T cells with autologous APCs in such a manner as to favor the growth of the T cells that have antigenic specificity for the mutated amino acid sequence over the T cells that do not have antigenic specificity for the mutated amino acid sequence. Accordingly, a population of T cells is provided that has a higher proportion of T cells that have antigenic specificity for the mutated amino acid sequence as compared to T cells that do not have antigenic specificity for the mutated amino acid sequence.

[0047] In an embodiment of the invention, the method further comprises obtaining multiple fragments of a tumor from the patient, separately co-culturing autologous T cells from each of the multiple fragments with the autologous APCs that present the mutated amino acid sequence as described herein with respect to other aspects of the invention, and separately assessing the T cells from each of the multiple fragments for antigenic specificity for the mutated amino acid sequence, as described herein with respect to other aspects of the invention.

[0048] In an embodiment of the invention in which T cells are co-cultured with autologous APCs expressing multiple mutated amino acid sequences (e.g., multiple mutated amino acid sequences encoded by a TMG construct or multiple mutated amino acid sequences in a pool of peptides pulsed onto autologous APCs), selecting the autologous T cells may further comprise separately assessing autologous T cells for antigenic specificity for each of the multiple mutated amino acid sequences. For example, the inventive method may further comprise separately inducing autologous APCs of the patient to present each

mutated amino acid sequence encoded by the construct (or included in the pool), as described herein with respect to other aspects of the invention (for example, by providing separate APC populations, each presenting a different mutated amino acid sequence encoded by the construct (or included in the pool)). The method may further comprise separately co-culturing autologous T cells of the patient with the different populations of autologous APCs that present each mutated amino acid sequence, as described herein with respect to other aspects of the invention. The method may further comprise separately selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a MHC molecule expressed by the patient, as described herein with respect to other aspects of the invention. In this regard, the method may comprise determining which mutated amino acid sequence encoded by a TMG construct that encodes multiple mutated amino acid sequences (or included in the pool) are immunologically recognized by the autologous T cells (e.g., by process of elimination).

[0049] The method may further comprise isolating a nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, from the selected autologous T cells, wherein the TCR, or the antigen-binding portion thereof, has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation. In an embodiment of the invention, prior to isolating the nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, the numbers selected autologous T cells that have antigenic specificity for the mutated amino acid sequence may be expanded. Expansion of the numbers of T cells can be accomplished by any of a number of methods as are known in the art as described in, for example, U.S. Patent 8,034,334; U.S. Patent 8,383,099; U.S. Patent Application Publication No. 2012/0244133; Dudley et al., *J. Immunother.*, 26:332-42 (2003); and Riddell et al., *J. Immunol. Methods*, 128:189-201 (1990). In an embodiment, expansion of the numbers of T cells is carried out by culturing the T cells with OKT3 antibody, IL-2, and feeder PBMC (e.g., irradiated allogeneic PBMC). In another embodiment of the invention, the numbers of selected autologous T cells that have antigenic specificity for the mutated amino acid sequence are not expanded prior to isolating the nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof.

[0050] The “the antigen-binding portion” of the TCR, as used herein, refers to any portion comprising contiguous amino acids of the TCR of which it is a part, provided that the antigen-binding portion specifically binds to the mutated amino acid sequence encoded by the

gene identified as described herein with respect to other aspects of the invention. The term “antigen-binding portion” refers to any part or fragment of the TCR of the invention, which part or fragment retains the biological activity of the TCR of which it is a part (the parent TCR). Antigen-binding portions encompass, for example, those parts of a TCR that retain the ability to specifically bind to the mutated amino acid sequence, or detect, treat, or prevent cancer, to a similar extent, the same extent, or to a higher extent, as compared to the parent TCR. In reference to the parent TCR, the functional portion can comprise, for instance, about 10%, 25%, 30%, 50%, 68%, 80%, 90%, 95%, or more, of the parent TCR.

[0051] The antigen-binding portion can comprise an antigen-binding portion of either or both of the α and β chains of the TCR of the invention, such as a portion comprising one or more of the complementarity determining region (CDR)1, CDR2, and CDR3 of the variable region(s) of the α chain and/or β chain of the TCR of the invention. In an embodiment of the invention, the antigen-binding portion can comprise the amino acid sequence of the CDR1 of the α chain (CDR1 α), the CDR2 of the α chain (CDR2 α), the CDR3 of the α chain (CDR3 α), the CDR1 of the β chain (CDR1 β), the CDR2 of the β chain (CDR2 β), the CDR3 of the β chain (CDR3 β), or any combination thereof. Preferably, the antigen-binding portion comprises the amino acid sequences of CDR1 α , CDR2 α , and CDR3 α ; the amino acid sequences of CDR1 β , CDR2 β , and CDR3 β ; or the amino acid sequences of all of CDR1 α , CDR2 α , CDR3 α , CDR1 β , CDR2 β , and CDR3 β of the inventive TCR.

[0052] In an embodiment of the invention, the antigen-binding portion can comprise, for instance, the variable region of the inventive TCR comprising a combination of the CDR regions set forth above. In this regard, the antigen-binding portion can comprise the amino acid sequence of the variable region of the α chain (V α), the amino acid sequence of the variable region of the β chain (V β), or the amino acid sequences of both of the V α and V β of the inventive TCR.

[0053] In an embodiment of the invention, the antigen-binding portion may comprise a combination of a variable region and a constant region. In this regard, the antigen-binding portion can comprise the entire length of the α or β chain, or both of the α and β chains, of the inventive TCR.

[0054] Isolating the nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, from the selected autologous T cells may be carried out in any suitable manner known in the art. For example, the method may comprise isolating RNA from the autologous T cells and sequencing the TCR, or the antigen-binding portion thereof, using

established molecular cloning techniques and reagents such as, for example, 5' Rapid Amplification of cDNA Ends (RACE) polymerase chain reaction (PCR) using TCR- α and - β chain constant primers.

[0055] In an embodiment of the invention, the method may comprise cloning the nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, into a recombinant expression vector using established molecular cloning techniques as described in, e.g., Green et al. (Eds.), *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratory Press; 4th Ed. (2012). For purposes herein, the term "recombinant expression vector" means a genetically-modified oligonucleotide or polynucleotide construct that permits the expression of an mRNA, protein, polypeptide, or peptide by a host cell, when the construct comprises a nucleotide sequence encoding the mRNA, protein, polypeptide, or peptide, and the vector is contacted with the cell under conditions sufficient to have the mRNA, protein, polypeptide, or peptide expressed within the cell. The vectors of the invention are not naturally-occurring as a whole. However, parts of the vectors can be naturally-occurring. The recombinant expression vectors can comprise any type of nucleotides, including, but not limited to DNA and RNA, which can be single-stranded or double-stranded, synthesized or obtained in part from natural sources, and which can contain natural, non-natural or altered nucleotides. The recombinant expression vectors can comprise naturally-occurring, non-naturally-occurring internucleotide linkages, or both types of linkages. Preferably, the non-naturally occurring or altered nucleotides or internucleotide linkages does not hinder the transcription or replication of the vector.

[0056] The recombinant expression vector of the invention can be any suitable recombinant expression vector, and can be used to transform or transfect any suitable host cell. Suitable vectors include those designed for propagation and expansion or for expression or both, such as plasmids and viruses. The vector can be selected from the group consisting of transposon/transposase, the pUC series (Fermentas Life Sciences), the pBluescript series (Stratagene, LaJolla, CA), the pET series (Novagen, Madison, WI), the pGEX series (Pharmacia Biotech, Uppsala, Sweden), and the pEX series (Clontech, Palo Alto, CA). Bacteriophage vectors, such as λ GT10, λ GT11, λ ZapII (Stratagene), λ EMBL4, and λ NM1149, also can be used. Examples of plant expression vectors include pBI01, pBI101.2, pBI101.3, pBI121 and pBIN19 (Clontech). Examples of animal expression vectors include pEUK-C1, pMAM and pMAMneo (Clontech). Preferably, the recombinant expression vector is a viral vector, e.g., a retroviral vector.

[0057] The TCR, or the antigen-binding portion thereof, isolated by the inventive methods may be useful for preparing cells for adoptive cell therapies. In this regard, an embodiment of the invention provides a method of preparing a population of cells that express a TCR, or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising isolating a TCR, or an antigen-binding portion thereof, as described herein with respect to other aspects of the invention, and introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into PBMC to obtain cells that express the TCR, or the antigen-binding portion thereof.

[0058] Introducing the nucleotide sequence (e.g., a recombinant expression vector) encoding the isolated TCR, or the antigen-binding portion thereof, into PBMC may be carried out in any of a variety of different ways known in the art as described in, e.g., Green et al. *supra*. Non-limiting examples of techniques that are useful for introducing a nucleotide sequence into PBMC include transformation, transduction, transfection, and electroporation.

[0059] In an embodiment of the invention, the method comprises introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into PBMC that are autologous to the patient. In this regard, the TCRs, or the antigen-binding portions thereof, identified and isolated by the inventive methods may be personalized to each patient. However, in another embodiment, the inventive methods may identify and isolate TCRs, or the antigen-binding portions thereof, that have antigenic specificity against a mutated amino acid sequence that is encoded by a recurrent (also referred to as “hot-spot”) cancer-specific mutation. In this regard, the method may comprise introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into PBMC that are allogeneic to the patient. For example, the method may comprise introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into the PBMC of another patient whose tumors express the same mutation in the context of the same MHC molecule.

[0060] In an embodiment of the invention, the PBMC include T cells. The T cells may be any type of T cell, for example, any of those described herein with respect to other aspects of the invention. Without being bound to a particular theory or mechanism, it is believed that less differentiated, “younger” T cells may be associated with any one or more of greater *in vivo* persistence, proliferation, and antitumor activity as compared to more differentiated, “older” T cells. Accordingly, the inventive methods may, advantageously, identify and

isolate a TCR, or an antigen-binding portion thereof, that has antigenic specificity for the mutated amino acid sequence and introduce the TCR, or an antigen-binding portion thereof, into “younger” T cells that may provide any one or more of greater *in vivo* persistence, proliferation, and antitumor activity as compared to “older” T cells (e.g., effector cells in a patient’s tumor) from which the TCR, or the antigen-binding portion thereof, may have been isolated.

[0061] In an embodiment of the invention, the method further comprises expanding the numbers of PBMC that express the TCR, or the antigen-binding portion thereof. The numbers of PBMC may be expanded, for example, as described herein with respect to other aspects of the invention. In this regard, the inventive methods may, advantageously, generate a large number of T cells having antigenic specificity for the mutated amino acid sequence.

[0062] Another embodiment of the invention provides a TCR, or an antigen-binding portion thereof, isolated by any of the methods described herein with respect to other aspects of the invention. An embodiment of the invention provides a TCR comprising two polypeptides (i.e., polypeptide chains), such as an alpha (α) chain of a TCR, a beta (β) chain of a TCR, a gamma (γ) chain of a TCR, a delta (δ) chain of a TCR, or a combination thereof. Another embodiment of the invention provides an antigen-binding portion of the TCR comprising one or more CDR regions, one or more variable regions, or one or both of the α and β chains of the TCR, as described herein with respect to other aspects of the invention. The polypeptides of the inventive TCR, or the antigen-binding portion thereof, can comprise any amino acid sequence, provided that the TCR, or the antigen-binding portion thereof, has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation.

[0063] Another embodiment of the invention provides an isolated population of cells prepared according to any of the methods described herein with respect to other aspects of the invention. The population of cells can be a heterogeneous population comprising the PBMC expressing the isolated TCR, or the antigen-binding portion thereof, in addition to at least one other cell, e.g., a host cell (e.g., a PBMC), which does not express the isolated TCR, or the antigen-binding portion thereof, or a cell other than a T cell, e.g., a B cell, a macrophage, a neutrophil, an erythrocyte, a hepatocyte, an endothelial cell, an epithelial cells, a muscle cell, a brain cell, etc. Alternatively, the population of cells can be a substantially homogeneous population, in which the population comprises mainly of PBMC (e.g., consisting essentially of) expressing the isolated TCR, or the antigen-binding portion thereof. The population also

can be a clonal population of cells, in which all cells of the population are clones of a single PBMC expressing the isolated TCR, or the antigen-binding portion thereof, such that all cells of the population express the isolated TCR, or the antigen-binding portion thereof. In one embodiment of the invention, the population of cells is a clonal population comprising PBMC expressing the isolated TCR, or the antigen-binding portion thereof, as described herein. By introducing the nucleotide sequence encoding the isolated TCR, or the antigen binding portion thereof, into PBMC, the inventive methods may, advantageously, provide a population of cells that comprises a high proportion of PBMC cells that express the isolated TCR and have antigenic specificity for the mutated amino acid sequence. In an embodiment of the invention, about 1% to about 100%, for example, about 1%, about 5%, about 10%, about 15%, about 20%, about 25%, about 30%, about 35%, about 40%, about 45%, about 50%, about 55%, about 60%, about 65%, about 70%, about 75%, about 80%, about 85%, about 90%, about 95%, about 96%, about 97%, about 98%, about 99%, or about 100%, or a range defined by any two of the foregoing values, of the population of cells comprises PBMC cells that express the isolated TCR and have antigenic specificity for the mutated amino acid sequence. Without being bound to a particular theory or mechanism, it is believed that populations of cells that comprise a high proportion of PBMC cells that express the isolated TCR and have antigenic specificity for the mutated amino acid sequence have a lower proportion of irrelevant cells that may hinder the function of the PBMC, e.g., the ability of the PBMC to target the destruction of cancer cells and/or treat or prevent cancer.

[0064] The inventive TCRs, or the antigen-binding portions thereof, and populations of cells can be formulated into a composition, such as a pharmaceutical composition. In this regard, the invention provides a pharmaceutical composition comprising any of the inventive TCRs, or the antigen-binding portions thereof, or populations of cells and a pharmaceutically acceptable carrier. The inventive pharmaceutical composition can comprise an inventive TCR, or an antigen-binding portion thereof, or population of cells in combination with another pharmaceutically active agent(s) or drug(s), such as a chemotherapeutic agents, e.g., asparaginase, busulfan, carboplatin, cisplatin, daunorubicin, doxorubicin, fluorouracil, gemcitabine, hydroxyurea, methotrexate, paclitaxel, rituximab, vinblastine, vincristine, etc.

[0065] Preferably, the carrier is a pharmaceutically acceptable carrier. With respect to pharmaceutical compositions, the carrier can be any of those conventionally used for the particular inventive TCR, or the antigen-binding portion thereof, or population of cells under consideration. Such pharmaceutically acceptable carriers are well-known to those skilled in

the art and are readily available to the public. It is preferred that the pharmaceutically acceptable carrier be one which has no detrimental side effects or toxicity under the conditions of use.

[0066] The choice of carrier will be determined in part by the particular inventive TCR, the antigen-binding portion thereof, or population of cells, as well as by the particular method used to administer the inventive TCR, the antigen-binding portion thereof, or population of cells. Accordingly, there are a variety of suitable formulations of the pharmaceutical composition of the invention. Suitable formulations may include any of those for oral, parenteral, subcutaneous, intravenous, intramuscular, intraarterial, intrathecal, or interperitoneal administration. More than one route can be used to administer the inventive TCR or population of cells, and in certain instances, a particular route can provide a more immediate and more effective response than another route.

[0067] Preferably, the inventive TCR, the antigen-binding portion thereof, or population of cells is administered by injection, e.g., intravenously. When the inventive population of cells is to be administered, the pharmaceutically acceptable carrier for the cells for injection may include any isotonic carrier such as, for example, normal saline (about 0.90% w/v of NaCl in water, about 300 mOsm/L NaCl in water, or about 9.0 g NaCl per liter of water), NORMOSOL R electrolyte solution (Abbott, Chicago, IL), PLASMA-LYTE A (Baxter, Deerfield, IL), about 5% dextrose in water, or Ringer's lactate. In an embodiment, the pharmaceutically acceptable carrier is supplemented with human serum albumin.

[0068] It is contemplated that the inventive TCRs, the antigen-binding portions thereof, populations of cells, and pharmaceutical compositions can be used in methods of treating or preventing cancer. Without being bound to a particular theory or mechanism, the inventive TCRs, or the antigen-binding portions thereof, are believed to bind specifically to a mutated amino acid sequence encoded by a cancer-specific mutation, such that the TCR, or the antigen-binding portion thereof, when expressed by a cell, is able to mediate an immune response against a target cell expressing the mutated amino acid sequence. In this regard, the invention provides a method of treating or preventing cancer in a mammal, comprising administering to the mammal any of the pharmaceutical compositions, TCRs, antigen-binding portions thereof, or populations of cells described herein, in an amount effective to treat or prevent cancer in the mammal.

[0069] The terms "treat," and "prevent" as well as words stemming therefrom, as used herein, do not necessarily imply 100% or complete treatment or prevention. Rather, there are

varying degrees of treatment or prevention of which one of ordinary skill in the art recognizes as having a potential benefit or therapeutic effect. In this respect, the inventive methods can provide any amount of any level of treatment or prevention of cancer in a mammal.

Furthermore, the treatment or prevention provided by the inventive method can include treatment or prevention of one or more conditions or symptoms of the cancer being treated or prevented. For example, treatment or prevention can include promoting the regression of a tumor. Also, for purposes herein, "prevention" can encompass delaying the onset of the cancer, or a symptom or condition thereof.

[0070] For purposes of the invention, the amount or dose of the inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition administered (e.g., numbers of cells when the inventive population of cells is administered) should be sufficient to effect, e.g., a therapeutic or prophylactic response, in the mammal over a reasonable time frame. For example, the dose of the inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition should be sufficient to bind to a mutated amino acid sequence encoded by a cancer-specific mutation, or detect, treat or prevent cancer in a period of from about 2 hours or longer, e.g., 12 to 24 or more hours, from the time of administration. In certain embodiments, the time period could be even longer. The dose will be determined by the efficacy of the particular inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition administered and the condition of the mammal (e.g., human), as well as the body weight of the mammal (e.g., human) to be treated.

[0071] Many assays for determining an administered dose are known in the art. For purposes of the invention, an assay, which comprises comparing the extent to which target cells are lysed or IFN- γ is secreted by T cells expressing the inventive TCR, or the antigen-binding portion thereof, upon administration of a given dose of such T cells to a mammal among a set of mammals of which is each given a different dose of the T cells, could be used to determine a starting dose to be administered to a mammal. The extent to which target cells are lysed or IFN- γ is secreted upon administration of a certain dose can be assayed by methods known in the art.

[0072] The dose of the inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition also will be determined by the existence, nature and extent of any adverse side effects that might accompany the administration of a particular inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical

composition. Typically, the attending physician will decide the dosage of the inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition with which to treat each individual patient, taking into consideration a variety of factors, such as age, body weight, general health, diet, sex, inventive TCR, the antigen-binding portion thereof, population of cells, or pharmaceutical composition to be administered, route of administration, and the severity of the condition being treated.

[0073] In an embodiment in which the inventive population of cells is to be administered, the number of cells administered per infusion may vary, for example, in the range of one million to 100 billion cells; however, amounts below or above this exemplary range are within the scope of the invention. For example, the daily dose of inventive host cells can be about 1 million to about 150 billion cells (e.g., about 5 million cells, about 25 million cells, about 500 million cells, about 1 billion cells, about 5 billion cells, about 20 billion cells, about 30 billion cells, about 40 billion cells, about 60 billion cells, about 80 billion cells, about 100 billion cells, about 120 billion cells, about 130 billion cells, about 150 billion cells, or a range defined by any two of the foregoing values), preferably about 10 million to about 130 billion cells (e.g., about 20 million cells, about 30 million cells, about 40 million cells, about 60 million cells, about 70 million cells, about 80 million cells, about 90 million cells, about 10 billion cells, about 25 billion cells, about 50 billion cells, about 75 billion cells, about 90 billion cells, about 100 billion cells, about 110 billion cells, about 120 billion cells, about 130 billion cells, or a range defined by any two of the foregoing values), more preferably about 100 million cells to about 130 billion cells (e.g., about 120 million cells, about 250 million cells, about 350 million cells, about 450 million cells, about 650 million cells, about 800 million cells, about 900 million cells, about 3 billion cells, about 30 billion cells, about 45 billion cells, about 50 billion cells, about 75 billion cells, about 90 billion cells, about 100 billion cells, about 110 billion cells, about 120 billion cells, about 130 billion cells, or a range defined by any two of the foregoing values).

[0074] For purposes of the inventive methods, wherein populations of cells are administered, the cells can be cells that are allogeneic or autologous to the mammal. Preferably, the cells are autologous to the mammal.

[0075] Another embodiment of the invention provides any of the TCRs, the antigen-binding portions thereof, isolated population of cells, or pharmaceutical compositions described herein for use in treating or preventing cancer in a mammal.

