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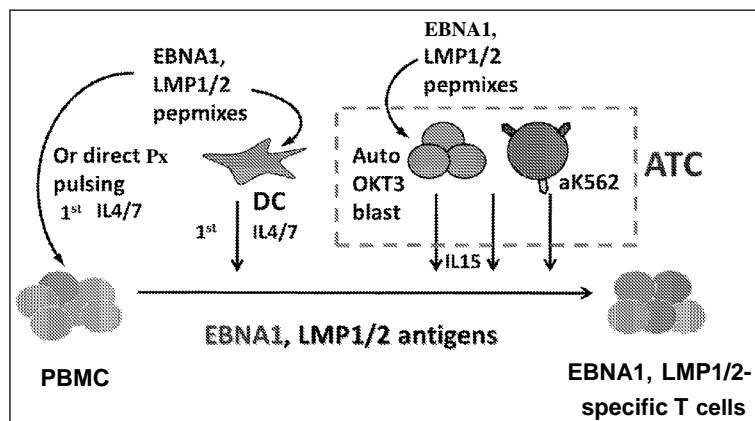
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(54) **Title:** PROCESS OF EXPANDING T CELLS

FIGURE 2: Diagram of the alternative methods demonstrated in this application of novel superior protocols for the generation of EBV specific T cells.



(57) **Abstract:** The present disclosure relates to a novel process for *in vitro* expansion of autologous or allogenic antigen specific T cells comprising the steps: a) culturing a population of autologous PCMB cells in the presence of: i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen OR a peptide/peptide mix relevant to a target antigen, and ii) at least one cytokine, and b) culturing a population of T cells from step a) in the presence of: i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen OR autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide/peptide mix relevant to a target antigen and an artificial co-stimulatory factor, and ii) optionally a cytokine, and characterized in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoding antigens or recombinant target antigens in the expansion of the relevant T cell population. The disclosure also extends to cell populations obtained from said process, pharmaceutical compositions comprising the said cell populations and use of the cells and compositions for treatment, particular the treatment or prophylaxis of virus infection and/or cancer, for example in immune compromised or immune competent patients.

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### Process of Expanding T cells

The present application claims priority from US provisional application serial number 61/569,577 incorporated herein by reference.

- 5 The present invention relates to a novel process for expanding T cells, such as autologous T cells, cell populations therefrom, pharmaceutical compositions comprising the said cell populations and use of the cells and compositions for treatment, particular the treatment or prophylaxis of virus infection and/or cancer, for example in immune compromised or immune competent human patients.

#### Background

- 10 While viruses are widely recognized as a cause of infectious disease, certain viruses are also associated with human cancer. The human immune system is central to the control of viral infections and also malignancies shown to be related to oncogenic viruses. Within the complex array of cells, antibodies and immunomodulatory molecules which constitute the human immune system, lymphocytes of thymic origin (T cells) operate in a central role to control viral infections and cancer. Hence, one  
15 approach to prevent or treat virus infections and cancer has been to take T-cells from these patients and stimulate and/or expand them *in vitro* before transfusing them back into the patient.

- In vivo* T cell activation and antigen-specific expansion is generally considered to result from a two signal process wherein the first signal is initiated by the ligation of the T cell receptor/CD3 complex with a major histocompatibility complex class I or class II molecule (MHC Class I or MHC Class II) presenting a  
20 peptide antigen. The MHC Class I or MHC Class II and peptide complex is expressed on the surface of a cell (the antigen presenting cell or APC). The peptide antigen originates from a molecule within the cell which undergoes endogenous processing and may be, *inter alia*, (1) a "self" antigen naturally occurring in the body; (2) a tumour antigen which results from a mutation related to cancer or (3) a viral antigen associated with infection or cancer. The recognition of the antigen by the T cell receptor is considered  
25 the first signal and the second signal arises from co-stimulation which results from the ligation of additional surface molecules on the T cell with additional molecules on the APC. The up-regulation and ligation of these co-stimulatory molecules between the T cell and APC may be necessary to effect or enhance T cell activation since the first signal may not be sufficient alone to achieve this. The two signal activation may lead to the expansion of the T cell so that greater numbers of the antigen-specific  
30 T cells will be available to control the pathogen or cancer giving rise to the immune response. The canonical understanding of this two signal activation is based on the same APC providing both the first signal and the second signal to the responding T cell such that the co-stimulation is directly associated with antigen recognition. The *in vitro* activation and expansion of T cells has traditionally been a long, complex and resource intensive process. A typical process may, for example take 8-12 weeks and often  
35 employs live "target" virus and/or viral vectors to achieve antigen presentation by antigen presenting cells (APC). Generally T cell activation/expansion requires static conditions rather than stirred or physically agitated culture systems.

One prior art method of culturing antigen specific T cells which recognize the LMP1 and LMP2 antigens of Epstein Barr Virus (EBV) may be summarised as follows:

#### 40 PREPARATORY STEPS

- In order to achieve the two signal T cell activation and expansion process in a controlled manner, it is useful to create an antigen-presenting cell through transfection of B cells taken from the patient.

This is referred to as establishing an autologous lymphoblastoid cell line and is undertaken through infection of the cell with EBV (EBV-LCL). It takes about 6-8 weeks to develop this cell line and thus it is one of the first stages that must be started as part of a culture process which relies upon the EBV-LCL for antigen presentation. It is prepared by culturing B cells from the patient with EBV virus in the presence of cyclosporin A to inhibit EBV-specific T cell outgrowth and elimination of the LCL.