[0076] The cancer may, advantageously, be any cancer, including any of acute lymphocytic cancer, acute myeloid leukemia, alveolar rhabdomyosarcoma, bone cancer, brain cancer, breast cancer, cancer of the anus, anal canal, or anorectum, cancer of the eye, cancer of the intrahepatic bile duct, cancer of the joints, cancer of the neck, gallbladder, or pleura, cancer of the nose, nasal cavity, or middle ear, cancer of the oral cavity, cancer of the vagina, cancer of the vulva, cholangiocarcinoma, chronic lymphocytic leukemia, chronic myeloid cancer, colon cancer, esophageal cancer, uterine cervical cancer, gastrointestinal carcinoid tumor, glioma, Hodgkin lymphoma, hypopharynx cancer, kidney cancer, larynx cancer, liver cancer, lung cancer, malignant mesothelioma, melanoma, multiple myeloma, nasopharynx cancer, non-Hodgkin lymphoma, cancer of the oropharynx, ovarian cancer, cancer of the penis, pancreatic cancer, peritoneum, omentum, and mesentery cancer, pharynx cancer, prostate cancer, rectal cancer, renal cancer, skin cancer, small intestine cancer, soft tissue cancer, stomach cancer, testicular cancer, thyroid cancer, cancer of the uterus, ureter cancer, urinary bladder cancer, solid tumors, and liquid tumors. Preferably, the cancer is an epithelial cancer. In an embodiment, the cancer is cholangiocarcinoma, melanoma, colon cancer, or rectal cancer.

[0077] The mammal referred to in the inventive methods can be any mammal. As used herein, the term "mammal" refers to any mammal, including, but not limited to, mammals of the order Rodentia, such as mice and hamsters, and mammals of the order Logomorpha, such as rabbits. It is preferred that the mammals are from the order Carnivora, including Felines (cats) and Canines (dogs). Preferably, the mammals are from the order Artiodactyla, including Bovines (cows) and Swines (pigs) or of the order Perssodactyla, including Equines (horses). Preferably, the mammals are of the order Primates, Ceboids, or Simoids (monkeys) or of the order Anthropoids (humans and apes). A more preferred mammal is the human. In an especially preferred embodiment, the mammal is the patient expressing the cancer-specific mutation.

[0078] The following examples further illustrate the invention but, of course, should not be construed as in any way limiting its scope.

EXAMPLES

[0079] The materials and methods for Examples 1-7 are set forth below.

Whole-exomic sequencing

[0080] Whole-exomic sequencing of cryopreserved tumor tissue (embedded in OCT) and normal peripheral blood cells was performed by Personal Genome Diagnostics (PGDx, Baltimore, MD) as described in Jones et al., *Science* 330: 228-231 (2010). The average number of distinct high quality sequence reads at each base was 155 and 160 for tumor and normal (PBMC) DNA, respectively.

Patient treatment and generation of tumor infiltrating lymphocytes (TIL) for adoptive cell therapy

[0081] Patient 3737 was enrolled in the institutional-review board (IRB)-approved protocol: "A Phase II Study Using Short-Term Cultured, Autologous Tumor-Infiltrating Lymphocytes Following a Lymphocyte Depleting Regimen in Metastatic Digestive Tract Cancers" (Trial registration ID: NCT01174121), which was designed to evaluate the safety and effectiveness of the adoptive transfer of autologous, *ex vivo* expanded tumor-infiltrating lymphocytes (TIL) in patients with gastrointestinal cancers.

[0082] TIL used for patient's first treatment was generated as described in Jin et al., *J. Immunother.*, 35: 283-292 (2012). Briefly, resected tumors were minced into approximately 1-2 mm fragments and individual fragments were placed in wells of a 24-well plate containing 2 ml of complete media (CM) containing high dose IL-2 (6000 IU/ml, Chiron, Emeryville, CA). CM consisted of RPMI supplemented with 10% in-house human serum, 2 mM L-glutamine, 25 mM HEPES and 10 µg/ml gentamicin. Additionally, a mixed tumor digest was also cultured in CM with high dose IL-2. After the initial outgrowth of T cells (between 2-3 weeks), 5×10^6 T cells from select cultures were rapidly expanded in gas-permeable G-Rex100 flasks using irradiated allogeneic PBMC at a ratio of 1 to 100 in 400 ml of 50/50 medium, supplemented with 5% human AB serum, 3000 IU/ml of IL-2, and 30 ng/ml of OKT3 antibody (Miltenyi Biotec, Bergisch Gladbach, Germany). 50/50 media was composed of a 1 to 1 mixture of CM with AIM-V media. All cells were cultured at 37 °C with 5% CO₂. The numbers of cells were rapidly expanded for two weeks prior to infusion. Patient 3737 underwent a non-myeloablative lymphodepleting regimen composed of cyclophosphamide and fludarabine prior to receiving 42.4 billion total T cells in conjunction with four doses of high dose IL-2.

[0083] TIL used for the patient's second treatment was generated in a similar manner as the first treatment with the following changes. The first treatment product (Patient 3737-TIL) was composed of a combination of 5 individual TIL cultures. These 5 cultures were individually assessed for expression of CD4 and V β 22, and reactivity against mutated ERBB2IP, and one culture was found to be highly enriched in V β 22+ ERBB2IP-mutation-reactive CD4+ T cells. This one TIL culture (after the initial outgrowth with high dose IL-2) was then rapidly expanded as described above. The patient underwent an identical non-myeloablative lymphodepleting regimen as the first treatment prior to receiving 126 billion total T cells in conjunction with four doses of high dose IL-2.

Generation of TMG constructs

[0084] Briefly, for each non-synonymous substitution mutation identified by whole exome sequencing, a "minigene" construct encoding the corresponding amino acid change flanked by 12 amino acids of the wild-type protein sequence was made. Multiple minigenes were genetically fused together to generate a TMG construct. These minigene constructs were codon optimized and synthesized as DNA String constructs (Life Technologies, Carlsbad CA). TMGs were then cloned into the pcDNA3.1 vector using In-Fusion technology (Clontech, Mountain View, CA). Site-directed mutagenesis was used to generate the nine "wild-type reversion" TMG-1 constructs (Gene Oracle, Mountain View, CA). The nucleotide sequence of all TMGs was verified by standard Sanger sequencing (Macrogen and Gene Oracle).

Generation of autologous APCs

[0085] Monocyte-derived, immature DCs were generated using the plastic adherence method. Briefly, autologous pheresis samples were thawed, washed, set to $5-10 \times 10^6$ cells/ml with neat AIM-V media (Life Technologies) and then incubated at approximately 1×10^6 cells/cm² in an appropriate sized tissue culture flask and incubated at 37 °C, 5% CO₂. After 90 minutes (min), non-adherent cells were collected, and the flasks were vigorously washed with AIM-V media, and then incubated with AIM-V media for another 60 min. The flasks were then vigorously washed again with AIM-V media and then the adherent cells were incubated with DC media. DC media comprised of RPMI containing 5% human serum (collected and processed in-house), 100 U/ml penicillin and 100 μ g/ml streptomycin, 2 mM L-glutamine, 800 IU/ml GM-CSF and 800 U/ml IL-4 (media supplements were from Life

Technologies and cytokines were from Peprtech). On day 3, fresh DC media was added to the cultures. Fresh or freeze/thawed DCs were used in experiments on day 5-7 after initial stimulation. In all experiments, flow cytometry was used to phenotype the cells for expression of CD11c, CD14, CD80, CD86, and HLA-DR (all from BD Bioscience) to ensure that the cells were predominantly immature DCs (CD11c+, CD14-, CD80^{low}, CD86+, and HLA-DR+; data not shown).

[0086] Antigen presenting B cells were generated using the CD40L and IL-4 stimulation method. Briefly, human CD19-microbeads (Miltenyi Biotec) were used to positively select B cells from autologous pheresis samples. CD19+ cells were then cultured with irradiated (6000 rad) 3T3 cells stably expressing CD40L (3T3-CD40L) at approximately a 1:1 ratio in B-cell media. B-cell media comprised of IMDM media (Life Technologies) supplemented with 7.5-10% human serum (in-house), 100 U/ml penicillin and 100 µg/ml streptomycin (Life Technologies), 10 µg/ml gentamicin (CellGro, Manassas, VA), 2 mM L-glutamine (Life Technologies), and 200 U/ml IL-4 (Peprtech). Fresh B-cell media was added starting on day 3, and media added or replaced every 2-3 days thereafter. Additional irradiated 3T3-CD40L feeder cells were also added as required. Antigen presenting B cells were typically used in experiments 2-3 weeks after initial stimulation.

Generation of in vitro transcribed RNA (IVT) RNA

[0087] Plasmids encoding the tandem minigenes were linearized with the restriction enzyme Sac II. A control pcDNA3.1/V5-His-TOPO vector encoding GFP was linearized with Not I. Restriction digests were terminated with EDTA, sodium acetate and ethanol precipitation. Complete plasmid digestion was verified by standard agarose gel electrophoresis. Approximately 1 µg of linearized plasmid was used for the generation of IVT RNA using the message machine T7 Ultra kit (Life Technologies) as directed by the manufacturer. RNA was precipitated using the LiCl₂ method, and RNA purity and concentrations were assessed using a NanoDrop spectrophotometer. RNA was then aliquoted into microtubes and stored at -80 °C until use.

RNA transfections

[0088] APCs (DCs or B cells) were harvested, washed 1x with PBS, and then resuspended in Opti-MEM (Life Technologies) at 10-30 x 10⁶ cells/ml. IVT RNA (4 µg or 8 µg) was aliquoted to the bottom of a 2 mm gap electroporation cuvette, and 50 µl or 100 µl of

APCs were added directly to the cuvette. The final RNA concentration used in electroporations was thus 80 µg/ml. Electroporations were carried out using a BTX-830 square wave electroporator. DCs were electroporated with 150 V, 10 ms, and 1 pulse, and B cells were electroporated with 150 V, 20 ms, and 1 pulse. Transfection efficiencies using these settings were routinely between 70-90% as assessed with GFP RNA (data not shown). All steps were carried out at room temperature. Following electroporation, cells were immediately transferred to polypropylene tubes containing DC- or B-cell media supplemented with the appropriate cytokines. Transfected cells were incubated overnight (12-14 h) at 37 °C, 5% CO₂. Cells were washed 1x with PBS prior to use in co-culture assays.

Peptide pulsing

[0089] Autologous B cells were harvested, washed, and then resuspended at 1×10^6 cells/ml in B-cell media supplemented with IL-4, and then incubated with 1 µg/ml of a 25-mer peptide overnight (12-14 h) at 37°C, 5% CO₂. After overnight pulsing, B cells were then washed 2x with PBS, and then resuspended in T-cell media and immediately used in co-culture assays. The peptides used were: mutated ERBB2IP (TSFLSINSKEETGHLNGNKYPNLE (SEQ ID NO: 73)); wild-type ERBB2IP (TSFLSINSKEETEHLNGNKYPNLE (SEQ ID NO: 45)); and, as a negative control, mutated ALK (RVLKGGSVRKL RHAKQLVLELGEEA (SEQ ID NO: 46)). The mutated ERBB2IP peptide was purchased from three different sources (GenScript, Piscataway, NJ, Peptide 2.0, Chantilly, VA, and SelleckChem, Houston TX) with all yielding the same *in vitro* results, while the wild-type ERBB2IP and mutated ALK peptides were purchased from Peptide 2.0. For culturing allogeneic EBV-B cells, RPMI media containing 10% FBS, 100 U/ml penicillin and 100 µg/ml streptomycin (Life Technologies), 10 µg/ml gentamicin (CellGro), and 2 mM L-glutamine was used instead of B-cell media.

T-cell sorting, expansion, and cloning

[0090] The BD FACSAria IIu and BD FACSJazz were used in all experiments requiring cell sorting. In indicated experiments, sorted T cells were expanded using excess irradiated (4000 rad) allogeneic feeder cells (pool of three different donor leukapheresis samples) in 50/50 media containing 30 ng/ml anti-CD3 antibody (OKT3) and 3000 IU/ml IL-2. Limiting dilution cloning was carried out in 96-well round bottom plates using the above stimulation

conditions with 5e4 feeder cells per well and 1-2 T cells per well. Media was exchanged starting at approximately 1 week post stimulation and then every other day or as required. Cells were typically used in assays, or further expanded, at approximately 2-3 weeks after the initial stimulation.

Co-culture assays: IFN- γ ELISPOT and ELISA, flow cytometry for cell surface activation markers, and intracellular cytokine staining (ICS)

[0091] When DCs were used as APCs, approximately 3.5×10^4 to 7×10^4 DCs were used per well of a 96-well flat or round-bottom plate. When B cells were used as APCs, approximately 2×10^5 cells were used per well of a 96-well round-bottom plate. In ELISPOT assays, 1×10^3 to 1×10^4 effector T cells were used per well, and in flow cytometry assays, 1×10^5 effector T cells were used per well. T cells were typically thawed and rested in IL-2 containing 50/50 media (3000 IU/ml IL-2) for two days and then washed with PBS (3x) prior to co-culture assays. All co-cultures were performed in the absence of exogenously added cytokines. For all assays, plate-bound OKT3 (0.1 μ g/ml or 1 μ g/ml) was used as a positive control.

[0092] In experiments involving HLA blocking antibodies, the following antibodies were used: pan-class-II (clone: IVA12), pan-class-I (clone: W6/32), HLA-DR (clone: HB55), HLA-DP (clone: B7/21), and HLA-DQ (clone: SPV-L3). Cells were blocked with 20-50 μ g/ml of the indicated antibody for 1-2 h at 37 °C, 5% CO₂ prior to co-culture with T cells. T4 are T cells that have been transduced with an HLA-DR4-restricted TCR that is reactive against an epitope in tyrosinase. DMF5 is an HLA-A2-restricted T-cell line reactive against MART-1. 624-CIITA is a HLA-A2 and HLA-DR4-positive melanoma cell line that stably expresses MHC-II due to ectopic expression of CIITA (class II, MHC, transactivator), and is positive for MART-1 and tyrosinase expression.

[0093] For IFN- γ ELISPOT assays, briefly, ELIIP plates (Millipore, MAIPSWU) were pre-treated with 50 μ l of 70% ethanol per well for 2 min, washed 3x with PBS, and then coated with 50 μ l of 10 μ g/ml IFN- γ capture antibody (Mabtech, clone: 1-D1K) and incubated overnight in the fridge. For OKT3 controls, wells were coated with a mixture of IFN- γ capture antibody (10 μ g/ml) and OKT3 (1 μ g/ml). Prior to co-culture, the plates were washed 3x with PBS, followed by blocking with 50/50 media for at least 1 h at room temperature (RT). After 20-24 h of co-culture, cells were flicked out of the plate, washed 6x

with PBS + 0.05% Tween-20 (PBS-T), and then incubated for 2 h at RT with 100 μ l/well of a 0.22 μ m filtered 1 μ g/ml biotinylated anti-human IFN- γ detection antibody solution (Mabtech, clone: 7-B6-1). The plate was then washed 3x with PBS-T, followed by a 1 h incubation with 100 μ l/well of streptavidin-ALP (Mabtech, Cincinnati, OH, diluted 1:3000). The plate was then washed 6x with PBS followed by development with 100 μ l/well of 0.45 μ m filtered BCIP/NBT substrate solution (KPL, Inc.). The reaction was stopped by rinsing thoroughly with cold tap water. ELISPOT plates were scanned and counted using an ImmunoSpot plate reader and associated software (Cellular Technologies, Ltd, Shaker Heights, OH).

[0094] Expression of the T-cell activation markers OX40 and 4-1BB was assessed by flow cytometry at approximately t=22-26 h post-stimulation. Briefly, cells were pelleted, washed with FACS buffer (1X PBS supplemented with 1% FBS and 2 mM EDTA), and then stained with the appropriate antibodies for approximately 30 min, at 4 °C in the dark. Cells were washed at least once with FACS buffer prior to acquisition on a BD FACSCanto II flow cytometer. All data were gated on live (PI negative), single cells.

[0095] Cytokine production was assessed using intracellular cytokine staining (ICS) and flow cytometry. Briefly, after target and effector cells were combined in the wells of a 96-well plate, both GolgiStop and GolgiPlug were added to the culture (BD Biosciences). GolgiStop and GolgiPlug were used at 1/2 of the concentration recommended by the manufacturer. At t=6 h post stimulation, cells were processed using the Cytofix/Cytoperm kit (BD Biosciences, San Jose, CA) according to the manufacturer's instructions. Briefly, cells were pelleted, washed with FACS buffer, and then stained for cell surface markers (described above). Cells were then washed 2x with FACS buffer prior to fixation and permeabilization. Cells were then washed with Perm/Wash buffer and stained with antibodies against cytokines for 30 min, at 4 °C in the dark. Cells were washed 2x with Perm/Wash buffer and resuspended in FACS buffer prior to acquisition on a FACSCantoII flow cytometer. All flow cytometry data were analyzed using FLOWJO software (TreeStar Inc).

[0096] IFN- γ in serum samples was detected using a human IFN- γ ELISA kit as directed by the manufacturer (Thermo Scientific, Waltham, MA).

Flow cytometry antibodies

[0097] The following titrated anti-human antibodies were used for cell surface staining: CCR7-FITC (clone: 150503), CD45RO-PE-Cy7 (clone: UCHL1), CD62L-APC (clone:

DREG-56), CD27-APC-H7 (clone: M-T271), CD4-eFluor 605NC (clone: OKT4), CD57-FITC (clone: NK-1), CD28-PE-Cy7 (clone: CD28.2), CD127-APC (clone: eBioRDR5), CD3-AF700 (clone: UCHT1), CD4-FITC, PE-Cy7, APC-H7 (clone: SK3), CD8-PE-Cy7 (clone: SK1), V β 22-PE (clone: IMMU 546), V β 5.2-PE (clone: 36213), OX40-PE-Cy7 or FITC (clone: Ber-ACT35), 4-1BB-APC (clone: 4B4-1), and CD107a-APC-H7 (clone: H4A3). All antibodies were from BD Biosciences, except CD4-eFluor605NC (eBioscience), V β 22-PE and V β 5.2-PE (Beckman Coulter), and 4-1BB-APC and OX40-PE-Cy7 (BioLegend). The following optimally titrated anti-human antibodies were used for intracellular cytokine staining: IFN- γ -FITC (clone: 4S.B3), IL-2-APC (clone: MQ1-17H12), TNF-PerCP-Cy5.5 or APC (clone: MAb11), IL-17-PE (clone: eBio64DEC17), and IL-4-PE-Cy7 (clone: 8D4-8). All ICS antibodies were from eBioscience except IL-4-PE-Cy7 (BD Bioscience). The IO Mark B Mark TCR V kit was used to assess the TCR-V β repertoire (Beckman Coulter).

Sequencing of the ERBB2IP mutation

[0098] Sanger sequencing was used to validate the *ERBB2IP* mutation found by whole-exomic sequencing. Total RNA was extracted from snap frozen T cells or tumor tissues (OCT block) using the RNeasy Mini kit (Qiagen). Total RNA was then reverse transcribed to cDNA using ThermoScript reverse transcriptase with oligo-dT primers (Life Technologies). Normal and tumor cDNA were then used as templates in a PCR with the following *ERBB2IP* primers flanking the mutation: *ERBB2IP* Seq Forward: 5'—TGT TGA CTC AAC AGC CAC AG—3' (SEQ ID NO: 47); and *ERBB2IP* Seq Reverse: 5'—CTG GAC CAC TTT TCT GAG GG—3' (SEQ ID NO: 48). Phusion DNA polymerase (Thermo Scientific) was used with the recommended 3-step protocol with a 58 °C annealing temperature (15 sec) and a 72 °C extension (30 sec). PCR products were isolated by standard agarose gel electrophoresis and gel extraction (Clontech). Products were directly sequenced using the same PCR primers (Macrogen).

Quantitative PCR

[0099] Total RNA was extracted from snap frozen T cells or tumor tissues (OCT block) using the RNeasy Mini kit (Qiagen, Venlo, Netherlands). Total RNA was then reverse transcribed to cDNA using qScript cDNA Supermix (Quanta Biosciences, Gaithersburg,

MD). Gene-specific Taqman primer and probe sets for human β -actin (catalogue #: 401846) and ERBB2IP (catalogue #: 4331182) were purchased from Life Technologies. Quantitative PCR was carried out with TAQMAN Fast Advanced Master Mix using the 7500 Fast Real Time PCR machine (both from Applied Biosystems). Specificity of amplified products was verified by standard agarose gel electrophoresis. All calculated threshold cycles (Ct) were 30 or below.

TCR-V β deep sequencing

[0100] TCR-V β deep sequencing was performed by immunoSEQ, Adaptive Biotechnologies (Seattle, WA) on genomic DNA isolated from peripheral blood, T cells, and frozen tumor tissue using the DNeasy blood and tissue kit (Qiagen). The number of total productive TCR reads per sample ranged from 279, 482 to 934,672. Only productive TCR rearrangements were used in the calculations of TCR frequencies.

TCR sequencing and construction of the ERBB2IP-mutation reactive TCR

[0101] T cells were pelleted and total RNA isolated (RNeasy Mini kit, Qiagen). Total RNA then underwent 5'RACE as directed by manufacturer (SMARTer RACE cDNA amplification kit, Clontech) using TCR- α and - β chain constant primers. Program 1 of the kit was used for the PCR, with a modification to the extension time (2 min instead of 3 min). The sequences of the α and β chain constant primers are: TCR- α , 5'—GCC ACA GCA CTG TGC TCT TGA AGT CC—3' (SEQ ID NO: 49); TCR- β , 5'—CAG GCA GTA TCT GGA GTC ATT GAG—3' (SEQ ID NO: 50). TCR PCR products were then isolated by standard agarose gel electrophoresis and gel extraction (Clontech). Products were then either directly sequenced or TOPO-TA cloned followed by sequencing of individual colonies (Macrogen). For sequencing of known V β 22+ T-cell clones, cDNA was generated from RNA using qScript cDNA Supermix (Quanta Biosciences). These cDNAs then were used as templates in a PCR using the TCR- β constant primer (above) and the V β 22-specific primer: 5'—CAC CAT GGA TAC CTG GCT CGT ATG C—3' (SEQ ID NO: 51). PCR products were isolated by standard agarose gel electrophoresis and gel extraction (Clontech). Products were directly sequenced (Macrogen) using the nested TCR- β chain constant primer: 5'—ATT CAC CCA CCA GCT CAG—3' (SEQ ID NO: 52).

[0102] Construction of the V β 22+ ERBB2IP-mutation TCR was done by fusing the V β 22+ TCR- α V-D-J regions to the mouse TCR- α constant chain, and the V β 22+ TCR- β -V-

D-J regions to the mouse TCR- β constant chains. The α and β chains were separated by a furin SGSG P2A linker. Use of mouse TCR constant regions promotes pairing of the introduced TCR and also facilitates identification of positively transduced T cells by flow cytometry using an antibody specific for the mouse TCR- β chain (eBioscience). The TCR construct was synthesized and cloned into the MSGV1 retroviral vector (Gene Oracle).

TCR transduction of peripheral blood T cells

[0103] Autologous pheresis samples were thawed and set to 2×10^6 cells/ml in T-cell media, which consists of a 50/50 mixture of RPMI and AIM-V media supplemented with 5% in-house human serum, 10 μ g/ml gentamicin (CellGro), 100 U/ml penicillin and 100 μ g/ml streptomycin, 1.25 μ g/ml amphotericin B (Fungizone) and 2 mM L-glutamine (all from Life Technologies). 2×10^6 cells (1 ml) were stimulated in a 24-well plate with 50 ng/ml soluble OKT3 (Miltenyi Biotec) and 300 IU/ml rhu IL-2 (Chiron) for 2 days prior to retroviral transduction. To generate transient retroviral supernatants, the retroviral vector MSGV1 encoding the V β 22-positive, ERBB2IP-mutation-specific TCR (1.5 μ g/well) and the envelope encoding plasmid RD114 (0.75 μ g/well) were co-transfected into the retroviral packaging cell line 293GP (1×10^6 cells per well of a 6-well poly-D-lysine-coated plates, plated the day prior to transfection) using lipofectamine 2000 (Life Technologies). Retroviral supernatants were collected at 42-48 h after transfection, diluted 1:1 with DMEM media, and then centrifuged onto retronectin-coated (10 μ g/ml, Takara), non-tissue culture-treated 6-well plates at 2,000 g for 2 h at 32 °C. Activated T cells (2×10^6 per well, at 0.5×10^6 cells/ml in IL-2 containing T-cell media) were then spun onto the retrovirus plates for 10 min at 300 g. Activated T cells were transduced overnight, removed from the plates and further cultured in IL-2 containing T-cell media. GFP and mock transduction controls were included in transduction experiments. Cells were typically assayed 10-14 days post-retroviral transduction.

EXAMPLE 1

[0104] This example demonstrates a method of identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence.

[0105] A 43-year old woman with widely metastatic cholangiocarcinoma (Patient (Pt.) 3737) who progressed through multiple chemotherapy regimens was enrolled onto a TIL-

based adoptive cell therapy (ACT) protocol for patients with gastrointestinal (GI) cancers. The clinical characteristics of patient 3737 are shown in Table 1.

TABLE 1

Sex	Age	Primary	Metastatic sites	Prior Therapy	Prior IL-2	Harvest site*	ECOG ⁺ Status	HLA-I	HLA-II
F	43	Intrahepatic cholangiocarcinoma (poorly differentiated)	Lungs, liver	Cisplatin + gemcitabine, taxotere	No	Lung	0	A*26 B*38 B*52 C*12	DRB1*0405 DRB1*1502 DQB1*0301 DQB1*0601 DPB1*0401 DPB1*10401

* Harvest site for generation of TIL and for whole exomic sequencing.

+ Performance status: ECOG, Eastern Cooperative Oncology Group

[0106] Lung metastases were resected and used as a source for whole-exomic sequencing and generation of T cells for treatment. Table 2 shows the somatic mutations identified by whole-exome sequencing of a metastatic lung nodule from patient 3737. The tumor nodule was estimated to be approximately 70% tumor by pathological analysis of a hematoxylin and eosin (H&E) stained section. Whole-exomic sequencing revealed 26 non-synonymous mutations (Table 2).

TABLE 2

Gene Symbol	Gene Description	Transcript Accession	Mutation Position		Mutation Type	Consequence	% Mutant Reads*
			Nucleotide (genomic)	Amino Acid (protein)			
ALK	anaplastic lymphoma receptor tyrosine kinase	CCDS33172.1	chr2_29996620-29996620_C_T	137R>H	Substitution	Nonsynonymous coding	30%
AR	androgen receptor	CCDS14387.1	chrX_66858483-66858483_C	NA	Insertion	Frameshift	31%
CD93	CD93 molecule	CCDS13149.1	chr20_23012929-23012929_C_T	634R>Q	Substitution	Nonsynonymous coding	26%
DIP2C	DIP2 disco-interacting protein 2 homolog C (Drosophila)	CCDS7054.1	chr10_365545-365545_C_T	NA	Substitution	Splice site acceptor	25%
ERBB2IP	erbb2 interacting protein	CCDS3990.1	chr5_65385316-65385316_A_G	805E>G	Substitution	Nonsynonymous coding	59%
FCER1A	Fc fragment of IgE; high affinity I; receptor for; α polypeptide	CCDS1184.1	chr1_157544227-157544227_G_C	219D>H	Substitution	Nonsynonymous coding	30%
GRXCR1	glutaredoxin; cysteine rich 1	CCDS43225.1	chr4_42590102-42590102_C_T	21A>V	Substitution	Nonsynonymous coding	18%
HLA-DOA	HLA class II histocompatibility antigen, DO α chain precursor	CCDS4763.1	chr6_33085209-33085209_C_T	NA	Substitution	Splice site donor	36%
KIF9	kinesin family member 9	CCDS2752.1	chr3_47287859-47287859_T_C	155T>A	Substitution	Nonsynonymous coding	20%

Gene Symbol	Gene Description	Transcript Accession	Mutation Position		Mutation Type	Consequence	% Mutant Reads*
			Nucleotide (genomic)	Amino Acid (protein)			
KLHL6	kelch-like 6 (Drosophila)	CCDS3245.2	chr3_184692410-184692413_CAGA_	NA	Deletion	Frameshift	20%
LHX9	LIM homeobox 9	CCDS1393.1	chr1_196164923-196164923_A_	NA	Deletion	Frameshift	21%
LONRF3	LON peptidase N-terminal domain and ring finger 3	CCDS35374.1	chrX_118007666-118007666_A_C	NA	Substitution	Splice site donor	10%
NAGS	N-acetylglutamate synthase	CCDS11473.1	chr17_39440355-39440355_G_A	412R>H	Substitution	Nonsynonymous coding	29%
NLRP2	NLR family; pyrin domain containing 2	CCDS12913.1	chr19_60186650-60186650_G_T	591S>I	Substitution	Nonsynonymous coding	32%
PDZD2	PDZ domain containing 2	CCDS34137.1	chr5_32124833-32124833_A_	NA	Deletion	Frameshift	30%
POU5F2	POU domain, class 5, transcription factor 2	NM_153216	chr5_93102847-93102847_A_C	60V>G	Substitution	Nonsynonymous coding	34%
RAC3	ras-related C3 botulinum toxin substrate 3 (rho family; small GTP binding protein Rac3)	CCDS11798.1	chr17_77584690-77584690_C_A	125T>N	Substitution	Nonsynonymous coding	27%
RAP1GDS1	RAP1; GTP-GDP dissociation stimulator 1	CCDS43253.1	chr4_99532209-99532209_C_A	198L>I	Substitution	Nonsynonymous coding	19%
RASA1	RAS p21 protein activator (GTPase activating protein) 1	CCDS34200.1	chr5_86703757-86703757_C_T	589R>C	Substitution	Nonsynonymous coding	63%
RETSAT	retinol saturase (all-trans-	CCDS1972.1	chr2_85424308-	553R>K	Substitution	Nonsynonymous	11%

Gene Symbol	Gene Description	Transcript Accession	Mutation Position		Mutation Type	Consequence	% Mutant Reads*
			Nucleotide (genomic)	Amino Acid (protein)			
	retinol 13;14-reductase)		85424308_C_T			coding	
SEC24D	SEC24 family; member D (S. cerevisiae)	CCDS3710.1	chr4_119872085-119872085_A_G	901M>T	Substitution	Nonsynonymous coding	18%
SENP3	SUMO1/sentrin/SMT3 specific peptidase 3	ENST00000321337	chr17_7408824-7408824_A_G	292M>V	Substitution	Nonsynonymous coding	33%
SLIT1	slit homolog 1 (Drosophila)	CCDS7453.1	chr10_98753840-98753840_G_C	1280N>K	Substitution	Nonsynonymous coding	45%
TARBP1	TAR (HIV-1) RNA binding protein 1	CCDS1601.1	chr1_232649342-232649342_C_A	655G>V	Substitution	Nonsynonymous coding	18%
TGM6	transglutaminase 6	CCDS13025.1	chr20_2332325-2332325_G_A	398D>N	Substitution	Nonsynonymous coding	51%
TTC39C	tetratricopeptide repeat domain 39C	CCDS32804.1	chr18_19966475-19966475_A_C	503N>T	Substitution	Nonsynonymous coding	24%

EXAMPLE 2

[0107] This example demonstrates a method of inducing autologous APCs of a patient to present the mutated amino acid sequence; co-culturing a population of autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence; and selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a MHC molecule expressed by the patient.

[0108] For each mutation identified in Example 1, a mini-gene construct was designed that encoded for the mutated amino acid flanked on each side by 12 amino acids from the endogenous protein. Multiple mini-genes were synthesized in tandem to generate tandem mini-gene (TMG) constructs (Table 3). In Table 3, the underlining denotes mutated amino acids and neo-sequences encoded by point mutations, or nucleotide insertions or deletions. For splice-site donor mutations (*HLA-DOA* and *LONRF3*), mutant minigene transcripts were designed that continued into the downstream intron until the next stop codon, based on the assumption that the mutations prevented splicing at that site. The splice-site acceptor mutation in *DIP2C* was not assessed.

TABLE 3

TMG	Mutated Gene	Mutated Minigene Amino Acid Sequence	TMG Amino Acid Sequence
1	ALK	RVLKGGSVRKLRL <u>HAKQLVLELGEEA</u> (SEQ ID NO: 1)	RVLKGGSVRKLRL <u>HAKQLVLELGEEA</u> QNAADSYSWVPE <u>QAESRAMENQYSPTSFLSINSKEETG</u> HLENGNKYPNLEF IPLLVLFAV <u>HTGLFISTQQQVTESDRPRKVRFRIV</u> SSHS GRVLKEVVEIYNESLFDLLS <u>ALPYVGPSVTPMTGKCLRDD</u> YLASL <u>HPRLHSIYVSEGY</u> PDIKQELLRCDI <u>ICKGGHSTVTD</u> LQVGTCLDLRDDKD <u>NIERLRDCKKLAPI</u> (SEQ ID NO: 26)
	CD93	QNAADSYSWVPE <u>QAESRAMENQYSP</u> (SEQ ID NO: 2)	
	ERBB2IP	TSFLSINSKEET <u>G</u> HLENGNKYPNLE (SEQ ID NO: 3)	
	FCER1A	FIPLLVLFAV <u>HTGLFISTQQQVT</u> (SEQ ID NO: 4)	
	GRXCR1	ESDRPRKVRFRIV <u>SSHS</u> GRVLKEVY (SEQ ID NO: 5)	
	KIF9	EIYNESLFDLLS <u>ALPYVGPSVTPMT</u> (SEQ ID NO: 6)	
	NAGS	GKCLRDDYLASL <u>HPRLHSIYVSEGY</u> (SEQ ID NO: 7)	
	NLRP2	PDIKQELLRCDI <u>ICKGGHSTVTDLQ</u> (SEQ ID NO: 8)	
	RAC3	VGTCLDLRDDKD <u>NIERLRDCKKLAPI</u> (SEQ ID NO: 9)	
2	RAP1GDS1	VKLLGIHCQNAAI <u>TEMCLVAFGNLA</u> (SEQ ID NO: 10)	VKLLGIHCQNAAI <u>TEMCLVAFGNLANLRKSSPGTSNKCL</u> RQVSSLVLHIELGRLHPCVMASL <u>KAQSPINLYLTG</u> LLPI HTLDVKST <u>LPAAVRCSESRLMTMDNFGKHHTL</u> SEAP LYVGGMPVMTMDNFGKHHTL <u>SEAPLYVGGMPVHD</u> GPFVFAEVNAN <u>YITWLWHEDES</u> RQAKEDFSGYDFE <u>TRL</u> HVRIHAALASPVRPGICPGD <u>GWRIPGLPHEF</u> (SEQ ID NO: 27)
	RASA1	NLRKSSPGTSNK <u>CLRQVSSLVLHIE</u> (SEQ ID NO: 11)	
	RETSAT	LGRHLHPCVMASL <u>KAQSPINLYLTG</u> (SEQ ID NO: 12)	
	SEC24D	LLPIHTLDVKST <u>LPAAVRCSESRL</u> (SEQ ID NO: 13)	
	SLIT1	MTMDNFGKHHTL <u>SEAPLYVGGMPV</u> (SEQ ID NO: 14)	
	TARBP1	AVDVEGMKTQYS <u>VKQRTENVLRIFL</u> (SEQ ID NO: 15)	
	TGM6	HDGPFVFAEVNAN <u>YITWLWHEDES</u> (SEQ ID NO: 16)	
	TTC39C	QAKEDFSGYDFE <u>TRLHVRIHAALAS</u> (SEQ ID NO: 17)	
	POU5F2	PAVRPGICPGD <u>GWRIPGLPHEF</u> (SEQ ID NO: 18)	
	3	SEN3	
LHX9		GTATTLTDLTNPL <u>LSL</u> (SEQ ID NO: 20)	
KLHL6		THIRRVPGAVSD <u>GRMGSWRAPPTLS</u> V PASPLTLQSHFRQARV (SEQ ID NO: 21)	
AR		RHLSQEFGLQITP <u>PGIPVHESTATLQH</u> <u>YSSGWAEKSKIL</u> (SEQ ID NO: 22)	
PDZD2		SPDSKIQMVSSSQ <u>KRALLCLIALLSRKQT</u> <u>WKIRTCLRRVRQKCF</u> (SEQ ID NO: 23)	
HLA-DOA		TLLSPQEAGATK <u>DECEGEEGAAGSRDLR</u> <u>SWVT</u> (SEQ ID NO: 24)	
LONRF3		EETGMPNKASKO <u>GPGSTQREGSLEEIPG</u> <u>LTNIYKLLTSVWGLLRVWVWGPALAF</u> T	

TMG	Mutated Gene	Mutated Minigene Amino Acid Sequence	TMG Amino Acid Sequence
		<u>CVTSEIAMRL</u> L (SEQ ID NO: 25)	

[0109] The TMG constructs were then used as templates for the generation of *in vitro* transcribed (IVT) RNA. Each of these IVT TMG RNAs was then individually transfected into autologous APCs (DCs) followed by a co-culture with TIL to determine whether any of the processed and presented mutated antigens were recognized by TIL. It was observed that 3737-TIL were reactive to a mutated antigen present in TMG-1, but not TMG-2 or TMG-3 (Fig. 1A). Moreover, the reactivity predominated in the CD4+ T-cell population as demonstrated by up-regulation of the activation markers OX40 and 4-1BB (Tables 4A and 4B). Tables 4A and 4B show the percentage of 3737-TIL detected by flow cytometry as having the indicated phenotype following coculture with DCs cultured with the non-specific stimulator OKT3 or DCs transfected with green fluorescent protein (GFP) RNA, or the indicated tandem mini-gene (TMG) construct encoding the various mutations identified by whole-exomic sequencing. Mock-transfected cells were treated with transfection reagent only without addition of nucleic acid. Data were gated on live CD3+ cells.

TABLE 4A

	4-1BB-/CD4-	4-1BB+/CD4-	4-1BB-/CD4+	4-1BB+/CD4+
Mock	49	0	51	0
GFP	49	0	51	0
TMG-1	47	4	38	11
TMG-2	47	0	53	0
TMG-3	48	0	52	0
OKT3	4	41	23	32

TABLE 4B

	OX40-/CD4-	OX40+/CD4-	OX40-/CD4+	OX40+/CD4+
Mock	49	0	51	0
GFP	48	0	51	1
TMG-1	49	2	16	33
TMG-2	47	0	53	0
TMG-3	48	0	52	0
OKT3	38	6	11	45

[0110] Although some 4-1BB up-regulation was observed in the CD4-negative (CD8+) T-cell population, these cells were sorted and no reactivity against the TMG was found. To determine which of the 9 mutations in TMG-1 was being recognized by 3737-TIL, 9 additional TMG-1 constructs were synthesized, each one containing a reversion of one of the mutations back to the wild-type sequence. Reactivity of 3737-TIL to TMG-1 was abrogated only when the ERBB2 interacting protein (*ERBB2IP*) mutation was reverted back to the wild-type sequence, indicating that the TIL specifically recognized the ERBB2IP^{E805G} mutation (Fig. 1B).

[0111] The ERBB2IP-mutation reactive T-cell response was characterized molecularly. An IFN- γ ELISPOT assay was performed, and the results were measured at 20 hours. Patient 3737-TIL were co-cultured with DCs transfected with TMG-1 that had been pre-incubated with nothing, or the indicated HLA-blocking antibodies (against MHC-I, MHC-II, HLA-DP, HLA-DQ, or HLA-DR) (Fig. 2A). As controls for antibody blocking, the HLA-A2 restricted MART-reactive T cell DMF5 (Fig. 2B) and the HLA-DR-restricted tyrosinase-reactive T cell T4 (Fig. 2C) were co-cultured with the MART and tyrosinase-positive 624-CIITA melanoma cell line that had been pre-incubated with nothing, or the indicated HLA-blocking antibodies. The response of 3737-TIL was blocked by anti-HLA-DQ antibodies (Fig. 2A).

[0112] Another IFN- γ ELISPOT assay was performed, and the results were measured at 20 hours. Patient 3737-TIL were co-cultured with autologous B cells or allogeneic EBV-B cells partially matched at the HLA-DQ locus that had been pulsed overnight with DMSO, mutated (mut) ALK or mut ERBB2IP 25-AA long peptides (Fig. 2D).

[0113] Another IFN- γ ELISPOT assay was performed, and the results were measured at 20 hours. Patient 3737-TIL were co-cultured with autologous B cells that had been pulsed overnight with the mut ERBB2IP 25-AA peptide, or the indicated truncated mut ERBB2IP peptides (Fig. 2E).

[0114] As shown in Figs. 2A-2E, the 3737-TIL response was restricted by the HLA-DQB1*0601 allele and the minimal epitope was located within the 13 amino acid sequence NSKEETGHLENGN (SEQ ID NO: 29).

EXAMPLE 3

[0115] This example demonstrates that autologous open repertoire peripheral blood T cells genetically modified with the TCR-V β 22 chain of the ERBB2IP-specific CD4+ T-cells

identified in Example 2 matched with its α chain conferred specific reactivity to the mutated ERBB2IP peptide.

[0116] The clonality of the mutated ERBB2IP-specific CD4+ T-cells identified in Example 2 were characterized by sorting them after antigen-specific activation, using OX40 as a marker of activation. These cells were then bulk expanded and cloned by limiting dilution. A flow cytometry-based survey of the TCR-V β repertoire demonstrated that the bulk-expanded population was > 95% V β 22+, and that 10/11 clones assessed were purely V β 22+. TCR sequence analysis revealed the same TCR β V-D-J sequence in 6/6 V β 22+ clones tested (Table 5), showing that the majority of the ERBB2IP-mutation reactive T cells was comprised of a dominant V β 22+ T-cell clone.