- o Prior to the expansion step with the EBV-LCLs, the culture system involves the initial activation and expansion of the LMP-1 and LMP-2 specific T cells with autologous dendritic cells which been transduced with a viral vector Ad5f35-LMP1-LMP2 (encoding the EBV proteins LMP1 and LMP2) (days 0 to 8).
- o day 9 to 12 the cytolytic T lymphocytes (CTLs) are harvested and re-suspended in fresh medium and re-stimulated with EBV-LCLs transduced with Ad5f35-LMP1-LMP2.
- o day 13 to 16 cytolytic T lymphocytes are fed with fresh medium and recombinant human IL-2
- o followed by weekly re-stimulation using CTL:LCL and twice weekly addition of IL-2 for a 4 to 8 week period.
- Dendritic cells for use in the process must also be prepared by taking PBMCs from a patient sample and activating them with IL-4 and GM-CSF to provide adherent PBMCs. These cells are then transduced with a viral vector Ad5f35-LMP1-LMP2 (i.e. encoding the EBV protein LMP1 and LMP2). Finally the dendritic cells are matured by the addition of IL-1 $\beta$ , IL-6, PGE-1 and TNF-a.

**SUMMARY OF CELL EXPANSION STEPS**

(also referred to as preparation of Cytotoxic T lymphocytes (CTLs))

- Once prepared the transduced dendritic cells are cultured with fresh PBMCs from the patient for a period of about 10 days.
- The T cells obtained from this step are then cultured with the transduced EBV-LCLs for a period of about 1 week.
- Then the T cells obtained from the latter step are then cultured with transduced EBV-LCLs in the presence of IL-2 for a further 10 days to provide an autologous T cell antigen specific product
- This process is repeated until sufficient T cells have expanded.

J Immunother Vol 33, Number 3, April 2010 describes a faster and more efficient way of culturing the cells over a period of 23 days employing a system from Wilson Wolf known as the GRex™ system. However, this rapid process still employs the traditional viral stimuli for the cells.

There are various problems with the prior art strategy:

- (i) the use of live virus such as EBV and viral vectors has the potential to cause an immunodominant response against the vector which may interfere with efficient generation of target virus specific CTL's;
- (ii) the use of live virus is an impediment to progression to phase 3 trials due to safety concerns;
- (iii) the requirement for B cells to manufacture the EBV-LCL, now that Rituxan (which depletes B cells) has become standard therapy for most lymphoma patients means the technique cannot be employed for many patients;
- (iv) the duration of manufacturing (a minimum of 6 weeks to establish the EBV-LCL and another 5 to 7 weeks for CTL expansion) is very inconvenient, impractical and economically challenging;

- (v) the complexity of cell manipulation provides many opportunities for error and contamination of the product, hence, the principles of good manufacturing practise (GMP) are difficult to comply with, and
- (vi) the autologous antigen presenting cells used to stimulate the T cell expansion can express antigens other than the target antigen, which may reduce the purity of the antigen-specific T cells which are desired for the therapeutic T cell product.

Nevertheless skilled persons have been reluctant to move away from the established processes because each step was thought necessary to generate a product with therapeutic characteristics and in particular to generate T cell populations that are suitable for recognising cells infected by live viruses and cancers expressing viral antigens, *in vivo*.

The present disclosure provides a method for the rapid and efficient production of antigen specific T cells with specificity to a target antigen.

**Summary of the Invention**

The present disclosure provides a process for *in vitro* expansion of antigen specific T cells such as autologous antigen specific T cells comprising the steps:

- a) culturing a population of autologous PBMCs in the presence of:
  - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR a peptide/peptide mix relevant to a target antigen(s), and
  - ii) at least one cytokine, and
- b) culturing a population of cells obtained from step a) in the presence of:
  - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) and an artificial co-stimulatory factor, and
  - ii) optionally a cytokine, and

characterised in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoded antigens in the expansion of the relevant T cell population.

In one embodiment there is provided a process for *in vitro* expansion of autologous antigen specific T cells comprising the steps:

- a) culturing a population of autologous PBMCs in the presence of:
  - i) dendritic cells which have been pulsed with a peptide mix relevant to a target antigen(s), and
  - ii) at least one cytokine, and
- b) culturing a population of cells obtained from step a) in the presence of:
  - i) autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide mix relevant to a target antigen(s) and an artificial co-stimulatory factor,
  - ii) a cytokine, and

characterised in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoded antigens in the expansion of the relevant T cell population.

In the method of the present disclosure the PBMCs or dendritic cells in step a) and the antigen presenting cells of step b) are generally pulsed (also referred to as loading) with peptides selected to present epitopes from the target antigen. These peptides are discussed in more detail below.

We have overcome problems of the prior art by:

- eliminating the need to generate EBV- LCL's, and therefore avoid the use of live virus, for antigen presentation (this allows the generation of antigen specific T cells from patients that have previously been B cell depleted, e.g. by Rituxan treatment)
- eliminating the need to use viral vector-, or DNA-, or RNA-encoded antigen to achieve antigen expression and presentation in antigen presenting cells
- providing an option to eliminate the use of DCs for antigen presentation
- providing a method for T cell activation in which the stimulatory signal is provided by an autologous cell population and the co-stimulatory signal is provided by a recombinant cell line or an artificial co-stimulatory complex
- providing an efficient and robust 2-step culture process to generate a total of, for example >10e7 CD3 T lymphocytes with suitable antigen specificity in three weeks or less.
- focusing stimulation of the T cell with specificity for clinically relevant virus antigen, such as EBV antigens that are otherwise dominated by antigens that are not expressed in type 2 latency tumors (lymphoma and NPC).