TABLE 5

TCR V β	V-D-J nucleotide sequence (CDR3 underlined)	V-D-J amino acid sequence (CDR3 underlined)	Number of V β 22 (TRBV2) clones with indicated V-D-J
V β 22 (TRBV2)	GAACCTGAAGTCACCCAGACTCCCAGCCATCAGGT CACACAGATGGGACAGGAAGTGATCTTGCGCTGT GTCCCCATCTCTAATCACTTATACTTCTATTGGTACA GACAAATCTTGGGGCAGAAAGTCGAGTTTCTGGTT TCCTTTTATAATAATGAAATCTCAGAGAAGTCTGAA ATATTCGATGATCAATTCTCAGTTGAAAGGCCTGAT GGATCAAATTTCACTCTGAAGATCCGGTCCACAAA GCTGGAGGACTCAGCCATGTACTTCTGT <u>GCCAGC</u> <u>AGCCTGGGTGACAGGGGTAATGAAAACTGT</u> <u>TTTTGGCAGTGGAACCCAGCTCTGTCTTGG</u> (SEQ ID NO: 39)	EPEVTQTPSHQVTQMG QEVILRCVPISNHLYFYR QILGQKVEFLVSFYNNEIS EKSEIFDDQFSVERPDGS NFTLKIRSTKLEDSAMYF <u>CASSLGDRGNEKLFFGS</u> GTQLSVL (SEQ ID NO: 40)	6/6

[0117] T-cell clones expressing this V β 22 TCR specifically produced the cytokine IFN- γ upon stimulation with the mutated ERBB2IP peptide (Table 6). CD4+ V β 22+ clones were co-cultured for 6 hours with OKT3 or autologous B cells pulsed overnight with wild-type (wt) ERBB2IP, mutated (mut) ALK, or mut ERBB2IP 25-AA long peptides. Table 6 shows the percentage of CD4+ V β 22+ and V β 22- clones that produce intracellular IFN- γ (IFN- γ +) or do not produce intracellular IFN- γ (IFN- γ -) after co-culture as measured by flow cytometry. Data are representative of 2 clones sharing the same TCR-V β sequence.

TABLE 6

	IFN- γ -/V β 22-	IFN- γ +/V β 22-	IFN- γ -/V β 22+	IFN- γ +/V β 22+
mutALK	1	0	98	1
wtERBB2IP	1	0	99	0
mutERBB2IP	1	0	19	80
OKT3	3	4	59	34

[0118] Moreover, autologous open repertoire peripheral blood T cells genetically modified with this TCR-V β 22 chain matched with its α chain (Table 7) conferred specific reactivity to the mutated ERBB2IP peptide (Tables 8A and 8B), demonstrating that this TCR specifically recognized the ERBB2IP^{E805G} mutation. Autologous open-repertoire peripheral blood T cells were transduced (Td) with the TCR derived from the V β 22+ clone (Table 8A), or were treated with transduction reagent only without addition of nucleic acid (Mock) (Table 8B), and then assessed for reactivity as described for Table 6. The endogenous V β 22+ TCR constant regions were swapped with mouse constant regions, allowing for the detection of the introduced TCR using antibodies against the mouse TCR β constant region (mTCR β). Plate-bound OKT3 was used as a control in all assays. Tables 8A and 8B show the percentage of mTCR β + and mTCR β - cells that produce intracellular IFN- γ (IFN- γ +) or do not produce intracellular IFN- γ (IFN- γ -) as measured by flow cytometry.

TABLE 7

TCR V α	V-J nucleotide sequence (CDR3 underlined)	V-J amino acid sequence (CDR3 underlined)
TRAV26-2	GATGCTAAGACCACACAGCCAAATTC AATGGAG AGTAACGAAGAAGAGCCTGTTCACTGCCTTGTA ACCACTCCACAATCAGTGGA ACTGATTACATACA TTGGTATCGACAGCTTCCCTCCAGGGTCCAGAG TACGTGATTCATGGTCTTACAAGCAATGTGAACA ACAGAATGGCCTCTCTGGCAATCGCTGAAGACA GAAAGTCCAGTACCTTGATCCTGCACCGTGCTAC CTTGAGAGATGCTGCTGTG TACTACTGCATCCT <u>GAGACGTCTTAACGACTACAAGCTCAGCTTT</u> GGAGCCGGAACCACAGTAACTGTAAGAGCAA (SEQ ID NO: 41)	DAKTTQPNSMESNEE EEPVHLP CNHSTISGTDYIHWYRQLPSQ GPEYVIHGLTSNVNRRMA SLAIAEDRKSSTLILHRATLRDA AVYYC <u>ILRRLNDYKLS</u> FGAGT TVTVRA (SEQ ID NO: 42)

TABLE 8A

	IFN- γ -/mTCR β -	IFN- γ +/mTCR β -	IFN- γ -/mTCR β +	IFN- γ +/mTCR β +
mutALK	16	0	84	0
wtERBB2IP	15	0	85	0
mutERBB2IP	19	0	69	12
OKT3	14	2	74	10

TABLE 8B

	IFN- γ -/mTCR β -	IFN- γ +/mTCR β -	IFN- γ -/mTCR β +	IFN- γ +/mTCR β +
mutALK	99	1	0	0
wtERBB2IP	100	0	0	0
mutERBB2IP	100	0	0	0
OKT3	83	17	0	0

EXAMPLE 4

[0119] This example demonstrates a method of treating cancer using the autologous cells identified in Example 2.

[0120] Patient 3737 underwent adoptive transfer of 42.4 billion TIL containing CD4+ ERBB2IP mutation-reactive T cells followed by four doses of IL-2 to enhance T-cell proliferation and function. For treatment, Patient 3737 underwent a resection of lung lesions. Tumors were then minced into small fragments and incubated with high dose IL-2 to expand tumor infiltrating lymphocytes (TIL). After an initial expansion of the numbers of cells in IL-2, the numbers of select TIL cultures were further expanded for 2 weeks using a rapid expansion protocol (REP) including irradiated allogeneic peripheral blood feeder cells, OKT3 and IL-2. Prior to cell infusion, the patient was pre-conditioned with cyclophosphamide (CTX: 60 mg/kg, once a day for two days) followed by fludarabine (Flu: 25 mg/m² for 5 days). Patient 3737-TIL included 42.4 billion TIL containing over 10 billion (25%) ERBB2IP-mutation reactive T cells, and was administered on day 0, followed by IL-2 (Aldesleukin, 7.2e5 IU/kg) every 8 hours. The patient received a total of 4 doses of IL-2.

[0121] 3737-TIL were co-cultured with DCs transfected with TMG-1 or TMG-1 encoding the wild-type (wt) ERBB2IP reversion, and flow cytometry was used to assess OX40 and V β 22 expression on CD4+ T cells at 24 hours post-stimulation. Plate-bound OKT3 stimulation was used as a positive control. Flow cytometry analysis demonstrated that

approximately 25% of the entire 3737-TIL product administered was comprised of the V β 22+, mutation-reactive T cells (Fig. 3A, Table 9), equating to the infusion of over 10 billion ERBB2IP-mutation specific CD4+ T cells. Table 9 shows the percentage of V β 22+ and V β 22- cells that express OX40 (OX40+) or do not express OX40 (OX40-) as measured by flow cytometry.

[0122] An IFN- γ ELISA assay was performed on patient 3737 serum samples pre- and post-adoptive cell transfer of 3737-TIL. The results are shown in Fig. 3B. As shown in Fig. 3B, elevated levels of IFN- γ were detected in the patient's serum for the first five days after cell infusion.

[0123] Although the patient had clear evidence of progressive disease prior to the cell infusion, tumor regression was observed by the two month follow-up, and all target lung and liver lesions continued to regress, reaching a maximum reduction of 30% at 7 months post-treatment (Fig. 3C). The patient experienced disease stabilization for approximately 13 months after cell infusion, after which disease progression was observed only in the lungs but not liver.

TABLE 9

	V β 22-/OX40-	V β 22-/OX40+	V β 22+/OX40-	V β 22+/OX40+
TMG-1 wtERBB2IP	45	0	55	0
TMG-1	33	12	3	52
OKT3	19	31	6	44

EXAMPLE 5

[0124] This example demonstrates the *in vitro* phenotype and function of the cells of Example 4.

[0125] To determine whether there was evidence that the CD4+ ERBB2IP-mutation-reactive T cells played a role in the disease stabilization, the *in vitro* phenotype and function of the cells were evaluated. 3737-TIL were co-cultured for 6 hours with autologous B cells pulsed overnight with wild-type (wt) ERBB2IP, mutated (mut) ALK or mut ERBB2IP 25-AA long peptides. Flow cytometry was used to assess expression of V β 22 and to detect intracellular production of IFN- γ (Table 10A), tumor necrosis factor (TNF) (Table 10B), and IL-2 (Table 10C) in the CD4+ population. The percentage of cells having the indicated phenotypes is shown in Tables 10A-10C. Table 10D displays the percentage of V β 22+ cells

that expressed the indicated number of cytokines. It was found that the V β 22+ ERBB2IP-mutation reactive CD4+ T cells were polyfunctional Th1 cells, as stimulation with the mutated ERBB2IP peptide induced the robust co-expression of IFN- γ , TNF, and IL-2 (Tables 10A-10C), but little to no IL-4 or IL-17.

TABLE 10A

	V β 22-/IFN- γ -	V β 22-/IFN- γ +	V β 22+/IFN- γ -	V β 22+/IFN- γ +
mutALK	45	0	55	0
wtERBB2IP	44	0	56	0
mutERBB2IP	40	8	6	47
OKT3	29	33	24	14

TABLE 10B

	V β 22-/TNF-	V β 22-/TNF+	V β 22+/TNF-	V β 22+/TNF+
mutALK	45	0	55	0
wtERBB2IP	43	1	56	0
mutERBB2IP	37	10	3	50
OKT3	10	52	6	32

TABLE 10C

	V β 22-/IL-2-	V β 22-/IL-2+	V β 22+/IL-2-	V β 22+/IL-2+
mutALK	45	0	55	0
wtERBB2IP	43	1	56	0
mutERBB2IP	38	10	5	47
OKT3	27	36	23	14

TABLE 10D

No. cytokines (gated on V β 22+)	mutALK	wtERBB2IP	OKT3	mutERBB2IP
0	99%	98%	12%	11%
1+	1%	2%	30%	
2+	None	None	24%	
3+	None	None	34%	89%

[0126] Further phenotypic characterization revealed that these cells were predominantly effector memory CD4⁺ T cells with cytolytic potential (Tables 11 and 12). Patient 3737-TIL were assessed by flow cytometry for expression of V β 22 (representing ERBB2IP-mutation-reactive T cells) and the T-cell differentiation markers CD28, CD45RO, CD57, CCR7, CD127, CD62L, and CD27. Data were gated on live CD3⁺CD4⁺ cells. Positive and negative quadrant gates were set using isotype stained or unstained cells. The percentage of cells having the indicated phenotypes is shown in Table 11. Human peripheral blood cells (containing T cells of all differentiation stages) were included in experiments to ensure that the antibodies were working.

TABLE 11

V β 22-/CD28-	V β 22-/CD28+	V β 22+/CD28-	V β 22+/CD28+
1	56	1	42
V β 22-/CD45RO-	V β 22-/CD45RO+	V β 22+/CD45RO-	V β 22+/CD45RO+
0	57	0	43
V β 22-/CD57-	V β 22-/CD57+	V β 22+/CD57-	V β 22+/CD57+
48	9	42	1
V β 22-/CCR7-	V β 22-/CCR7+	V β 22+/CCR7-	V β 22+/CCR7+
57	0	43	0
V β 22-/CD127-	V β 22-/CD127+	V β 22+/CD127-	V β 22+/CD127+
25	32	21	22
V β 22-/CD62L-	V β 22-/CD62L+	V β 22+/CD62L-	V β 22+/CD62L+
49	8	42	1
V β 22-/CD27-	V β 22-/CD27+	V β 22+/CD27-	V β 22+/CD27+
57	0	43	0

[0127] Patient 3737-TIL were co-cultured for 6 hours with OKT3 or autologous B cells pulsed overnight with wild-type (wt) ERBB2IP, mutated (mut) ALK or mut ERBB2IP 25-AA long peptides. Antibodies specific for the degranulation marker CD107a were added at the beginning of the co-culture. Flow cytometry was used to assess expression of V β 22 and to detect cell surface mobilization of CD107a. Data were gated on the CD4⁺ population. The percentage of cells having the indicated phenotypes is shown in Table 12.

TABLE 12

	V β 22-/CD107a-	V β 22-/CD107a+	V β 22+/CD107a-	V β 22+/CD107a+
mutALK	51	0	48	1
wtERBB2IP	51	1	48	0
mutERBB2IP	53	6	19	22
OKT3	42	19	26	13

[0128] There also appeared to be a minor population of polyfunctional V β 22-negative, ERBB2IP-mutation-reactive CD4⁺ T cells present in 3737-TIL (Tables 9 and 10). These V β 22-negative cells were sorted by FACS and then rested in IL-2 containing media for 2 days prior to being co-cultured with autologous B cells pulsed overnight with wild-type (wt) ERBB2IP, mutated (mut) ALK or mut ERBB2IP 25-AA long peptides. Flow cytometry was used to assess expression of V β 22 and to detect intracellular production of IL-2 (Table 13C), TNF (Table 13B), and IFN- γ (Table 13A) in the CD4⁺ population at 6 hours (h) post-stimulation. The percentage of cells having the indicated phenotypes are shown in Tables 13A-13C.

TABLE 13A

	V β 22-/IFN- γ -	V β 22-/IFN- γ +	V β 22+/IFN- γ -	V β 22+/IFN- γ +
mutALK	99	0	1	0
wtERBB2IP	99	0	1	0
mutERBB2IP	85	14	0	1
OKT3	50	49	0	1

TABLE 13B

	V β 22-/TNF-	V β 22-/TNF+	V β 22+/TNF-	V β 22+/TNF+
mutALK	99	0	1	0
wtERBB2IP	97	2	1	0
mutERBB2IP	78	21	0	1
OKT3	9	90	0	1

TABLE 13C

	V β 22-/IL-2-	V β 22-/IL-2+	V β 22+/IL-2-	V β 22+/IL-2+
mutALK	99	0	1	0
wtERBB2IP	97	2	1	0
mutERBB2IP	78	21	0	1
OKT3	36	63	0	1

[0129] Flow cytometry was used to assess expression OX40 and V β 22 in the CD4⁺ population at 24 h post stimulation. Cells that upregulated OX40 were sorted and the numbers of the cells were expanded, and the TCR-V β repertoire was analyzed by flow cytometry. The results are shown in Fig. 3D. Sorting of the V β 22-negative cells followed by activation of these cells revealed that one or more additional clonotypes reactive to this epitope were present in 3737-TIL (Tables 13A-13C), the most dominant clonotype of which was V β 5.2 (Fig. 3D).

[0130] The sorted cells described in Fig. 3D were co-cultured for 6 h with autologous B cells pulsed overnight with wt ERBB2IP, mut ALK or mut ERBB2IP 25-AA long peptides. Flow cytometry was used to assess expression of V β 5.2 and to detect intracellular production of IL-2 (Table 14C), TNF (Table 14B), and IFN- γ (Table 14A) in the CD4⁺ population. Table 15 displays the percentage of V β 5.2⁺ cells that expressed the indicated number of cytokines.

TABLE 14A

	V β 5.2-/IFN- γ -	V β 5.2-/IFN- γ +	V β 5.2+/IFN- γ -	V β 5.2+/IFN- γ +
mutALK	51	0	49	0
wtERBB2IP	54	0	46	0
mutERBB2IP	42	13	20	25
OKT3	28	23	25	24

TABLE 14B

	V β 5.2-/TNF-	V β 5.2-/TNF+	V β 5.2+/TNF-	V β 5.2+/TNF+
mutALK	50	2	48	0
wtERBB2IP	52	2	46	0
mutERBB2IP	33	21	3	43
OKT3	5	46	3	46

TABLE 14C

	V β 5.2-/IL-2-	V β 5.2-/IL-2+	V β 5.2+/IL-2-	V β 5.2+/IL-2+
mutALK	51	1	48	0
wtERBB2IP	54	1	45	0
mutERBB2IP	38	17	14	31
OKT3	31	21	27	21

TABLE 15

No. cytokines (gated on V β 5.2+)	mutALK	wtERBB2IP	mutERBB2IP	OKT3
0	98%	98%	3%	6%
1+	2%	2%	11%	25%
2+	None	None	36%	36%
3+	None	None	50%	33%

[0131] V β 22-negative cells that upregulated OX40 upon stimulation with mutated ERBB2IP were sorted and the numbers of cells were expanded. RNA from these cells was then isolated and underwent rapid amplification of 5' complementary DNA ends (5'RACE) with TCR- β constant chain primers to identify the expressed TCR-V β sequences. TOPO-TA cloning was performed on the polymerase chain reaction (PCR) products and individual colonies were then sequenced. Flow cytometry demonstrated that 40-50% of these T cells were V β 5.2 (TRBV5-6). By Sanger sequencing, 3/7 TOPO-TA colonies were V β 5.2 (TRBV5-6) with the sequence shown in Table 16. Table 16 displays the most frequent TCR β V-D-J sequence of V β 22-negative ERBB2IP-mutation-reactive T cells.

TABLE 16

TCR V β	V-D-J nucleotide sequence (CDR3 underlined)	V-D-J amino acid sequence (CDR3 underlined)	Number of TOPO- TA clones with indicated V-D-J
V β 5.2 (TRBV5-6)	GACGCTGGAGTCACCCAAAGTCCCACACACCTGAT CAAAACGAGAGGACAGCAAGTGA CTCTGAGATGC TCTCCTAAGTCTGGGCATGACACTGTGCCTGGTAC CAACAGGCCCTGGGTCAGGGGCCCCAGTTTATCTT TCAGTATTATGAGGAGGAAGAGAGACAGAGAGGC AACTCCCTGATCGATTCTCAGGTCACCAGTTCCT AACTATAGCTCTGAGCTGAATGTGAACGCCTTGTT GCTGGGGGACTCGGCCCTCTATCTCTGTGCCAGCA <u>GCAAAGGCCCGGGAGGCAACTACGAGCAGTACTT</u> CGGGCCGGGCACCAGGCTCACGGTCACAG (SEQ ID NO: 43)	DAGVTQSPTHLIKTR GQQVTLRCSPKSGHD TVSWYQQALGQGPQ FIFQYEEEEERQRGNF PDRFSGHQFPNYSSE LNVNALLLGDALYLC <u>ASSKGPGGNYEQYFG</u> PGTRLTVT (SEQ ID NO: 44)	3/7

[0132] The majority of the V β 5.2+ cells produced multiple cytokines in an antigen-specific manner (Tables 14A-14C, 15, and 16). There also appeared to be a minor population of V β 5.2-negative (and V β 22-negative) CD4+ T cells that recognized mutated ERBB2IP (Tables 14A-14C and 15). Thus, the TIL used to treat patient 3737 contained at least three different polyfunctional CD4+ T-cell clones that recognized the same mutation in ERBB2IP, showing that this mutation was highly immunogenic.

EXAMPLE 6

[0133] This example demonstrates the *in vivo* persistence of the cells of Example 4.

[0134] To determine whether there was evidence that the CD4+ ERBB2IP-mutation-reactive T cells played a role in the disease stabilization, the *in vivo* persistence of the cells was evaluated. TCR-V β deep sequencing revealed that these clonotypes were rare or not detectable in the peripheral blood prior to ACT (Figs. 4A and 4B). Ten days after ACT, both clones were present at greater than 2% of the total T cells in the peripheral blood, but declined to less than 0.3% by day 34 post-cell infusion (Figs. 4A and 4B). Three lung metastases, which were resected nearly a year and a half after ACT, were infiltrated by the ERBB2IP-mutation-reactive T cells (Figs. 4A and 4B), showing that these cells contributed to the cancer regression and disease stabilization. The V β 22+ ERBB2IP-mutation-reactive clone was the most frequent clone detected in tumor nodule-3 (Tu-3-Post) and represented nearly 8% of total T cells in the tumor (Figs. 4A and 4B), whereas this clone was the second

and twelfth most frequent in tumor nodules-1 and -2, respectively. The V β 5.2+ ERBB2IP-mutation-reactive clone was also enriched compared to its frequency in blood in all three tumor nodules (Figs. 4A and 4B). Thus, patient 3737 experienced tumor regression with stabilization of disease for more than one year after receiving over 10 billion ERBB2IP-mutation-specific polyfunctional Th1 cells which infiltrated and persisted in the metastatic lesions.

[0135] Reverse transcriptase quantitative PCR (RT-qPCR) analysis of *ERBB2IP* expression in Patient 3737-TIL (T cells) and tumors pre-(Tu-Pre) and post adoptive cell transfer was performed. Three separate metastatic lung lesions (Tu-1, -2, -3-Post) were resected approximately 17 months post cell infusion. The results are shown in Fig. 4C, and are relative to *β -actin* (*ACTB*). A 350 base pair (bp) segment of the ERBB2IP gene containing the mutation was PCR-amplified from the cDNA samples described for Fig. 4C and Sanger sequenced. The location of the mutation was at nucleotide position 2414 of the coding sequence, corresponding to a change at position 805 of the amino acid sequence. Relatively high levels of *ERBB2IP* expression in both the original and recurrent lung lesions, as determined by quantitative RT-PCR, were observed (Fig. 4C), and Sanger sequencing validated the presence of the ERBB2IP mutation in all tumor lesions.

[0136] Immunohistochemistry analyses of T-cell infiltrates and MHC expression pre- and post-ACT were performed. Post-ACT tumors were harvested approximately 17 months after the first ACT. A positive control (tonsil) was included for all stains. The T-cell infiltrate and MHC expression of the tumors *in situ* are summarized in Tables 17 and 18, respectively.

TABLE 17

Tumor Nodule	CD3		CD8		CD4	
	Tumor	Stroma	Tumor	Stroma	Tumor	Stroma
Pre-1A	0-1	1	0-1	1	0-1	1
Pre-2A	0-1	1	0-1	1	0	0
Pre-3A	0	0-1	0	0-1	0	0
Pre-3B	0-1	1	0-1	0-1	0-1	1
Post-1A	1	1	1	1	0-1	1
Post-1B	1	2	1-2	2	1	2
Post-2A	0-1	1	0-1	1	0-1	0-1

0, no infiltrate
 1, rare to few
 2, moderately dense
 3, very dense

TABLE 18

Tumor Nodule	HLA-I	HLA-II (HLA-DR)
Pre-1A	1-2, >50%	0
Pre-2A	1-2, >50%	0
Pre-3A	1, >50%	0
Pre-3B	2, >50%	0
Post-1A	2-3, >50%	0
Post-1B	3, >50%	0
Post-2A	2, >50%	0

> 50% denotes greater than 50% of the tumor cells were positive.

0, negative
 1, weakly positive
 2, moderately positive
 3, strongly positive

EXAMPLE 7

[0137] This example demonstrates the contribution of mutation-reactive Th1 cells to the anti-tumor response of Example 4.

[0138] To specifically evaluate the contribution of mutation-reactive Th1 cells to the anti-tumor response *in vivo*, a TIL product that was comprised of > 95% of the V β 22+ ERBB2IP-mutation-reactive Th1 cells (about 120 billion mutation-reactive cells) was generated and adoptively transferred into patient 3737.

[0139] Flow cytometric analysis of the TIL-product used for re-treatment was performed. Table 19 shows that after gating on CD3, 97% were CD4+/CD8-, and of these, 98% were V β 22+ after further gating on CD4+ cells (Table 20).. Re-treatment TIL were co-cultured for

6 h with autologous B cells pulsed overnight with wild-type (wt) or mutated (mut) ERBB2IP 25-AA long peptides. Flow cytometry was used to detect intracellular TNF production in the CD4+ population (Table 20).

TABLE 19

CD8-/CD4-	CD8-/CD4+	CD8+/CD4-	CD8+/CD4+
0	97	3	0

TABLE 20

	V β 22-/TNF-	V β 22-/TNF+	V β 22+/TNF-	V β 22+/TNF+
wtERBB2IP	2	0	98	0
mutERBB2IP	1	3	3	93

[0140] Again, the patient experienced a decrease in target lesions, but unlike the first treatment, tumor regression was observed even at the first month follow-up and continued as of the follow-up at 4 months after the second treatment (Fig. 4D). Tumor regression was continuing as of the follow up at 8 months after the second treatment.

[0141] Six months after the second administration of mutation-reactive cells, computerized tomography (CT) scans of the lungs of Patient 3737 were taken, and the resulting images are shown in Figs. 7A-C. These images were compared to those taken prior to the second administration of mutation-reactive cells (Figs. 7D-7F). As shown in Figs. 7A-7F, an approximately 36% decrease in cancerous lesions was observed which provided a partial response (PR) by Response Evaluation Criteria In Solid Tumors (RECIST) criteria.

[0142] Eight months after the second administration of mutation-reactive cells, positron emission tomography (PET) scans of the liver and lungs of Patient 3737 were taken. It was observed that the target lesions continued to shrink. The radio-labeled glucose analogue, FDG (fluorodeoxyglucose), was administered to assess the uptake of glucose by the tumors in order to measure the metabolic activity of the tumors. The PET scans demonstrated no glucose uptake in 2 liver lesions and only some uptake in lung lesions.

EXAMPLES 8-10

[0143] The materials and methods for Examples 8-10 are set forth below.

Patient materials and cell lines

[0144] All patient materials were obtained in the course of a National Cancer Institute Institutional Review Board approved clinical trial. Patient 2359 and Patient 2591 were enrolled in clinical trials (Trial registration ID: NCT00096382 and ID: NCT00335127, respectively) that have been described in Dudley et al., *J. Clin. Oncol.*, 26: 5233-9 (2008). The patients underwent resections from which both a TIL line and a tumor cell line were established. TILs used for this study were generated by methods described in Dudley et al., *J. Immunother.*, 26: 332-42 (2003). Briefly, tumor fragments were excised and cultured in media containing IL-2. The numbers of TIL cultures that expanded were screened for recognition of autologous or HLA-matched tumor, and the numbers of reactive TILs were expanded using a rapid expansion protocol (REP) with IL-2, anti-CD3 antibody and irradiated feeder cells to large numbers for patient infusion (Riddell et al., *Science*, 257: 238-41 (1992)). A small portion of TILs underwent a second REP. For co-culture assays, T cells and tumor cells were cultured at 1:1 ratio in a 96-well plate with 200 μ L medium (AIM-V medium supplemented with 5% human serum) for 16 hours (hr).

[0145] To evaluate the antigen reactivity of TIL with clinical activity, two metastatic melanoma patients who experienced durable complete responses to adoptive TIL therapy were studied. Patient 2359 had a primary cutaneous melanoma at the right knee that metastasized to the thigh, iliac and inguinal lymph nodes. This individual experienced a complete regression of all metastatic lesions in response to autologous TIL transfer that was ongoing for over eight years following treatment. Patient 2591 had a primary back melanoma that metastasized to the abdominal wall, mesenteric lymph nodes, right colon, and supraclavicular lymph nodes. This individual experienced a complete regression of all metastatic lesions in response to autologous TIL transfer and remained disease free nine years after treatment.

Whole-exome sequencing

[0146] The method has been described in Robbins et al., *Nat. Med.*, 19: 747-52 (2013). Genomic DNA purification, library construction, exome capture of approximately 20,000 coding genes and next-generation sequencing of tumor and normal samples were performed at Personal Genome Diagnostics (Baltimore, MD). In brief, genomic DNA from tumor and normal samples was fragmented and used for Illumina TRUSEQ library construction

(Illumina, San Diego, CA). Exonic regions were captured in solution using the Agilent SURESELECT 50 Mb kit (version 3) according to the manufacturer's instructions (Agilent, Santa Clara, CA). Paired-end sequencing, resulting in 100 bases from each end of each fragment, was performed using a HISEQ 2000 Genome Analyzer (Illumina). Sequence data were mapped to the reference human genome sequence, and sequence alterations were determined by comparison of over 50 million bases of tumor and normal DNA. Over 8 billion bases of sequence data were obtained for each sample, and a high fraction of the bases were from the captured coding regions. Over 43 million bases of target DNA were analyzed in the tumor and normal samples, and an average of 42–51 reads were obtained at each base in the normal and tumor DNA samples.

[0147] Bioinformatic analyses were carried out by Personal Genome Diagnostics and the Genome Technology Access Center, Genomics and Pathology Services of the Washington University School of Medicine. The tags were aligned to the human genome reference sequence (hg18) using the ELAND algorithm of the CASAVA 1.6 software (Illumina). The chastity filter of the BASECALL software of Illumina was used to select sequence reads for subsequent analyses. The ELANDv2 algorithm of the CASAVA 1.6 software was then applied to identify point mutations and small insertions and deletions. Known polymorphisms recorded in dbSNP were removed from the analysis. Potential somatic mutations were filtered and visually inspected as described in Jones et al., *Science*, 330: 228-31 (2010).

The construction of tandem minigene library

[0148] Non-synonymous mutations from melanoma samples were identified from whole-exome sequencing data. Tandem minigene constructs that encode polypeptides containing 6 identified mutated amino acid residues flanked on their N- and C- termini, 12 amino acids on both sides, were synthesized (Integrated DNA Technologies, Coralville, Iowa), and then cloned into pcDNA3.1 expression vector using the IN-FUSION Advantage PCR Cloning Kit (Clontech), according to the manufacturer's instructions.

IFN- γ ELISPOT assay

[0149] The responses directed against tumor cell lines and peptide-pulsed target cells were quantified in an IFN- γ ELISPOT assay using 96-well PVDF-membrane filter plates

(EMD Millipore, Billerica, MA) coated with 15 $\mu\text{g/ml}$ of the monoclonal anti-IFN- γ antibody 1D1K (Mabtech, Inc., Cincinnati, OH). Bound cytokine was detected using 1 $\mu\text{g/ml}$ of the biotinylated anti-IFN- γ antibody 7-B6-1 (Mabtech). HEK293 cells expressing HLA-A*0201, HLA-A*0205 or HLA-C*0701 were pulsed with peptides for 2 h at 37 °C. The following peptides were used: MART-1: AAGIGILTV (SEQ ID NO: 54), mutated KIF2C: RLFPGTLTIKI (SEQ ID NO: 55), mutated POLA2: TRSSGSHFVF (SEQ ID NO: 56). T cells were co-cultured overnight with target cells or medium containing 50 ng/ml PMA plus 1 μM ionomycin (PMA/I). The numbers of spots per 10^5 T cells were calculated.

EXAMPLE 8

[0150] This example demonstrates that TIL 2359 recognize a mutated antigen as assessed by minigene library screening.

[0151] The reactivity of TIL 2359 was evaluated using TMG constructs that were generated based on the non-synonymous mutations identified by exomic analysis of tumor and normal DNA. Each TMG construct encoded up to six individual minigene fragments that corresponded to the mutated codon flanked on either side by the 12 additional codons present in the normal gene product. One example is illustrated in Fig. 5A.

[0152] COS-7 cells were transiently transfected individually with one of twelve tandem minigenes encoding the 71 minigenes based on exomic DNA sequences containing non-synonymous point mutations identified from Mel 2359. These COS-7 cells were also co-transfected with HLA-A*0205, the dominant HLA restriction element used for autologous tumor cell recognition by this TIL. Co-culture of these transfectants with TIL 2359 resulted in the recognition of one of the 12 TMG constructs, RJ-1 (Fig. 5B). RJ-1 encoded mutated fragments of the *EPHB2*, *KIF2C*, *SLC44A5*, *ABCA4*, *DENND4B*, and *EPRS* genes, as shown in Fig. 5A. Subsequently, six RJ-1 variant constructs were generated, each of which encoded the WT rather than the mutated residue present in one of the six minigenes (Fig. 5C). TIL 2359 recognized COS-7 cells co-transfected with HLA-A*0205 plus five of the six individually transfected RJ-1 variants, but failed to recognize the variant encoding the WT *KIF2C* sequence, indicating that this minigene encoded a mutated epitope recognized by TIL 2359 (Fig. 5C). To further test this observation, COS-7 cells were co-transfected with either WT or mutated full-length *KIF2C* cDNA transcripts that were amplified from Mel 2359, together with either HLA-A*0101, HLA-A*0201 or HLA-A*0205 cDNA. The co-culture

experiment indicated that TIL 2359 T cells recognized COS-7 cells co-transfected with the mutated but not WT *KIF2C* gene product, in a HLA-A*0205-restricted manner (Fig. 5D).

[0153] The mutated *KIF2C* coding region contained a single C to A transversion at nucleotide 46 that resulted in a substitution of threonine for alanine at position 16 in the native *KIF2C* protein. Exomic sequencing results indicated that DNA from Mel 2359 exclusively corresponded to the mutated but not the normal residue at position 46, results confirmed by direct Sanger sequencing of Mel 2359 DNA, indicating the loss of heterozygosity at this locus. In an attempt to identify the mutated *KIF2C* epitope recognized by TIL 2359, peptides encompassing the *KIF2C* mutation that were predicted to bind with high affinity to HLA-A*0205 were synthesized (Hoof et al., *Immunogenetics*, 61: 1-13 (2009)), and pulsed on HEK293 cells that stably expressed HLA-A*0205 (Table 21). HEK293-A*0205 cells pulsed with a decamer corresponding to residues 10-19 stimulated the release of high levels of IFN- γ from TIL 2359 T cells, and the peptide was recognized at a minimum concentration of 0.1 nM. In contrast, the corresponding WT peptide did not induce significant IFN- γ release at a concentration as high as 10 μ M (Fig. 5E).

TABLE 21

Amino acid position	Mutated Peptide	Predicted HLA-A*0205 binding affinity (nM)	Co-culture result [IFN- γ (pg/mL)]
10-19	RLFPGLTIKI (SEQ ID NO: 59)	55.21	10690
10-17	RLFPGLTI (SEQ ID NO: 60)	132.35	121.5
15-25	LTIKIQRSNGL (SEQ ID NO: 61)	251.33	31.5
7-17	LQARLFPGLTI (SEQ ID NO: 62)	293.83	27
7-16	LQARLFPGLT (SEQ ID NO: 63)	1549.33	24

EXAMPLE 9

[0154] This example demonstrates that TIL 2591 recognize a mutated antigen identified by minigene library screening.

[0155] The mutated T-cell antigen recognized by TIL 2591 was identified by synthesizing 37 TMG constructs encoding the 217 minigenes based on exomic DNA sequences containing non-synonymous point mutations identified from Mel 2591. TIL 2591 recognized autologous tumor cells in the context of multiple HLA restriction elements. Therefore, HEK293 cell lines stably expressing each of the six MHC class I HLA molecules isolated from Mel 2591 were transiently transfected individually with the 37 TMG constructs, followed by an overnight co-culture with TIL 2591. Initial results indicated that TIL 2591 recognized HLA-C*0701⁺ HEK293 cells (HEK293-C*0701) cells that were transiently transfected with minigene DW-6, but failed to respond significantly to the other minigene constructs (Fig. 6A). Each of the six individual mutated minigenes in the DW-6 tandem construct (Fig. 6B) were then individually reverted to the WT sequence (Fig. 6C). Evaluation of responses to the WT variants indicated that TIL 2591 recognized COS-7 cells transfected with each of the DW-6 variants, with the exception of a construct encoding the WT *POLA2* fragment (Fig. 6C). To test these findings, COS-7 cells were transfected with either a WT or mutated full-length *POLA2* cDNA construct, together with HLA-C*0401, HLA-C*0701 or HLA-C*0702 cDNA. TIL 2591 T cells only recognized target cells transfected with HLA-C*0701 plus the mutated *POLA2* construct, but not the corresponding WT transcript (Fig. 6D). The single C to T transition at nucleotide 1258 of the *POLA2* coding region resulted in a substitution of leucine for phenylalanine at position 420 of the WT *POLA2* protein. Sanger sequencing indicated that both genomic DNA and cDNA derived from Mel 2591 RNA contained both the WT and mutated nucleotide at position 1258, whereas genomic DNA isolated from PBMC of patient 2591 corresponded to the WT sequence, indicating that this represented a heterozygous somatic mutation in Mel 2591 cells.

[0156] An HLA-C*0701 binding algorithm was then used to identify candidate *POLA2* peptides overlapping with the mutated leucine residue at position 420 (Table 22). Co-culture results indicated that HLA-C*0701⁺ HEK293 cells pulsed with a decamer corresponding to residues 413-422 of mutated *POLA2* stimulated the release of IFN- γ from TIL 2591 T cells at a minimum concentration of 10 nM. In contrast, the corresponding WT peptide did not induce significant IFN- γ release at a concentration as high as 10 μ M (Fig. 6E).

TABLE 22

Amino acid position	Mutated Peptide	Predicted HLA-C*0701 binding affinity (nM)	Co-culture result [IFN- γ (pg/mL)]
413-422	TRSSGSHFVF (SEQ ID NO: 68)	147.35	1106
413-423	TRSSGSHFVFV (SEQ ID NO: 69)	280.38	50
413-421	TRSSGSHFV (SEQ ID NO: 70)	285.90	60
413-420	TRSSGSHF (SEQ ID NO: 71)	518.82	48
420-429	FVFPVPSLRDV (SEQ ID NO: 72)	599.44	39

[0157] The proportion of T cells in TIL 2359 and 2591 recognizing the mutated KIF2C and POLA2, respectively, was then estimated using IFN- γ enzyme-linked immunosorbent spot (ELISPOT) assays. TIL 2359 generated approximately 2,000 spots per 100,000 T cells in response to HLA-A*0205⁺ cells pulsed with the mutated KIF2C epitope, similar to that observed in response to the autologous melanoma (Table 23). TIL 2591 generated greater than 7,000 spots in response to the HLA-A2 restricted MART-1 epitope, while only small fractions of T cells reacted against the HLA-C*0701-restricted mutated POLA2 epitope (Table 23).

TABLE 23

	TIL 2359 Spots per 1 x 10⁵ cells
Mel 2359	1698
293-A*0205	189
293-A*0205 + KIF2Cmut	2057
	TIL 2591 Spots per 1 x 10⁵ cells
Mel 2591	11344
293-A*02	999
293-A*02 + MART-1	7404
293-C*0701	906
293-C*0701 + POLA2 mut	1280

EXAMPLE 10

[0158] This example demonstrates a method of identifying T cells reactive against a mutated antigen present in gastrointestinal (GI) cancer identified by minigene library screening.

[0159] Whole-exome sequencing was performed on metastatic lesions from GI cancer patients to identify mutations. Next, minigene constructs that encoded each mutation were generated and transfected into autologous APCs to allow for the processing and presentation of all the mutations expressed by the tumor. These APCs were then co-cultured with tumor infiltrating lymphocytes (TIL) and T-cell reactivity against the mutations was determined by IFN- γ ELISPOT and 4-1BB and OX40 upregulation by flow cytometry.

[0160] In one patient with colon cancer, 119 mutations were evaluated for mutation-reactivity. Several, but not all, TIL cultures were found to contain highly variable proportions of CD8⁺ T cells that specifically recognized a mutation in CASP8 (67 F \rightarrow V). Upon further expansion *in vitro*, these mutation-reactive CD8⁺ T cells were markedly outgrown by other cells in culture. Administration of 40.3 x 10⁹ TIL, which was estimated to contain about 0.31% (approximately 127 million) mutation-reactive cells, to the patient did not result in a clinical response at the first follow-up approximately six weeks after administration of cells. The patient died about six weeks later. Without being bound to a particular theory or mechanism, it is believed that any one or more of the very late stage of the disease prior to treatment, the patient's poor overall condition, and the patient's poor

tolerance of the lymphodepleting chemotherapy administered prior to adoptive cell therapy may have been contributing factors in the patient's death. A TCR that was reactive against mutated CASP8 was isolated from the TIL, and T cells transduced to express the TCR were reactive against DCs pulsed with mutated CASP8.

[0161] In another patient with rectal cancer, 155 mutations were evaluated for mutation-reactivity. At least 3 different mutation-reactivities were found, two comprising CD8+ T-cell responses and one CD4+ response. Administration of mutation-reactive TIL to the patient initially resulted in a mixed response at approximately 1.5 months after treatment, but the patient later developed progressive disease at approximately 3.5 months after treatment. A potentially mutation-reactive TCR was isolated from the CD4+ TIL and from the CD8+ TIL.

[0162] In a third patient (cholangiocarcinoma), T cells reactive against 38 mutations tested were not detected. For this patient, the "mutation call" threshold was lowered, and an additional 125 putative mutations will be evaluated. The "mutation call" is an arbitrarily set threshold at which a sequence is identified as a mutation using bioinformatics. In this case, as a first pass, the threshold was relatively high (for example, providing a high level of confidence that the mutations identified were true mutations). The threshold was then lowered, providing a lower level of confidence that the mutations identified were true mutations, however, the possibility that the mutations identified were true mutations remained.

[0163] These data show that the ability of the human immune system to mount a T-cell response against somatic mutations in metastatic GI cancers may not be a rare event. The study is ongoing.

[0164] All references, including publications, patent applications, and patents, cited herein are hereby incorporated by reference to the same extent as if each reference were individually and specifically indicated to be incorporated by reference and were set forth in its entirety herein.

[0165] The use of the terms "a" and "an" and "the" and "at least one" and similar referents in the context of describing the invention (especially in the context of the following claims) are to be construed to cover both the singular and the plural, unless otherwise indicated herein or clearly contradicted by context. The use of the term "at least one" followed by a list of one or more items (for example, "at least one of A and B") is to be

construed to mean one item selected from the listed items (A or B) or any combination of two or more of the listed items (A and B), unless otherwise indicated herein or clearly contradicted by context. The terms “comprising,” “having,” “including,” and “containing” are to be construed as open-ended terms (i.e., meaning “including, but not limited to,”) unless otherwise noted. Recitation of ranges of values herein are merely intended to serve as a shorthand method of referring individually to each separate value falling within the range, unless otherwise indicated herein, and each separate value is incorporated into the specification as if it were individually recited herein. All methods described herein can be performed in any suitable order unless otherwise indicated herein or otherwise clearly contradicted by context. The use of any and all examples, or exemplary language (e.g., “such as”) provided herein, is intended merely to better illuminate the invention and does not pose a limitation on the scope of the invention unless otherwise claimed. No language in the specification should be construed as indicating any non-claimed element as essential to the practice of the invention.