15 The presently claimed invention has significant advantages for the manufacture of the autologous T cell products and potentially makes the therapy available to a wider population of patients. It also minimised the amount of time, intervention and resource required to produce a therapeutic product, and also advantageously minimises the opportunity for contamination.

20 Moreover, the specificity and properties of the therapeutic product obtained are at least equivalent to the product produced by the prior art methods and in a number of aspects may have improved properties.

Autologous cells from certain patients, such as cancer patients are different from cells obtained from healthy individuals because patient cells often are found to be anergic, i.e. incapable of delivering an immune response against antigens associated with the infection or cancer. Immune suppression is often considered to be systemic whereas anergy is usually described on an antigen-specific basis wherein a specific clone of T cells is no longer able to deliver an immune response against a target antigen. The cancer microenvironment, for example can create anergy such that T cells which recognize cancer antigens are no longer functional.

Evidence of the immune anergy or suppression is, for example the inability to clear virus infection and/or the presence of virus associated cancer cells. In healthy individuals these cells are cleared by the immune system (Teague, R. M., B. D. Sather, J. A. Sacks, M. Z. Huang, M. L. Dossett, J. Morimoto, X. Tan, S. E. Sutton, M. P. Cooke, C. Ohlen, and P. D. Greenberg. 2006. Interleukin-15 rescues tolerant CD8+ T cells for use in adoptive immunotherapy of established tumors. Nat. Med. 12:335-341 and Chemnitz, J. M., D. Eggle, J. Driesen, S. Classen, J. L. Riley, S. bey-Pascher, M. Beyer, A. Popov, T. Zander, and J. L. Schultze. 2007. RNA fingerprints provide direct evidence for the inhibitory role of TGF beta and PD-1 on CD4+ T cells in Hodgkin lymphoma. Blood 110:3226-3233).

35 Additionally, in patients with Nasopharyngeal Carcinoma (NPC), the results of autologous T cell immunotherapy with antigen-specific cytotoxic T lymphocytes (CTLs) have been relatively ineffective. In one trial only 1 of 11 patients had a complete response, and this may be explained by the inability to reactivate LMP-specific T cells from these patients. In fact the inventors hypothesise that NPC energizes T cells with specificity for the viral tumour antigens.

Thus in some patients the prior art methods were unable to reactivate appropriate antigen specific T cells adequately.

Efficacy of infused T cells depends not only on their ability to recognize the targeted tumor antigens, but also to recognize multiple epitopes within those antigens to prevent tumor escape due to epitope loss, virus strain variation and T cell driven mutation. Hence there is a need to develop a manufacturing strategy which reproducibly reactivates and expands CTLs that recognize a broad repertoire of epitopes from the antigens, such as LMP1- LMP2-, EBNA1 and BARP1- that are expressed in NPC and in EBV-positive lymphomas.

Prior to the work by the present inventors, who are leaders in the field, it was not known whether peptides could be used to generate an autologous antigen specific T cell population for prophylaxis and treatment of viral infections and cancer associated with viruses. Nor was it known that dendritic cells or T-APCs employed in the present process could be rendered useful as antigen presenting cells employing said peptides. What is more the T cell responses to the peptides seem to be relevant in the context of naturally processed peptides which are recognized by the immune system.

The present invention represents a very significant advancement in the preparation of (autologous) antigen specific T cell preparation and this is likely to result in practical benefits for patients and medical practitioners.

The factors that are important in expanded T cell populations of the present disclosure are:

- the avidity of the T cells for each epitope recognized,
- the number of epitopes recognized within each antigen,
- the number of antigens recognized,
- the fold expansion of T cells and the frequency of T cells with the desired specificity.

#### Brief Description of the Figures

**FIGURE 1** shows a diagrammatic representation of a prior art process for generating EBV specific T cells. This method required the production of autologous dendritic cells (DC) for the first round of activation/expansion and Lymphoblastoid Cell Lines (LCL) for the subsequent rounds of activation/expansion. Both, Dcs and LCLs were transfected with adenoviral vectors containing the EBV antigens of choice in order to stimulate the growth of EBV specific T cells.

**FIGURE 2** shows a diagrammatic representation of the various embodiments of the invention. The diagram shows the process of the invention by demonstrating the use of peptides to generate EBNA1, LMP1 and LMP2 specific T cells. In all new methods the use of Adenoviral vectors as the way of providing the antigen(s) was replaced by the addition of exogenous peptide(s). Also the use of LCL for the second round of activation/expansion was abolished and replaced with either peptide loaded autologous DCs or peptide loaded antigen presenting autologous T cells (T-APC) and aK562 cells. Another embodiment shows that the first stimulation can be performed without the use of DC and utilizes the antigen presenting cells present in the PBMC population.

The generation of PBMCs, DCs and T-APCs is described in this document under Protocol 1, 2 and 3 respectively.

**FIGURE 3** figures in this series show the stimulation of T cell expansion by aK562 using T-APCs in step b) of Claim 1 of the invention. This demonstrates the use of a novel system to achieve a powerful antigen specific T cell stimulation for the second round of activation/expansion. It is based on the surprising discovery that the first (antigen specific, stimulatory) signal and the second (co-stimulatory) signal can be provided by two separate components. In the example shown the first signal is provided by peptide loaded T-APCs and the second signal is provided by the aK562 cells that have been modified to present co-stimulatory molecules but not MHC. This is as explained in step b) of claim 1 of the invention.