[0166] Preferred embodiments of this invention are described herein, including the best mode known to the inventors for carrying out the invention. Variations of those preferred embodiments may become apparent to those of ordinary skill in the art upon reading the foregoing description. The inventors expect skilled artisans to employ such variations as appropriate, and the inventors intend for the invention to be practiced otherwise than as specifically described herein. Accordingly, this invention includes all modifications and equivalents of the subject matter recited in the claims appended hereto as permitted by applicable law. Moreover, any combination of the above-described elements in all possible variations thereof is encompassed by the invention unless otherwise indicated herein or otherwise clearly contradicted by context.

[0167] This invention was made with Government support under project number ZIABC010984 by the National Institutes of Health, National Cancer Institute. The Government has certain rights in the invention.

CLAIMS

1. A method of isolating a T cell receptor (TCR), or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising:

identifying one or more genes in the nucleic acid of a cancer cell of a patient, each gene containing a cancer-specific mutation that encodes a mutated amino acid sequence, wherein the cancer cell is obtained from blood, primary tumor, or tumor metastasis of the patient;

inducing autologous antigen presenting cells (APCs) of the patient to present the mutated amino acid sequence;

co-culturing autologous T cells of the patient with the autologous APCs that present the mutated amino acid sequence;

selecting the autologous T cells that (a) were co-cultured with the autologous APCs that present the mutated amino acid sequence and (b) have antigenic specificity for the mutated amino acid sequence presented in the context of a major histocompatibility complex (MHC) molecule expressed by the patient; and

isolating a nucleotide sequence that encodes the TCR, or the antigen-binding portion thereof, from the selected autologous T cells, wherein the TCR, or the antigen-binding portion thereof, has antigenic specificity for the mutated amino acid sequence encoded by the cancer-specific mutation.

2. The method of claim 1, wherein inducing autologous APCs of the patient to present the mutated amino acid sequence comprises pulsing APCs with peptides comprising the mutated amino acid sequence or a pool of peptides, each peptide in the pool comprising a different mutated amino acid sequence.

3. The method of claim 1, wherein inducing autologous APCs of the patient to present the mutated amino acid sequence comprises introducing a nucleotide sequence encoding the mutated amino acid sequence into the APCs.

4. The method of claim 3, wherein the nucleotide sequence introduced into the autologous APCs is a tandem minigene (TMG) construct, each minigene comprising a different

gene, each gene including a cancer-specific mutation that encodes a mutated amino acid sequence.

5. The method of any one of claims 1-4, further comprising obtaining multiple fragments of a tumor from the patient, separately co-culturing autologous T cells from each of the multiple fragments with the autologous APCs that present the mutated amino acid sequence, and separately assessing the T cells from each of the multiple fragments for antigenic specificity for the mutated amino acid sequence.

6. The method of any one of claims 1-5, wherein selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selectively growing the autologous T cells that have antigenic specificity for the mutated amino acid sequence.

7. The method of any one of claims 1-6, wherein selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selecting the T cells that express any one or more of programmed cell death 1 (PD-1), lymphocyte-activation gene 3 (LAG-3), T cell immunoglobulin and mucin domain 3 (TIM-3), 4-1BB, OX40, and CD107a.

8. The method of any one of claims 1-7, wherein selecting the autologous T cells that have antigenic specificity for the mutated amino acid sequence comprises selecting the T cells (i) that secrete a greater amount of one or more cytokines upon co-culture with APCs that present the mutated amino acid sequence as compared to the amount of the one or more cytokines secreted by a negative control or (ii) in which at least twice as many of the numbers of T cells secrete one or more cytokines upon co-culture with APCs that present the mutated amino acid sequence as compared to the numbers of negative control T cells that secrete the one or more cytokines.

9. The method of claim 8, wherein the one or more cytokines comprise interferon (IFN)- γ , interleukin (IL)-2, tumor necrosis factor alpha (TNF- α), granulocyte/monocyte colony stimulating factor (GM-CSF), IL-4, IL-5, IL-9, IL-10, IL-17, and IL-22.

10. The method of any one of claims 1-9, wherein identifying one or more genes in the nucleic acid of the cancer cell comprises sequencing the whole exome, the whole genome, or the whole transcriptome of the cancer cell.

11. The method of any one of claims 1-10, wherein the cancer cell is obtained from the blood of the patient.

12. The method of any one of claims 1-10, the cancer cell is obtained from the primary tumor of the patient.

13. The method of any one of claims 1-10, the cancer cell is obtained from the tumor metastasis of the patient.

14. A method of preparing a population of cells that express a TCR, or an antigen-binding portion thereof, having antigenic specificity for a mutated amino acid sequence encoded by a cancer-specific mutation, the method comprising:

isolating a TCR, or an antigen-binding portion thereof, according to the method of any one of claims 1-13, and

introducing the nucleotide sequence encoding the isolated TCR, or the antigen-binding portion thereof, into peripheral blood mononuclear cells (PBMC) to obtain cells that express the TCR, or the antigen-binding portion thereof.

15. The method of claim 14, further comprising expanding the numbers of PBMC that express the TCR, or the antigen-binding portion thereof.

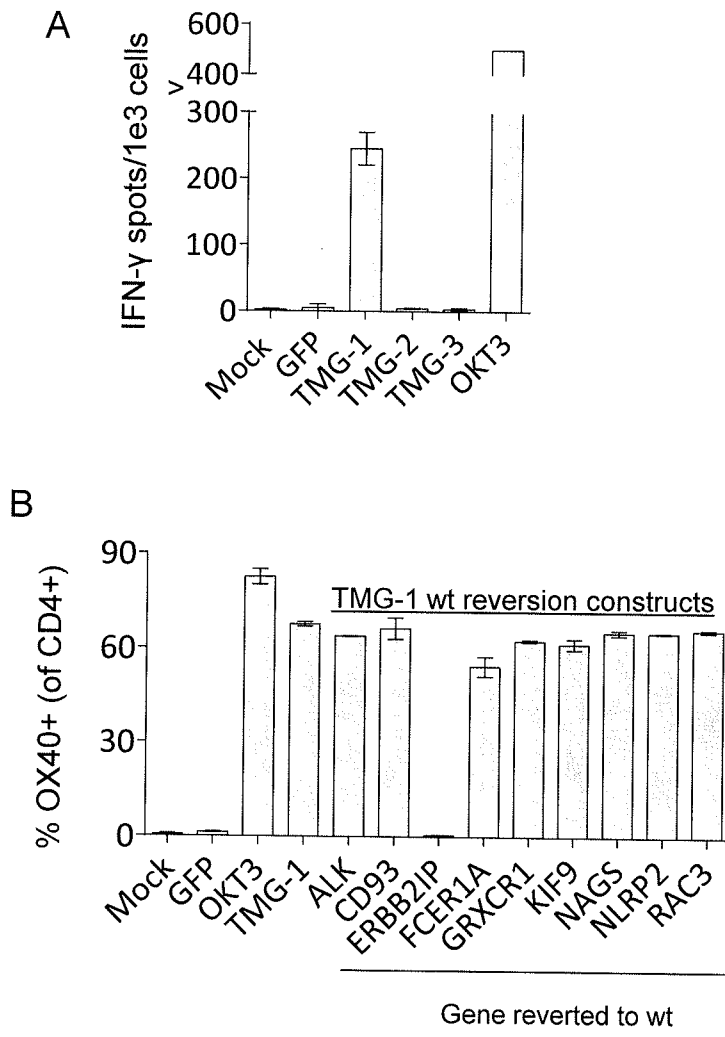


FIG. 1

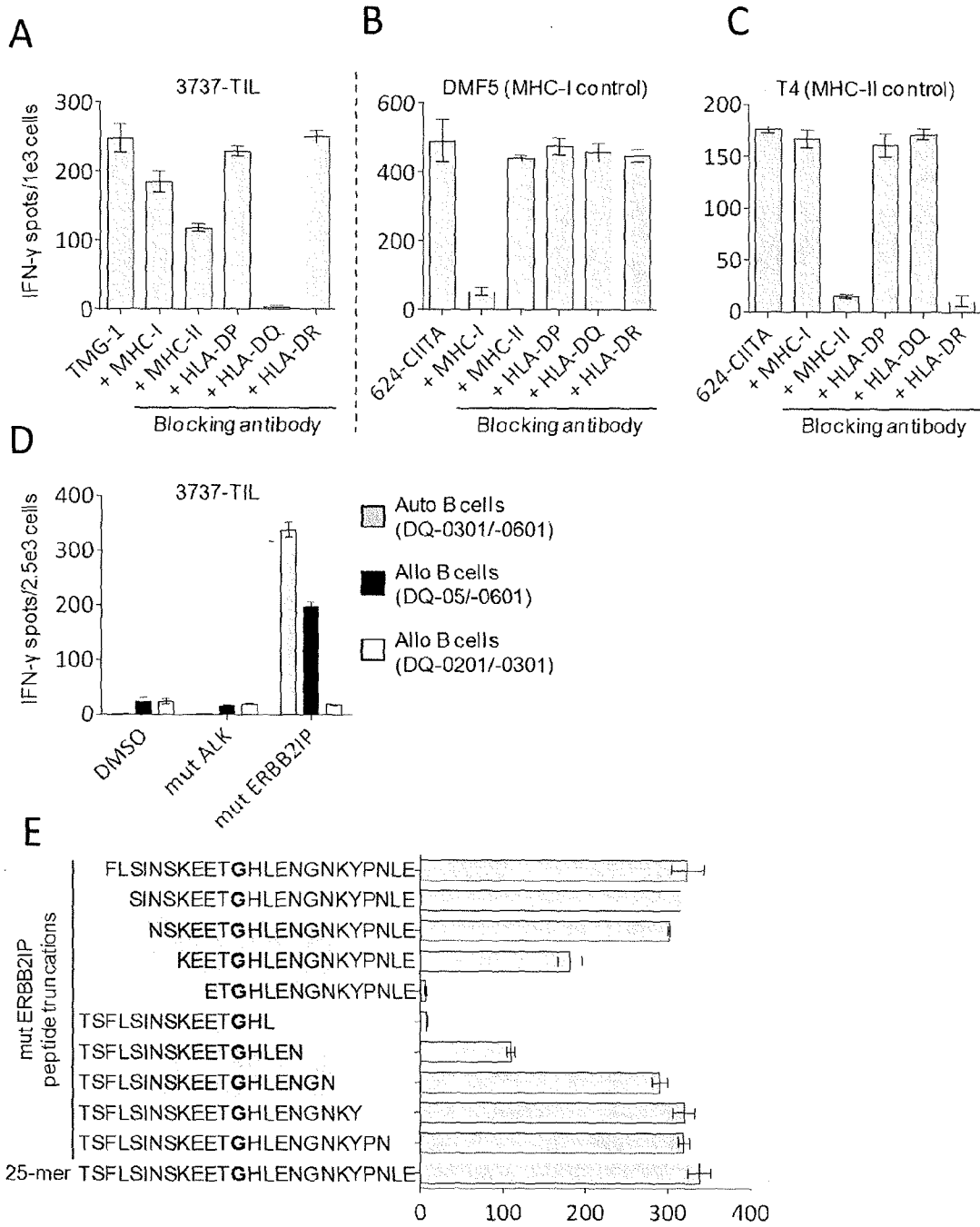


FIG. 2

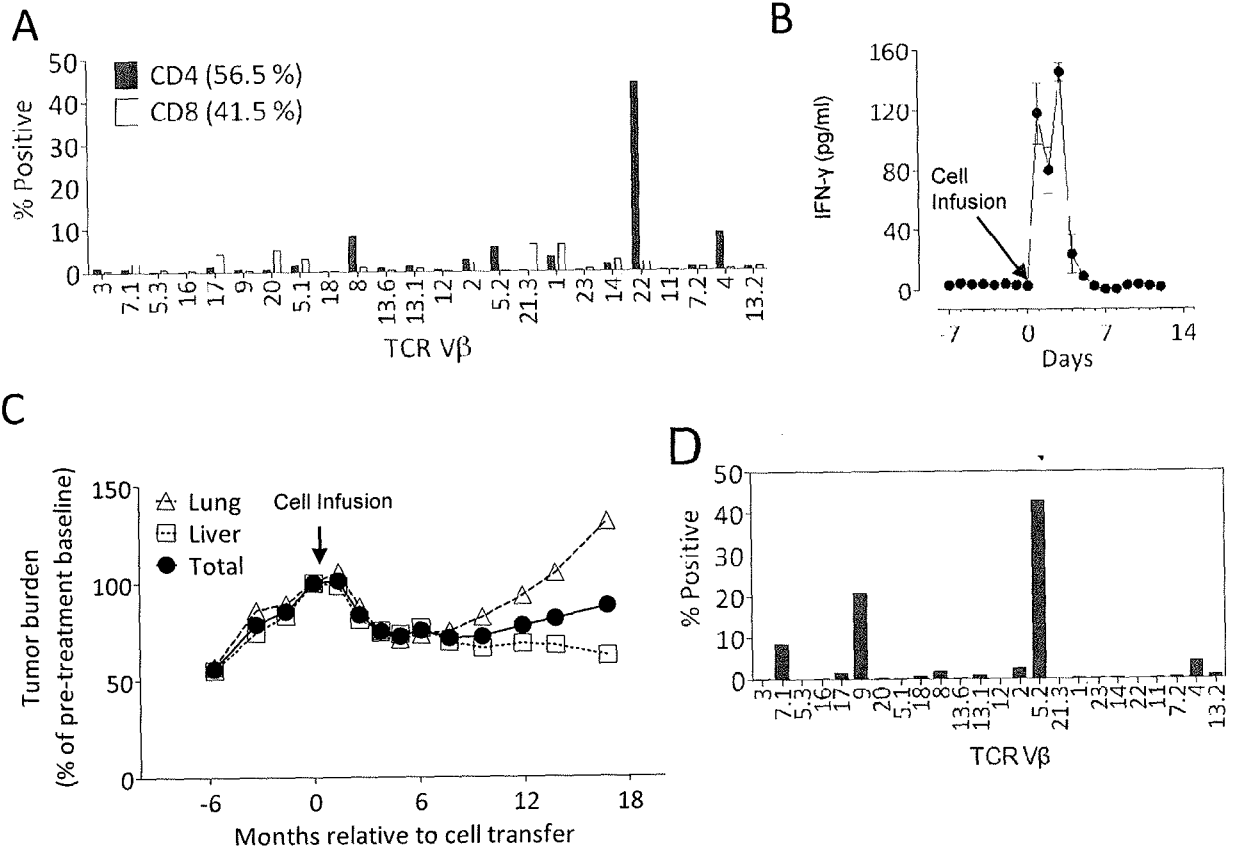


FIG. 3

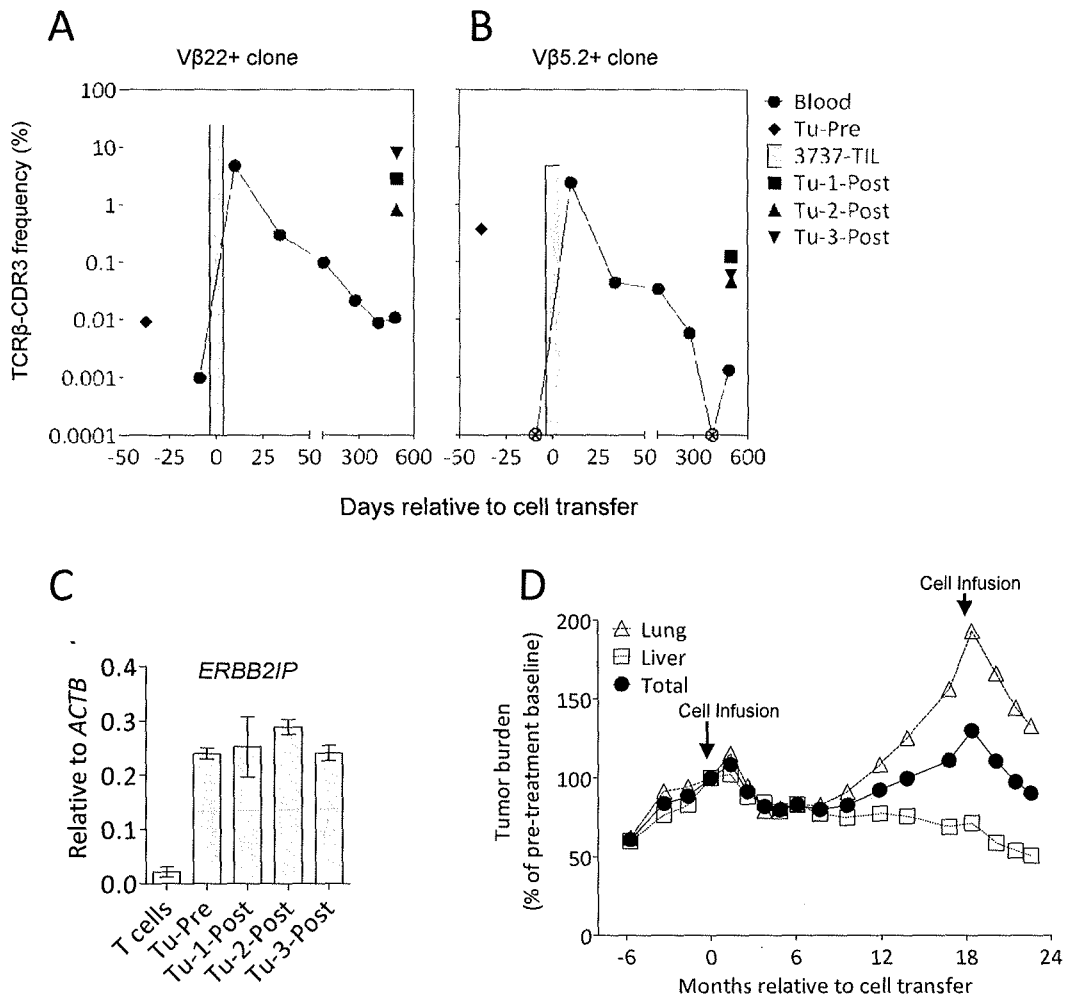


FIG. 4

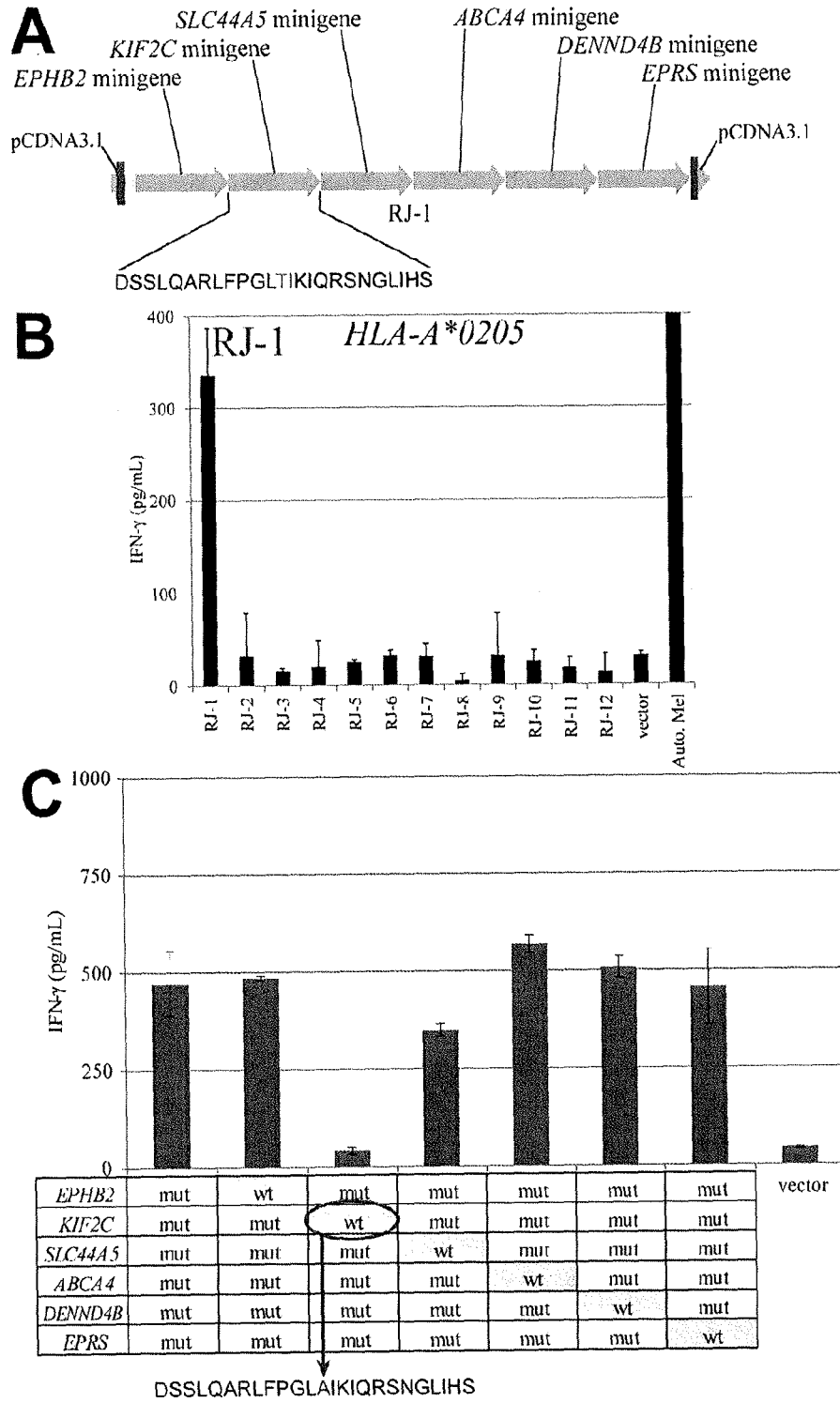


FIG. 5A-5C

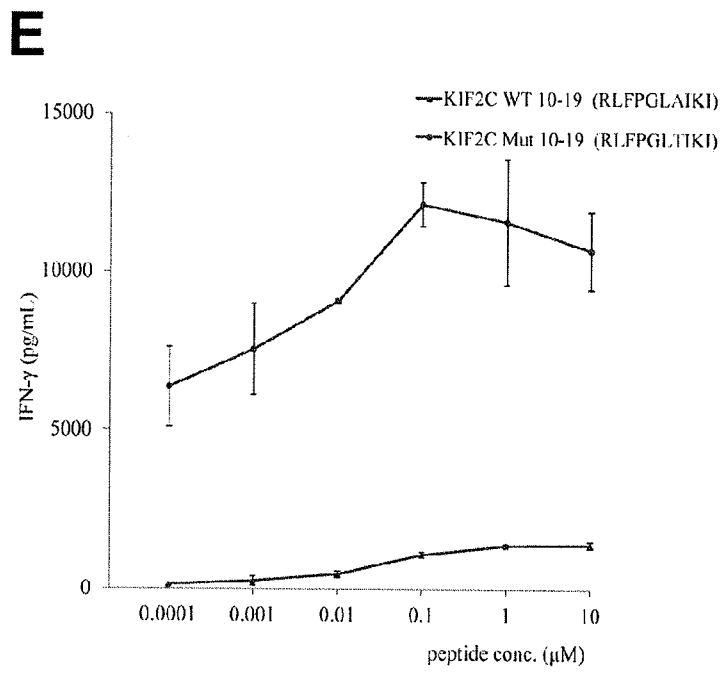
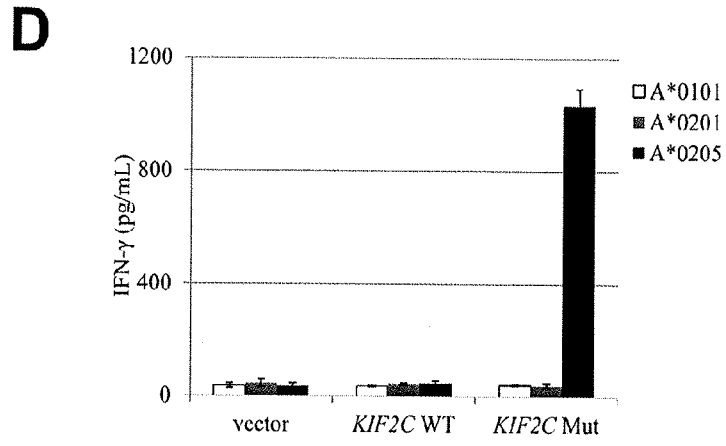


FIG. 5D-5E

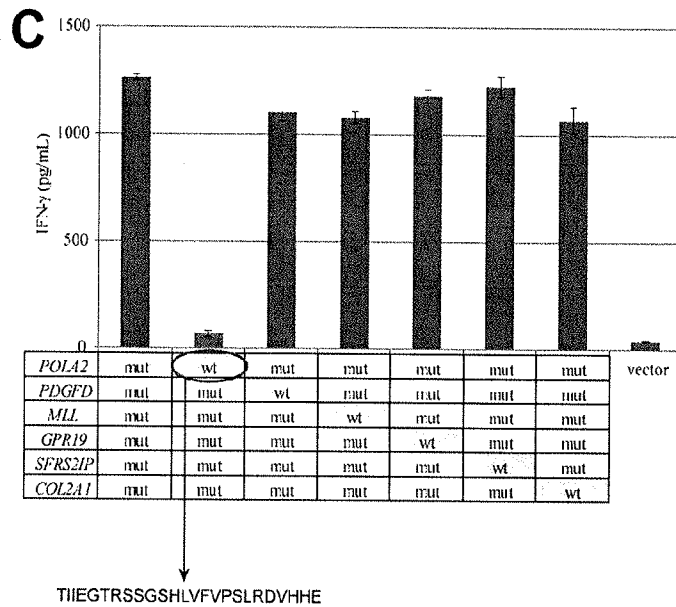
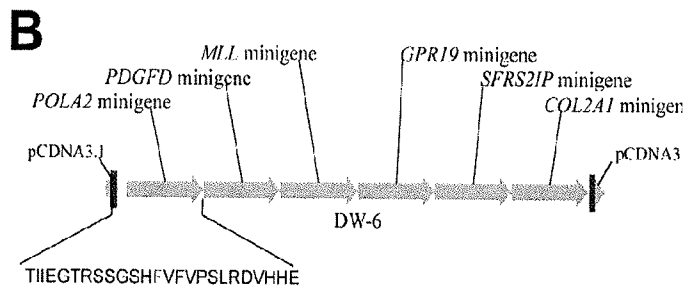
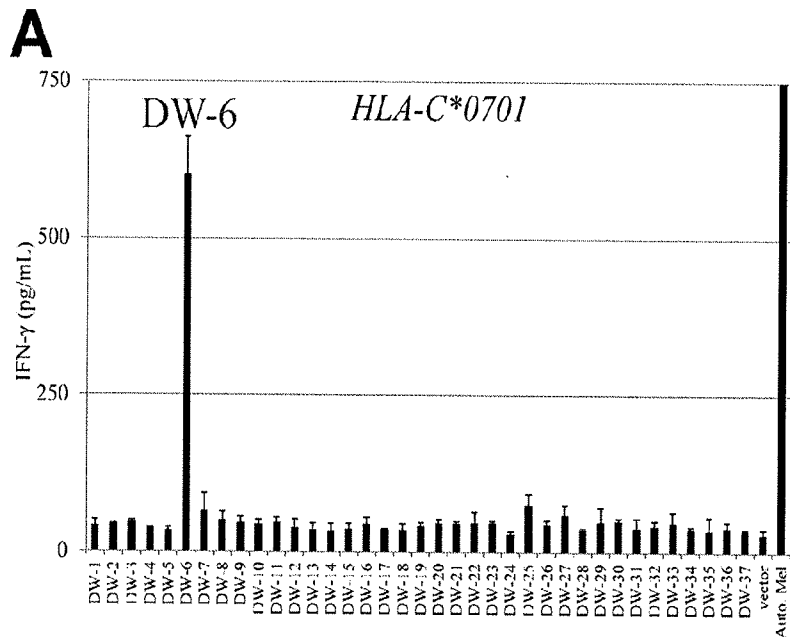


FIG. 6A-6C

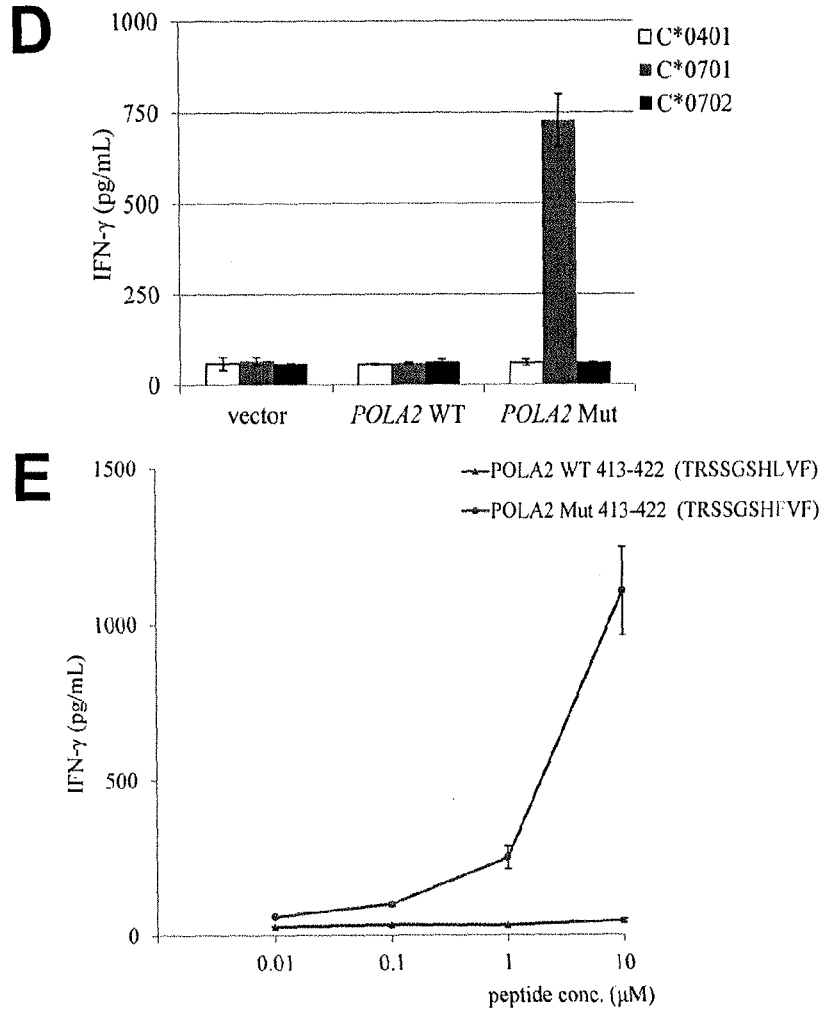


FIG. 6D-6E

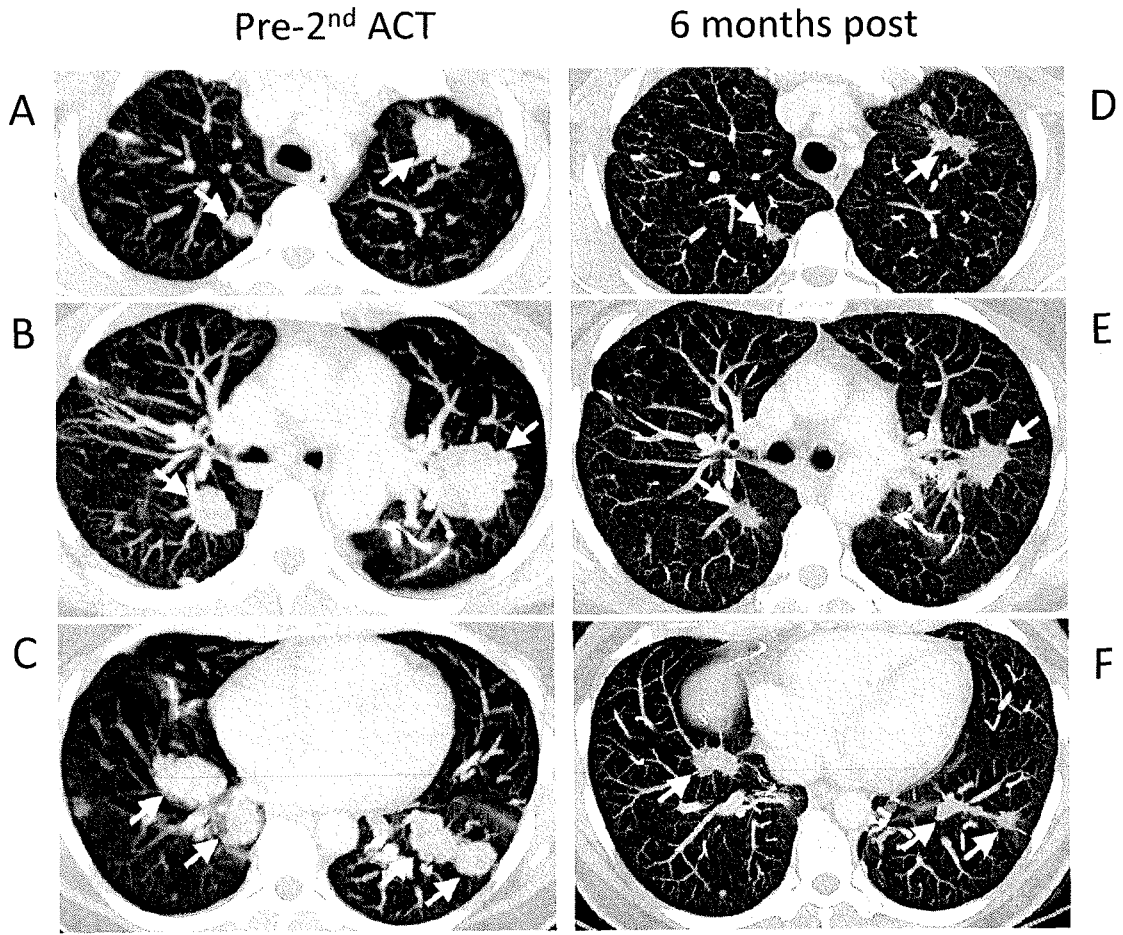


FIG. 7

Sequence Listing

2023282185 18 Dec 2023

1	Sequence Listing Information	
1-1	File Name	34707AU3 ST26 Sequence Listing.xml
1-2	DTD Version	V1_3
1-3	Software Name	WIPO Sequence
1-4	Software Version	2.3.0
1-5	Production Date	2023-12-14
1-6	Original free text language code	
1-7	Non English free text language code	
2	General Information	
2-1	Current application: IP Office	AU
2-2	Current application: Application number	2023282185
2-3	Current application: Filing date	2023-12-12
2-4	Current application: Applicant file reference	34707AU3
2-5	Earliest priority application: IP Office	WO
2-6	Earliest priority application: Application number	PCT/US2014/058796
2-7	Earliest priority application: Filing date	2014-10-02
2-8en	Applicant name	United States of America, as represented by the Secretary, Department of Health and Human Services, C/O National Institute of Health, Office of Technology Transfer
2-8	Applicant name: Name Latin	
2-9en	Inventor name	TRAN, Eric
2-9	Inventor name: Name Latin	
2-10en	Invention title	Methods of isolating T cell receptors having antigenic specificity for a cancer specific mutation
2-11	Sequence Total Quantity	73

3-1	Sequences		
3-1-1	Sequence Number [ID]	1	
3-1-2	Molecule Type	AA	
3-1-3	Length	25	
3-1-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-1-5	Residues	RVLKGGSVRK LRHAKQLVLE LGEEA	25
3-2	Sequences		
3-2-1	Sequence Number [ID]	2	
3-2-2	Molecule Type	AA	
3-2-3	Length	25	
3-2-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-2-5	Residues	QNAADSYSWV PEQAESRAME NQYSP	25
3-3	Sequences		
3-3-1	Sequence Number [ID]	3	
3-3-2	Molecule Type	AA	
3-3-3	Length	25	
3-3-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-3-5	Residues	TSFLSINSKE ETGHLENGNK YPNLE	25
3-4	Sequences		
3-4-1	Sequence Number [ID]	4	
3-4-2	Molecule Type	AA	
3-4-3	Length	25	
3-4-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-4-5	Residues	FIPLLVVILF AVHTGLFIST QQQVT	25
3-5	Sequences		
3-5-1	Sequence Number [ID]	5	
3-5-2	Molecule Type	AA	
3-5-3	Length	25	
3-5-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-5-5	Residues	ESDRPRKVRF RIVSSHSGRV LKEVY	25
3-6	Sequences		
3-6-1	Sequence Number [ID]	6	
3-6-2	Molecule Type	AA	
3-6-3	Length	25	
3-6-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-6-5	Residues	EIYNESLFDL LSALPYVGPS VTPMT	25
3-7	Sequences		
3-7-1	Sequence Number [ID]	7	

3-7-2	Molecule Type	AA	
3-7-3	Length	25	
3-7-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-7-5	Residues	GKKLRDDYLA SLHPRLHSIY VSEGY	25
3-8	Sequences		
3-8-1	Sequence Number [ID]	8	
3-8-2	Molecule Type	AA	
3-8-3	Length	25	
3-8-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-8-5	Residues	PDIKQELLRC DIICKGGHST VTDLQ	25
3-9	Sequences		
3-9-1	Sequence Number [ID]	9	
3-9-2	Molecule Type	AA	
3-9-3	Length	25	
3-9-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-9-5	Residues	VGTKLDRDD KDNIERLRDK KLAPI	25
3-10	Sequences		
3-10-1	Sequence Number [ID]	10	
3-10-2	Molecule Type	AA	
3-10-3	Length	25	
3-10-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-10-5	Residues	VKLLGIHCQN AAITEMCLVA FGNLA	25
3-11	Sequences		
3-11-1	Sequence Number [ID]	11	
3-11-2	Molecule Type	AA	
3-11-3	Length	25	
3-11-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-11-5	Residues	NLRKSSPGTS NKCLRQVSSL VLHIE	25
3-12	Sequences		
3-12-1	Sequence Number [ID]	12	
3-12-2	Molecule Type	AA	
3-12-3	Length	25	
3-12-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic	
		source 1..25	
		mol_type=protein	
		organism=synthetic construct	
	NonEnglishQualifier Value		
3-12-5	Residues	LGRLHPCVMA SLKAQSPIPN LYLTG	25
3-13	Sequences		
3-13-1	Sequence Number [ID]	13	
3-13-2	Molecule Type	AA	
3-13-3	Length	25	

3-13-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-13-5	NonEnglishQualifier Value Residues	LLPIHTLDVK STTLPAAVRC SESRL	25
3-14	Sequences		
3-14-1	Sequence Number [ID]	14	
3-14-2	Molecule Type	AA	
3-14-3	Length	25	
3-14-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-14-5	Residues	MTMDNFGKHY TLKSEAPLYV GGMPV	25
3-15	Sequences		
3-15-1	Sequence Number [ID]	15	
3-15-2	Molecule Type	AA	
3-15-3	Length	25	
3-15-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-15-5	Residues	AVDVEGMKTQ YSVKQRTENV LRIFL	25
3-16	Sequences		
3-16-1	Sequence Number [ID]	16	
3-16-2	Molecule Type	AA	
3-16-3	Length	25	
3-16-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-16-5	Residues	HDGPFVFAEV NANYITWLWH EDESR	25
3-17	Sequences		
3-17-1	Sequence Number [ID]	17	
3-17-2	Molecule Type	AA	
3-17-3	Length	25	
3-17-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-17-5	Residues	QAKEDFSGYD FETRLHVRIH AALAS	25
3-18	Sequences		
3-18-1	Sequence Number [ID]	18	
3-18-2	Molecule Type	AA	
3-18-3	Length	25	
3-18-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-18-5	Residues	PAVRPGICPG PDGWRIPLGP LPHEF	25
3-19	Sequences		
3-19-1	Sequence Number [ID]	19	
3-19-2	Molecule Type	AA	
3-19-3	Length	25	
3-19-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic	

3-19-5	NonEnglishQualifier Value Residues	source 1..25 mol_type=protein organism=synthetic construct VAQELFQGS D LGVAEEAERP GEKAG	25
3-20	Sequences		
3-20-1	Sequence Number [ID]	20	
3-20-2	Molecule Type	AA	
3-20-3	Length	15	
3-20-4	Features Location/Qualifiers	REGION 1..15 note=Synthetic source 1..15 mol_type=protein organism=synthetic construct	
3-20-5	NonEnglishQualifier Value Residues	GTATTLTDLT NPLSL	15
3-21	Sequences		
3-21-1	Sequence Number [ID]	21	
3-21-2	Molecule Type	AA	
3-21-3	Length	45	
3-21-4	Features Location/Qualifiers	REGION 1..45 note=Synthetic source 1..45 mol_type=protein organism=synthetic construct	
3-21-5	NonEnglishQualifier Value Residues	THIRRIVPGA VSDGRMGSWR APPTLSVPAS PLTLQSHFR QQARV	45
3-22	Sequences		
3-22-1	Sequence Number [ID]	22	
3-22-2	Molecule Type	AA	
3-22-3	Length	40	
3-22-4	Features Location/Qualifiers	REGION 1..40 note=Synthetic source 1..40 mol_type=protein organism=synthetic construct	
3-22-5	NonEnglishQualifier Value Residues	RHLSQEFQWL QITPPGIPVH ESTATLQHYS SGWAEKSKIL	40
3-23	Sequences		
3-23-1	Sequence Number [ID]	23	
3-23-2	Molecule Type	AA	
3-23-3	Length	44	
3-23-4	Features Location/Qualifiers	REGION 1..44 note=Synthetic source 1..44 mol_type=protein organism=synthetic construct	
3-23-5	NonEnglishQualifier Value Residues	SPDSKIQMVS SSQKRALLCL IALLSRKQW KIRTCLRRVR QKCF	44
3-24	Sequences		
3-24-1	Sequence Number [ID]	24	
3-24-2	Molecule Type	AA	
3-24-3	Length	32	
3-24-4	Features Location/Qualifiers	REGION 1..32 note=Synthetic source 1..32 mol_type=protein organism=synthetic construct	
3-24-5	NonEnglishQualifier Value Residues	TLLSPQEAGA TKDECEGEEG AAGSRDLRSW VT	32
3-25	Sequences		
3-25-1	Sequence Number [ID]	25	
3-25-2	Molecule Type	AA	
3-25-3	Length	67	
3-25-4	Features Location/Qualifiers	REGION 1..67 note=Synthetic source 1..67 mol_type=protein	