**Figure 3A** shows the expansion of T cells using aK562 and T-APCs according in step b) of Claim 1 of the invention. This figure shows the result in 6 normal donors of using the different ratios of CTL:T-APC:aK562 during the second stimulation of the process and the resulting cell expansion. The 1:1:5 ratio was shown to be the most optimal in the majority of donors.

**Figure 3B** shows the optimal CTL to aK562 to T-APC ratio for T cell expansion in step b) of Claim 1 of the invention. The results were generated by comparing (C) the fold expansion (as individually demonstrated in figure 3A), (D) the percentage of CD56+, CD3- cells in the culture and (E) the response in an Interferon - $\gamma$  (IFN $\gamma$ ) ELISPOT of the final cell product generated using the different cell to cell ratios (SCF=Spot forming colonies).

**Figure 3C** shows a comparison of interferon gamma secreting cells in cultures of EBV specific T cells employing various ratios of CTLs to T-APCs to aK562 cells and different antigens. The number of cells in the final culture producing IFN $\gamma$  in an ELISPOT assay is shown in 2 individual donors following the second stimulation at the different culture ratios. This is shown for the 3 individual EBV antigens of interest and a control without antigen.

**Figure 3D** shows that T-APC can act as antigen presenting cells for HLA class I and II restricted antigens.

A. Upon activation of PBMC with OKT3 and CD28 antibodies (T-APC), T cells will start to upregulate HLA class II as well as co-stimulatory molecules such as CD80, CD83, CD86 and 4-1BBL. Even though HLA class II will reach up to 100% by the end of a week, the level of co-stimulatory molecules is transient and remains low.

B. aK562 is an HLA(-)ve leukemia cell line that has been engineered to express stable and high level of CD80, CD83, CD86 and 4-1BBL.

**FIGURE 4** figures in this series show the results using the prior art and the various embodiments of the invention to expand EBV specific cells from healthy donors

**FIGURE 4A** is a schematic representation of the steps of the prior art process and various embodiments of the invention and their nomenclature. This diagram shows the prior art in a shaded box and the nomenclature used to reference the first and second stimulations. The cell type used as the APC is shown first and then the way the antigen was delivered (Ad vector or Peptides -Px). For the embodiment using T-APC and aK562 plus peptide this is abbreviated in the data to ATC.

**FIGURE 4B** shows expansion of EBV-specific T cells using the prior art and the various embodiments of the invention. This figure shows the summary of results from 9 healthy donors. Following expansion of EBV specific T cells using the prior art and the 3 other methods outlined in



figure 4A, cells were counted at day 9, 16 and 23 and the results displayed below as millions of cells. Both methods employing aK562 in combination with DCs or T-APCs show significantly better expansion than the prior art.

5 **FIGURE 4C** shows a comparison of interferon gamma secreting cells in cultures of EBV specific T cells using the prior art and the various embodiments of the invention. Following culture using the prior art method and the various embodiments of the invention, the cell populations were re-stimulated with peptides in an EELISPOT assay. Peptides from the three antigens of interest (EBNA-1, LMP1 and LMP2) were used in the assay and the response was either the same or enhanced when compared to the prior art. One representative example (of a total of 9 healthy donors) is shown, followed by a graph collating the information from all normal donors for the embodiments using ATC.

10 **FIGURE 4D** shows the T cell receptor affinity for EBV specific T cells expanded by the prior art and the various embodiments of the invention. Cells were cultured to day 16 using the different methods outlined in figure 4A they were then re-stimulated with increasing dilutions of peptide in an ELISPOT assay to determine if the new embodiments of the invention would be detrimental to the avidity of the T cell receptors to peptide. In fact the new methods show a similar and in the case of EBNA1 significantly increased avidity for the individual EBV peptides.

15 **FIGURE 4E** shows the distribution of various T cell markers for cells that were expanded using the prior art and the various embodiments of the invention. At day 16 T cells produced using the 4 different methods were harvested and immune-phenotyped using flow cytometry. This shows that the composition of the cell products is unchanged between the prior art method and the embodiments of the invention. There is a difference between the prior art and the new embodiments in terms of the expression of CD62L. This is expressed on naive and central memory T cells and is down regulated by effector memory T cells therefore this again could be seen as an advantage for the embodiments of the invention.

20 **FIGURE 4F** shows that culture of cells using the prior art skews responses towards antigens expressed in LCL cells rather than on the latency type 2 antigens present in EBV+ cancer cells from lymphoma and NPC patients. Cells were generated using the prior art and embodiments of the invention. These were then re-stimulated with a variety of EBV peptides in an ELISPOT assay. This shows that the prior art method skews the response towards specific LCL dominant epitopes whereas the new methods show increased activity against cancer associated antigens such as EBNA1.

25 **FIGURE 5** figures in this series show the results using the prior art and the various embodiments of the invention to expand EBV specific cells from NPC patients

30 **FIGURE 5A** is a schematic representation of the steps of the prior art process and various embodiments of the invention and their nomenclature. This diagram shows the prior art in a shaded box and the nomenclature used to reference the first and second stimulations. The cell type used as the APC is shown first and then the way the antigen was delivered (Ad vector or Peptide -Px). For the embodiment using T-APC and aK562 plus peptide this is abbreviated in the data to ATC.