3-25-5	NonEnglishQualifier Value Residues	organism=synthetic construct EETGMPNKAS KQGPSTQRE GSLEEIPGLT NIYKLLTSVW GLLRLWVWGP ALAFTSCVTS 60 EIAMRLL 67
3-26	Sequences	
3-26-1	Sequence Number [ID]	26
3-26-2	Molecule Type	AA
3-26-3	Length	225
3-26-4	Features Location/Qualifiers	REGION 1..225 note=Synthetic source 1..225 mol_type=protein organism=synthetic construct
3-26-5	NonEnglishQualifier Value Residues	RVLKGGSVRK LRHAKQLVLE LGEEAQNAAD SYSWVPEQAE SRAMENQYSP TSFLSINSKE 60 ETGHLENGNK YPNLEFIPLL VVILFAVHTG LFISTQQQVT ESDRPRKVERF RIVSSHSGRV 120 LKEVYEIYNE SLFDLLSALP YVGPSVTPMT GKCLRDDYLA SLHPRLHSIY VSEGYPDIKQ 180 ELLRCDIICK GGHSTVTDLQ VGTKLCLRDD KDNIERLRDK KLAPI 225
3-27	Sequences	
3-27-1	Sequence Number [ID]	27
3-27-2	Molecule Type	AA
3-27-3	Length	225
3-27-4	Features Location/Qualifiers	REGION 1..225 note=Synthetic source 1..225 mol_type=protein organism=synthetic construct
3-27-5	NonEnglishQualifier Value Residues	VKLLGIHCQN AAITEMCLVA FGNLANLRKS SPGTSNKCLR QVSSVLVHIE LGRHLPCVMA 60 SLKAQSPIPN LYLTGLLPIH TLDVKSTTLP AAVRCSESRL MTMDNFGKHY TLKSEAPLYV 120 GGMPVMTMDN FGKHYTLKSE APLYVGGMPV HDGPFVFAEV NANYITWLWH EDESRAKED 180 FSGYDFETRL HVRIHAALAS PAVRPGICPG PDGWRIPLGP LPHEF 225
3-28	Sequences	
3-28-1	Sequence Number [ID]	28
3-28-2	Molecule Type	AA
3-28-3	Length	268
3-28-4	Features Location/Qualifiers	REGION 1..268 note=Synthetic source 1..268 mol_type=protein organism=synthetic construct
3-28-5	NonEnglishQualifier Value Residues	VAQELFQGS D LGVAEEAERP GEKAGGTATT LTDLTNPLSL THIRRVPGA VSDGRMGSWR 60 APPTLSVPAS PLTLQSHFR QQARVRHLSQ EFGWLQITPP GIPVHESTAT LQHYSSGWAE 120 KSKILSPDSK IQMVSSSQKR ALLCLIALLS RKQTKWIRTC LRRVRQKCFE LLSPQEAGAT 180 KDECEGEEGA AGSRDLRSWV TEETGMPNKA SKQGPSTQR EGSLEEIPGL TNIYKLLTSV 240 WGLRLWVWG PALAFTSCVT SEIAMRLL 268
3-29	Sequences	
3-29-1	Sequence Number [ID]	29
3-29-2	Molecule Type	AA
3-29-3	Length	13
3-29-4	Features Location/Qualifiers	REGION 1..13 note=Synthetic source 1..13 mol_type=protein organism=synthetic construct
3-29-5	NonEnglishQualifier Value Residues	NSKEETGHLE NGN 13
3-30	Sequences	
3-30-1	Sequence Number [ID]	30
3-30-2	Molecule Type	AA
3-30-3	Length	23
3-30-4	Features Location/Qualifiers	REGION 1..23 note=Synthetic source 1..23 mol_type=protein organism=synthetic construct
3-30-5	NonEnglishQualifier Value Residues	FLSINSKEET GHLENGNKYP NLE 23

3-31	Sequences		
3-31-1	Sequence Number [ID]	31	
3-31-2	Molecule Type	AA	
3-31-3	Length	21	
3-31-4	Features Location/Qualifiers	REGION 1..21 note=Synthetic source 1..21 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-31-5	Residues	SINSKEETGH LENGNKYPNL E	21
3-32	Sequences		
3-32-1	Sequence Number [ID]	32	
3-32-2	Molecule Type	AA	
3-32-3	Length	19	
3-32-4	Features Location/Qualifiers	REGION 1..19 note=Synthetic source 1..19 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-32-5	Residues	NSKEETGHLE NGNKYPNLE	19
3-33	Sequences		
3-33-1	Sequence Number [ID]	33	
3-33-2	Molecule Type	AA	
3-33-3	Length	17	
3-33-4	Features Location/Qualifiers	REGION 1..17 note=Synthetic source 1..17 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-33-5	Residues	KEETGHLENG NKYPNLE	17
3-34	Sequences		
3-34-1	Sequence Number [ID]	34	
3-34-2	Molecule Type	AA	
3-34-3	Length	15	
3-34-4	Features Location/Qualifiers	REGION 1..15 note=Synthetic source 1..15 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-34-5	Residues	TSFLSINSKE ETGHL	15
3-35	Sequences		
3-35-1	Sequence Number [ID]	35	
3-35-2	Molecule Type	AA	
3-35-3	Length	17	
3-35-4	Features Location/Qualifiers	REGION 1..17 note=Synthetic source 1..17 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-35-5	Residues	TSFLSINSKE ETGHLEN	17
3-36	Sequences		
3-36-1	Sequence Number [ID]	36	
3-36-2	Molecule Type	AA	
3-36-3	Length	19	
3-36-4	Features Location/Qualifiers	REGION 1..19 note=Synthetic source 1..19 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-36-5	Residues	TSFLSINSKE ETGHLENGN	19
3-37	Sequences		
3-37-1	Sequence Number [ID]	37	

3-37-2	Molecule Type	AA	
3-37-3	Length	21	
3-37-4	Features	REGION 1..21	
	Location/Qualifiers	note=Synthetic source 1..21 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-37-5	Residues	TSFLSINSKE ETGHLENGNK Y	21
3-38	Sequences		
3-38-1	Sequence Number [ID]	38	
3-38-2	Molecule Type	AA	
3-38-3	Length	23	
3-38-4	Features	REGION 1..23	
	Location/Qualifiers	note=Synthetic source 1..23 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-38-5	Residues	TSFLSINSKE ETGHLENGNK YPN	23
3-39	Sequences		
3-39-1	Sequence Number [ID]	39	
3-39-2	Molecule Type	DNA	
3-39-3	Length	346	
3-39-4	Features	misc_feature 1..346	
	Location/Qualifiers	note=Synthetic source 1..346 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-39-5	Residues	gaacctgaag tcaccagac tcccagccat caggtcacac agatgggaca ggaagtgatc 60 ttgcgctgtg tccccatctc taatcactta tacttctatt ggtacagaca aatcttgggg 120 cagaaagtgc agtttctggt ttccctttat aataatgaaa tctcagagaa gtctgaaata 180 ttcgatgatc aattctcagt tgaaaggcct gatggatcaa atttcaactc gaagatccgg 240 tccacaaaagc tggaggactc agccatgtac ttctgtgcca gcagctggg tgacaggggt 300 aatgaaaaac tgttttttgg cagtggaaacc cagctctctg tcttgg 346	
3-40	Sequences		
3-40-1	Sequence Number [ID]	40	
3-40-2	Molecule Type	AA	
3-40-3	Length	114	
3-40-4	Features	REGION 1..114	
	Location/Qualifiers	note=Synthetic source 1..114 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-40-5	Residues	EPEVTQTPSH QVTQMGQEVI LRCVPISNHL YFYRQILGQ KVEFLVSFYN NEISEKSEIF 60 DDQFSVERPD GSNFTLKIRS TKLEDSAMYF CASSLGDRGN EKLFFGSGTQ LSVL 114	
3-41	Sequences		
3-41-1	Sequence Number [ID]	41	
3-41-2	Molecule Type	DNA	
3-41-3	Length	331	
3-41-4	Features	misc_feature 1..331	
	Location/Qualifiers	note=Synthetic source 1..331 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-41-5	Residues	gatgctaaga ccacacagcc aaattcaatg gagagtaacg aagaagagcc tgttcacttg 60 ccttgtaacc actccacaat cagtggaaact gattacatac attggtatcg acagcttccc 120 tcccagggtc cagagtacgt gattcatggt cttacaagca atgtgaacaa cagaatggcc 180 tctctggcaa tcgctgaaga cagaaagtcc agtaccttga tcttgcaccg tgctaccttg 240 agagatgctg ctgtgtaacta ctgcatcctg agacgtctta acgactacaa gctcagcttt 300 ggagccggaa ccacagtaac tgtaagagca a 331	
3-42	Sequences		
3-42-1	Sequence Number [ID]	42	
3-42-2	Molecule Type	AA	
3-42-3	Length	110	
3-42-4	Features	REGION 1..110	

	Location/Qualifiers	note=Synthetic source 1..110 mol_type=protein organism=synthetic construct	
3-42-5	NonEnglishQualifier Value Residues	DAKTTQPNMS ESNEEEPVHL PCNHSTISGT DYIHWYRQLP SQGPPEYVIHG LTSNVNMRMA 60 SLAIAEDRKS STLILHRATL RDAAVYYCIL RRLNDYKLSF GAGTTVTVRA 110	
3-43	Sequences		
3-43-1	Sequence Number [ID]	43	
3-43-2	Molecule Type	DNA	
3-43-3	Length	343	
3-43-4	Features	misc_feature 1..343	
	Location/Qualifiers	note=Synthetic source 1..343 mol_type=other DNA organism=synthetic construct	
3-43-5	NonEnglishQualifier Value Residues	gacgctggag tcacccaaag tcccacacac ctgatcaaaa cgagaggaca gcaagtgact 60 ctgagatgct ctctaagtc tgggcatgac actgtgtcct ggtaccaaca ggccctgggt 120 caggggcccc agtttatctt tcagtattat gaggaggaag agagacagag aggcaacttc 180 cctgatcgat tctcaggtca ccagttccct aactatagct ctgagctgaa tgtgaacgcc 240 ttgttgctgg gggactcggc cctctatctc tgtgccagca gcaaaggccc gggaggcaac 300 tacgagcagt acttcggggc gggcaccagg ctcacggta cag 343	
3-44	Sequences		
3-44-1	Sequence Number [ID]	44	
3-44-2	Molecule Type	AA	
3-44-3	Length	114	
3-44-4	Features	REGION 1..114	
	Location/Qualifiers	note=Synthetic source 1..114 mol_type=protein organism=synthetic construct	
3-44-5	NonEnglishQualifier Value Residues	DAGVTQSPTH LIKTRGQQVT LRCSPKSGHD TVSWYQQALG QGPQFIFQYY EEEERQRGNF 60 PDRFSGHQFP NYSSELNVNA LLLGDSALYL CASSKGPGGN YEQYFGPGR LTVT 114	
3-45	Sequences		
3-45-1	Sequence Number [ID]	45	
3-45-2	Molecule Type	AA	
3-45-3	Length	25	
3-45-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-45-5	NonEnglishQualifier Value Residues	TSFLSINSKE ETEHLENGNK YPNLE 25	
3-46	Sequences		
3-46-1	Sequence Number [ID]	46	
3-46-2	Molecule Type	AA	
3-46-3	Length	25	
3-46-4	Features	REGION 1..25	
	Location/Qualifiers	note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-46-5	NonEnglishQualifier Value Residues	RVLKGGSVRK LRHAKQLVLE LGEEA 25	
3-47	Sequences		
3-47-1	Sequence Number [ID]	47	
3-47-2	Molecule Type	DNA	
3-47-3	Length	20	
3-47-4	Features	misc_feature 1..20	
	Location/Qualifiers	note=Synthetic source 1..20 mol_type=other DNA organism=synthetic construct	
3-47-5	NonEnglishQualifier Value Residues	tgttgactca acagccacag 20	
3-48	Sequences		

3-48-1	Sequence Number [ID]	48	
3-48-2	Molecule Type	DNA	
3-48-3	Length	20	
3-48-4	Features	misc_feature 1..20	
	Location/Qualifiers	note=Synthetic source 1..20 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-48-5	Residues	ctggaccact tttctgaggg	20
3-49	Sequences		
3-49-1	Sequence Number [ID]	49	
3-49-2	Molecule Type	DNA	
3-49-3	Length	26	
3-49-4	Features	misc_feature 1..26	
	Location/Qualifiers	note=Synthetic source 1..26 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-49-5	Residues	gccacagcac tgtgctcttg aagtcc	26
3-50	Sequences		
3-50-1	Sequence Number [ID]	50	
3-50-2	Molecule Type	DNA	
3-50-3	Length	24	
3-50-4	Features	misc_feature 1..24	
	Location/Qualifiers	note=Synthetic source 1..24 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-50-5	Residues	caggcagtat ctggagtcac tgag	24
3-51	Sequences		
3-51-1	Sequence Number [ID]	51	
3-51-2	Molecule Type	DNA	
3-51-3	Length	25	
3-51-4	Features	misc_feature 1..25	
	Location/Qualifiers	note=Synthetic source 1..25 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-51-5	Residues	caccatggat acctggctcg tatgc	25
3-52	Sequences		
3-52-1	Sequence Number [ID]	52	
3-52-2	Molecule Type	DNA	
3-52-3	Length	18	
3-52-4	Features	misc_feature 1..18	
	Location/Qualifiers	note=Synthetic source 1..18 mol_type=other DNA organism=synthetic construct	
	NonEnglishQualifier Value		
3-52-5	Residues	attcaccac cagctcag	18
3-53	Sequences		
3-53-1	Sequence Number [ID]	53	
3-53-2	Molecule Type	AA	
3-53-3	Length	15	
3-53-4	Features	REGION 1..15	
	Location/Qualifiers	note=Synthetic source 1..15 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-53-5	Residues	ETGHLENGNK YPNLE	15
3-54	Sequences		
3-54-1	Sequence Number [ID]	54	
3-54-2	Molecule Type	AA	

3-54-3	Length	9	
3-54-4	Features Location/Qualifiers	REGION 1..9 note=Synthetic source 1..9 mol_type=protein organism=synthetic construct	
3-54-5	NonEnglishQualifier Value Residues	AAGIGILTV	9
3-55	Sequences		
3-55-1	Sequence Number [ID]	55	
3-55-2	Molecule Type	AA	
3-55-3	Length	10	
3-55-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-55-5	Residues	RLFPGTLIKI	10
3-56	Sequences		
3-56-1	Sequence Number [ID]	56	
3-56-2	Molecule Type	AA	
3-56-3	Length	10	
3-56-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-56-5	Residues	TRSSGSHFVF	10
3-57	Sequences		
3-57-1	Sequence Number [ID]	57	
3-57-2	Molecule Type	AA	
3-57-3	Length	25	
3-57-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-57-5	Residues	DSSLQARLFP GLTIKIQRSN GLIHS	25
3-58	Sequences		
3-58-1	Sequence Number [ID]	58	
3-58-2	Molecule Type	AA	
3-58-3	Length	10	
3-58-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-58-5	Residues	RLFPLAIKI	10
3-59	Sequences		
3-59-1	Sequence Number [ID]	59	
3-59-2	Molecule Type	AA	
3-59-3	Length	10	
3-59-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
	NonEnglishQualifier Value		
3-59-5	Residues	RLFPGTLIKI	10
3-60	Sequences		
3-60-1	Sequence Number [ID]	60	
3-60-2	Molecule Type	AA	
3-60-3	Length	8	
3-60-4	Features	REGION 1..8	

	Location/Qualifiers	note=Synthetic source 1..8 mol_type=protein organism=synthetic construct	
3-60-5	NonEnglishQualifier Value Residues	RLFPGLTI	8
3-61	Sequences		
3-61-1	Sequence Number [ID]	61	
3-61-2	Molecule Type	AA	
3-61-3	Length	11	
3-61-4	Features Location/Qualifiers	REGION 1..11 note=Synthetic source 1..11 mol_type=protein organism=synthetic construct	
3-61-5	NonEnglishQualifier Value Residues	LTIKIQRSNG L	11
3-62	Sequences		
3-62-1	Sequence Number [ID]	62	
3-62-2	Molecule Type	AA	
3-62-3	Length	11	
3-62-4	Features Location/Qualifiers	REGION 1..11 note=Synthetic source 1..11 mol_type=protein organism=synthetic construct	
3-62-5	NonEnglishQualifier Value Residues	LQARLFPGLT I	11
3-63	Sequences		
3-63-1	Sequence Number [ID]	63	
3-63-2	Molecule Type	AA	
3-63-3	Length	10	
3-63-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
3-63-5	NonEnglishQualifier Value Residues	LQARLFPGLT	10
3-64	Sequences		
3-64-1	Sequence Number [ID]	64	
3-64-2	Molecule Type	AA	
3-64-3	Length	25	
3-64-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-64-5	NonEnglishQualifier Value Residues	TIIEGTRSSG SHFVFPVPSLR DVHHE	25
3-65	Sequences		
3-65-1	Sequence Number [ID]	65	
3-65-2	Molecule Type	AA	
3-65-3	Length	25	
3-65-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-65-5	NonEnglishQualifier Value Residues	DSSLQARLFP GLAIKIQRSN GLIHS	25
3-66	Sequences		
3-66-1	Sequence Number [ID]	66	
3-66-2	Molecule Type	AA	
3-66-3	Length	25	
3-66-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25	

		mol_type=protein organism=synthetic construct	
3-66-5	NonEnglishQualifier Value Residues	TIIIEGTRSSG SHLVFVPSLR DVHHE	25
3-67	Sequences		
3-67-1	Sequence Number [ID]	67	
3-67-2	Molecule Type	AA	
3-67-3	Length	10	
3-67-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
3-67-5	NonEnglishQualifier Value Residues	TRSSGSHLVF	10
3-68	Sequences		
3-68-1	Sequence Number [ID]	68	
3-68-2	Molecule Type	AA	
3-68-3	Length	10	
3-68-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	
3-68-5	NonEnglishQualifier Value Residues	TRSSGSHFVF	10
3-69	Sequences		
3-69-1	Sequence Number [ID]	69	
3-69-2	Molecule Type	AA	
3-69-3	Length	11	
3-69-4	Features Location/Qualifiers	REGION 1..11 note=Synthetic source 1..11 mol_type=protein organism=synthetic construct	
3-69-5	NonEnglishQualifier Value Residues	TRSSGSHFVF V	11
3-70	Sequences		
3-70-1	Sequence Number [ID]	70	
3-70-2	Molecule Type	AA	
3-70-3	Length	9	
3-70-4	Features Location/Qualifiers	REGION 1..9 note=Synthetic source 1..9 mol_type=protein organism=synthetic construct	
3-70-5	NonEnglishQualifier Value Residues	TRSSGSHFV	9
3-71	Sequences		
3-71-1	Sequence Number [ID]	71	
3-71-2	Molecule Type	AA	
3-71-3	Length	8	
3-71-4	Features Location/Qualifiers	REGION 1..8 note=Synthetic source 1..8 mol_type=protein organism=synthetic construct	
3-71-5	NonEnglishQualifier Value Residues	TRSSGSHF	8
3-72	Sequences		
3-72-1	Sequence Number [ID]	72	
3-72-2	Molecule Type	AA	
3-72-3	Length	10	
3-72-4	Features Location/Qualifiers	REGION 1..10 note=Synthetic source 1..10 mol_type=protein organism=synthetic construct	

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3-72-5	NonEnglishQualifier Value Residues	FVFPVPSLRDV	10
3-73	Sequences		
3-73-1	Sequence Number [ID]	73	
3-73-2	Molecule Type	AA	
3-73-3	Length	25	
3-73-4	Features Location/Qualifiers	REGION 1..25 note=Synthetic source 1..25 mol_type=protein organism=synthetic construct	
3-73-5	NonEnglishQualifier Value Residues	TSFLSINSKE ETGHLENGK YPNLE	25