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**FIGURE 5B** shows expansion of EBV-specific T cells using the prior art and the various embodiments of the invention. Following expansion of EBV specific T cells using the prior art and the 3 other methods outlined in figure 5A, cells were counted at day 9, 16 and 23 and the results displayed below as millions of cells or fold expansion. This is representative of 3 and 4 NPC donors respectively. Both methods employing aK562 in combination with DCs or T-APCs show significantly better expansion than the prior art.

**FIGURE 5C** shows a comparison of interferon gamma secreting cells in cultures of EBV specific T cells using the prior art and the various embodiments of the invention. Following culture using the prior art method and the various embodiments the cell populations were re-stimulated with peptides in an ELISPOT assay. Peptides from the three antigens of interest were used- EBNA-1, LMP1 and LMP2 and the response in NPC patients was enhanced when compared to the prior art. One representative example is shown.

**FIGURE 5D** shows the T cell receptor affinity for EBV specific T cells expanded by the prior art and the various embodiments of the invention. Cells were cultured to day 16 using the different methods outlined in figure 5A. They were then re-stimulated with increasing dilutions of peptide in an ELISPOT assay to determine if the new embodiments of the invention would be detrimental to the avidity of the T cell receptors to peptide. In fact the new methods show a similar and in the case of EBNA1 significantly increased avidity for the individual EBV peptides.

**FIGURE 5E** shows the distribution of various T cell markers for cells that were expanded using the prior art and the various embodiments of the invention. At day 31 T cells produced using the 4 different methods described in Figure 5A were harvested and immune-phenotyped using flow cytometry. This shows that the composition of the cell products is unchanged between the prior art method and the embodiments of the invention. There is a difference between the prior art and the new embodiments in terms of the expression of CD62L. This is expressed on naive and central memory T cells and is down regulated by effector memory T cells therefore this again could be seen as an advantage for the embodiments of the invention.

There is some change as to the methods outlined in figure 5A. This is due to the NPC patient cells being subjected to more than one re-stimulation step. However the nomenclature for each stimulation remains the same. This data is representative of 4 NPC patients.

**Figure 5F** shows that T Cells from NPC patients expanded employing the new embodiments of the invention(CD3+/CD19-) kill LCL (EBV+ cancer cell-line, CD3-/CD19+) better in co-culture for 4 and 10 days as T cells expanded by the prior art. T cells and LCL were incubated at a 1:1 ratio.

**FIGURE 6** figures in this series show the results using the prior art and the various embodiments of the invention to expand EBV specific cells from lymphoma patients

**FIGURE 6A** is a schematic representation of the steps of the prior art process and various embodiments of the invention and their nomenclature. This diagram shows the prior art in a shaded box and the nomenclature used to reference the first and second stimulations. The cell type used as the APC is shown first and then the way the antigen

was delivered (Ad vector or Peptides -Px). For the embodiment using T-APC and aK562 plus peptide this is abbreviated in the data to ATC as previously but also has been alternatively abbreviated to KATpx and ATpk in this section of the data.

5 **FIGURE 6B** shows expansion of EBV-specific T cells using the prior art and the various embodiments of the invention. Following expansion of EBV specific T cells using the prior art and the 3 other methods outlined in figure 6A, cells were counted at day 9 and 16 and the results displayed below as millions of cells or fold expansion. This is representative of results with cells from 4 lymphoma patients.

10 **FIGURE 6C** shows a comparison of interferon gamma secreting cells in cultures of EBV specific T cells using the prior art and the various embodiments of the invention. Following culture using the prior art method and the various embodiments the cell populations were re-stimulated with peptides in an ELISPOT assay. Peptides from the three antigens of interest were used- EBNA-1, LMPI and LMP2 and the response in lymphoma patients was enhanced when compared to the prior art. One representative example is shown.  
15 These results show that when peptide pulsed PBMCs or DCs were used for the first expansion, the number of antigen specific T cells was significantly increased after 9 days relative to culture where the prior art was used.

**Figure 6D** shows that T Cells from lymphoma patients expanded employing the new embodiments of the invention (CD3+/CD19-) kill LCL (EBV+ cancer cell-line, CD3-/CD19+) equally well or better in co-culture for 2 or 4 days as T cells expanded by the prior art. T cells and LCL were incubated at a 1:1 ratio.

20 **FIGURE 7** figures in this series show the results using the prior art and the various embodiments of the invention to expand VZV specific cells from healthy donors. PBMCs were pulsed with overlapping peptide libraries (15 mers overlapping by 11 amino acids) (pepmixes) spanning the VZV proteins, gE, ORF10, IE61, IE62 and IE63 in the presence of IL-4 and IL-7. On days 9, 16 and 23 they were restimulated with autologous activated T cells (AATCs, T-APCs) pulsed with the same pepmixes in the presence of aK562 cells expressing co-stimulatory molecules CD80, CD83, CD86 and 4-1BB ligand (K562-cs), at a ratio of CTLs to T-APCs to K562-cs of 1:1:5. Their rate of proliferation was measured by counting and antigen-specificity was measured in gamma interferon ELISPOT assays.  
25

30 **FIGURE 7A** is a schematic representation of the steps of the prior art process and various embodiments of the invention and their nomenclature

**FIGURE 7B** shows expansion of EBV-specific T cells using the prior art and the various embodiments of the invention. Growth rate of T cells expanded using the protocol described above. >500 fold expansion can be achieved over 22 days

35 **FIGURE 7C** shows a comparison of interferon gamma secreting cells in cultures of EBV specific T cells using the prior art and the various embodiments of the invention. Frequency of T cells from 6 to 9 healthy VZV-seropositive donors that secrete gamma interferon in response to stimulation with VZV peptides after activation with pepmix pulsed PBMCs or DCs on day 0 and pepmix-pulsed autologous activated T cells (T-APCs) plus K-562-cs cells on days 9, 16 and 23. Each symbol represents one donor. Embodiments of the current invention either using peptide pulsed PBMCs or DCs during the first expansion  
40

and T-APCs in combination with aK562s for subsequent expansions result in a significant expansion of target CTLs across the various VZV antigens used.

#### Detailed Description of the Invention

Autologous T cells are cells derived from the patient i.e. cells that are natural to the patient as opposed to cells from a donor. Certain tumours associated with viral infection have developed a way to grow in the patient despite the presence of virus-specific T cells. This involves the expression of molecules that are inhibitory to T cells and the modulation of virus gene expression. Thus the T cells in these patients may have reduced function towards the cancer cells, which may be described as a form of anergy.

In autologous T cell therapy a sample of T cells are removed from the patient for activation and expansion *ex vivo*. Once the antigen-specific T cell population has been prepared it is infused into the patient where the T cell cells will further expand and will eliminate cells presenting their target antigens by direct (cytotoxic) and indirect (immune regulatory) mechanisms.

"T cell" is a term commonly employed in the art and intended to include thymocytes, immature T lymphocytes, mature T lymphocytes, resting T lymphocytes or activated T lymphocytes. A T cell can be a T helper (Th) cell, for example a T helper 1 (Th1) or a T helper 2 (Th2) cell. The T cell can be a CD4+ T cell, CD8+ T cell, CD4+CD8+ T cell, CD4-CD8- T cell or any other subset of T cells.

Antigen specific T cell as employed herein is intended to refer to T cells that recognise a particular antigen and responds thereto, for example by proliferating and/or producing cytokines in response thereto.

In one or more embodiments the process does not employ recombinant target antigens for stimulating specificity. Recombinant antigens herein refers to whole or large fragments of antigens prepared by recombinant techniques. In contrast the peptides employed are small fragments of antigen and are generally synthetic.

The present invention relates to *ex vivo* processing of cells and the T cell products obtained therefrom.

Usually the present invention does not include the step of obtaining the sample from the patient.

The step of obtaining a sample from the patient is a routine technique of taking a blood sample (which can be processed and optionally provided as an apheresis product). This process presents little risk to patients and does not need to be performed by a doctor but can be performed by appropriately trained support staff. In one embodiment the sample derived from the patient is approximately 200ml of blood or less, for example 150ml or less such as in the range 100-150ml. Generally at least about 60ml of blood is required.

Typically the PBMCs for T cell expansion, DC generation and T-APC generation are obtained from the blood or apheresis product by Ficoll density gradient separation known to those skilled in the art.

Ficoll density gradient separation employs a synthetic sucrose polymer the concentration of which varies through the solution to exploit the separation of different cells during sedimentation. Suitable reagents are available, for example from GE Healthcare, such as Ficoll Paque™PLUS.

In one embodiment the centre responsible for taking the blood sample or for shipping the blood sample processes the sample by Ficoll density gradient separation prior to transportation.

In one embodiment the blood sample or processed blood sample is transported at ambient

temperature, for example above 4°C and below about 30°C.

In one embodiment the blood sample or processed blood sample is filled into a container, such as bag, comprising two chambers, wherein one chamber contains additives, such as preservatives and/or

anticoagulants and the blood or processed blood is filled into the second chamber, after which a seal between the first and second chamber is broken and the contents of the two chambers are mixed. Culturing cells as employed herein is intended to refer to activating and expanding and/or differentiating cells *in vitro*.

- 5 It is known to the skilled person, that expansion of T cells is generally performed in a suitable T cell expansion medium. Generally the process of step a) can be performed without changing the medium. Generally the process of step b) can be performed without changing the media. However, media should be changed if indicated by a glucometer, for example that is if the glucose in the system falls below 100mg/dl. Thus the process of the present disclosure is efficient in that it minimizes the amount
- 10 of intervention required to expand the T cells.

T cell expansion may be evaluated by counting viable CD3+ cells.

- Viable cells can be tested by cell staining with, for example Trypan blue (and light microscopy) or 7-amino-actinomycin D, vital dye emitting at 670nm (or ViaProbe a commercial ready-to-use solution of 7AAD) and flow cytometry, employing a technique known to those skilled in the art. Where the stain
- 15 penetrates into the cells the cells are considered not viable. Cells which do not take up dye are considered viable. An exemplary method may employ about 5  $\mu$ L of 7AAD and about 5  $\mu$ L of Annexin-V (a phospholipid-binding protein which binds to external phospholipid phosphatidylserine exposed during apoptosis) per approximate 100 $\mu$ L of cells suspension. This mixture may be incubated at ambient temperature for about 15 minutes in the absence of light. The analysis may then be performed
- 20 employing flow cytometry. See for example MG Wing, AMP Montgomery, s. Songsivilai and JV Watson. An Improved Method for the Detection of Cell Surface Antigens in Samples of Low Viability using Flow Cytometry. J Immunol Methods 126: 21-27 1990.

T cell expansion media generally comprises serum, media and any cytokines employed in the relevant expansion step (i.e. step a) or step b)).

- 25 In one embodiment the serum employed is, for example 15% serum or less such as 10% serum, in particular human serum is employed.

In one embodiment the media is Advanced RPMI or RPMI 1640, available from Life Technologies.

In one embodiment the cytokines employed are discussed below.

In one embodiment the cell expansion medium comprises 10% human AB serum, 200 mM

- 30 L-glutamine, 45% Earle's Ham's amino acids (EHAA or Click's medium) and 45% advanced RPMI or RPMI-1640.

In one embodiment the media employed does not require the use of serum.

Cell expansion as employed herein refers to increasing the number of the target cells in a population of cells as a result of cell division.

- 35 In one embodiment in step a) the PBMCs are treated directly with a peptide/peptide mix. It was very surprising that autologous PBMCs could activate T cells in the presence of peptides in a manner similar to when autologous dendritic cells are present, in particular because the message from the literature is that dendritic cells are the optimal antigen presenting cells. Chen ML, Wang FH, Lee PK, Lin CM. Immunol Lett. 2001 Jan 1; 75(2):91-6.

- 40 In another embodiment in step a) of the present process dendritic cells are employed which are prepared from the patients PBMCs.

The blood sample is not generally subject to initial physical selection of cells, for example selection of a sub-population of cells (or T cells) from an apheresis population.

In one embodiment 1 to  $2 \times 10^7$  PBMCs are stimulated with 0.5 to  $1 \times 10^6$  peptide-pulsed DCs in the presence of cytokines in the GRex40 in 30mls of medium. A harvest of cells, for example in the range  
5 50 to  $80 \times 10^7$  antigen-specific responder T cells after 9 to 14 days of culture. However this may be lower in patients with anergic T cells.

Medium (45% advanced RPMI, 45% EHAA, 10% FCs and 200mM L-glutamine) will be changed only if indicated by a drop in glucose below 100 mg/dl (on glucometer).

Dendritic cells are often referred to, by those skilled in the art, as professional antigen presenting cells.

10 The term refers to the fact that dendritic cells are optimal in delivery the two signal activation process to T cells, i.e., in addition to presenting antigen on the cell surface, dendritic cells also provide a strong co-stimulatory signal. Both signals, stimulation by antigen presentation and co-stimulation are required to achieve T cell activation.

Dendritic cells for use in the process of the present invention may be generated from a sample of the  
15 patients PBMCs by pulsing (or loading) with a peptide mixture the details of which are discussed below. Pulsing or loading as employed herein is intended to refer simply to exposing the relevant cells, such as PBMCs or dendritic cells, to the peptide mix in an appropriate medium.

Dendritic cells for use in the process may be prepared by taking PBMCs from a patient sample and adhering them to plastic. Generally the monocyte population sticks and all other cells can be washed  
20 off. The adherent population is then differentiated with IL-4 and GM-CSF to produce monocyte derived dendritic cells. These cells may be matured by the addition of IL-1 $\beta$ , IL-6, PGE-1 and TNF-a (which upregulates the important co-stimulatory molecules on the surface of the dendritic cell) and are then transduced with a peptide mixture as described herein to provide the required dendritic cells.

Reference to generating and maturing DC in this way is found in Jonuleit H, Kuhn U, Muller G, et al. Pro-inflammatory cytokines and prostaglandins induce maturation of potent immunostimulatory dendritic  
25 cells under fetal calf serum-free conditions. Eur J Immunol. 1997;27:3135-3142.

Peptides may be added at 1 to 100ngpeptide/ $15 \times 10^6$  PBMCs or ATCs (see discussion below) or 1 to 100 ng peptide per  $2 \times 10^6$  DCs for each peptide library/pepmix.

In one embodiment PBMCs are stimulation with IL-4 and GM-CSF for 3 to 5 days followed by 1 or 2 days  
30 of maturation with GM-CSF, IL-4, TNF-a, IL-1b, PGE-1 or PGE-2 and IL-6 followed by pulsing with said peptides.

Thus in one embodiment the dendritic cells in step a) are autologous.

In one embodiment the dendritic cell response produced is balanced, CD4 and CD8 response.

Balanced CD4 and CD8 response as employed herein is intended to refer to the fact that the CD4 cells  
35 or CD8 are not depleted in the expansion process. However a balanced population as employed herein may still be skewed in that there may be more CD4 cells than CD8 cells or *vice versa*.

In one embodiment the ratio of PBMCs to dendritic cells in step a) in the range 10:1 to 50:1 respectively, for example 15:1, 16:1, 17:1, 18:1, 19:1, 20:1, 21:1, 22:1, 23:1, 24:1, 25:1, such as about  
40 20:1.

Dendritic cells provide powerful activation of T cells and when the frequency of antigen-specific T cells in the culture is low (less than 1 in 100 T cells may be specific for the antigens of interest), for example in step a) it is efficient to employ dendritic cells. However, as the numbers and frequency of antigen

specific T cells expand and the ratio of specific T cells to dendritic cells increases the efficiency of the activation of the dendritic cells decreases unless the number of dendritic cells can be increased. The antigen-specific T cells will frequently deliver a cytotoxic response to the dendritic cell following activation. The killing of the dendritic cells will then reduce the activation signal available to continue activating and expanding the target antigen-specific T cells.

Whilst dendritic cells are very effective in stimulating T cells to expand into populations specific to a target antigen it is difficult to generate large quantities of dendritic cells. Thus whilst step b) may employ dendritic cells in practice there are advantages to employing different antigen presenting cells and an artificial co-stimulator factor in step b). In some instances when the number of dendritic cells to T cells is too small no activation of the T cells is observed. This is discussed in more detail below.

Advantageously, dendritic cells are thought to be capable of activating both memory T cells and naive T cells. The presence of memory T cells in the cell populations according to the present invention may be important.

Minor population as employed herein is intended to refer to the fact that the absolute numbers of cells in the minor population is significantly lower than the number of cells in the desired population, for example 30 percent or less of the total population.

The peptide mixtures described below may be used for one or more purposes selected from pulsing of dendritic cells, pulsing of antigen presenting cells or may be employed directly with PBMCs in step a).

The peptide mixes are from a relevant viral antigen, for example an EBV viral antigen. Epstein-Barr virus, frequently referred to as EBV, is a member of the herpes virus family and chronically infects over 95% of the world population.

In one embodiment the peptides are from an antigen of papilloma virus.

In one embodiment the peptides are from an antigen of hepatitis C virus.

In one embodiment the peptides are from an antigen of vaccinia virus (VV).

In one embodiment the peptides are from an antigen of varicella zoster virus (VZV).

In one embodiment the peptides are from an antigen of human immunodeficiency virus.

In one embodiment the peptides are from an antigen of Hepatitis B, HHV-8, CMV, HTLV-1, SV40 and/or merckel cell virus.

In one embodiment the peptides are from a combination of viruses, for example any two or more described herein, such as EBV and VZV, EBV and VV, VZV and VV or EBV, VZV and VV.

Instead of culturing the autologous T cells in the presence of cells that are infected with the relevant virus, such as EBV, or in the presence of adenovirus vectors encoding viral proteins the cells are cultured in the presence of antigen presenting cells that were pulsed with a peptide or a mixture of peptides. This reduces the risk of contamination of the final product with pathogens which is important because there is no method of sterilizing the T cell product that will be infused into the patient.

In one embodiment the peptide mix or some of the peptides in the mix cover part or all of the sequence of the antigen LMP1 (Latent Membrane Protein 1 Uniprot number P03230).

In one embodiment the peptide mix or some of the peptides in the mix cover part or all of the sequence of the antigen LMP2 (Latent Membrane Protein 2 Uniprot number Q1HVJ2).

In one embodiment the peptide mix or some of the peptides in the mix cover part or all of the sequence of the antigen EBNA 1, 2, 3, 4, 5 or 6 or a combination of the same, in particular EBNA 1.

Epstein-Barr nuclear antigen 1 (EBNA1) is a multifunctional, dimeric viral protein associated with Epstein-Barr virus. It is the only viral protein of EBV that is found in all EBV-related malignancies and therefore is a significant antigen to target. It is important in establishing and maintaining the altered state that cells take when infected with EBV. EBNA1 possesses a glycine-alanine repeat sequence that separates the protein into amino- and carboxy-terminal domains. This sequence also seems to stabilize the protein, preventing proteasomal breakdown, as well as impairing antigen processing and MHC class I-restricted antigen presentation. This thereby inhibits the CD8-restricted cytotoxic T cell response against virus-infected cells. The EBNA1 transcript area originates at the Qp promoter during latency phases I and II. It is the only viral protein expressed during the first latency phase. The EBNA1 pepmix activates HLA class II-restricted cytotoxic CD4 T cells.

In one embodiment the peptide mix or some of the peptides in the mix cover part or all of the sequence of the antigen BARF 1 (BamHI A rightward reading frame I, Uniprot number Q777A5). BARF1 is a 221 amino acid protein encoded by the BARF 1 gene which is located in the BamHI-A fragment of the EBV genome. BARF1 is expressed in various EBV-associated epithelioid malignancies, for example in NK/T lymphomas and in Burkitt's lymphoma.

Other potential EBV antigens include LP and BHRF1.

The major virion/envelope proteins for vaccinia virus are described in PNAS January 7, 2003 vol 100 no. 1 page 217-222 (Drexler et al). These include A10L (major core protein p4a), H3L (heparin binding protein), C7L (host range protein 2), D8L (cell surface binding protein), B22R (unknown function) and G5R (unknown function).

Varicella zoster virus immunogens include gE, ORF10, IE61, IE62 and IE63.

Peptides from each one of the specific target antigens listed *supra* may independently be employed in step a) and/or step b) of the process.

Generally some or all of the epitopes/antigens/peptides employed or expressed for the purpose of providing a primary signal for T activation in step a) and step b) will be common to both steps.

The peptides may cover part or all of the target antigen, for example may be overlapping to increase the opportunity of presenting the amino acids of an epitope in an immunologically relevant way. Alternatively or additionally peptides of known epitopes may be included and if desired over-represented, that is to say may be a more significant percentage of the peptides presented.

Antigens in HIV include gag, pol, env, nef, gp180, gp120 and the like.

"Covers part or all of the sequence of the antigen" as employed herein is intended to refer to the fact that there is identity or significant similarity between the peptide and the relevant portion of the full length antigen, for example the peptide is substantially identical to a contiguous region of the antigen.

Selected to present epitopes as employed herein is intended to refer to the fact that the linear sequence of an epitope is known and included into a peptide mix (that is peptides are selected encoding known epitopes) or, for example where the antigen sequence has not been epitope mapped then the peptides are designed to cover part or all of the sequence in an overlapping manner to maximise the opportunity of presenting one or more appropriate epitopes. Of course a mixture of these two strategies can be employed if desired, for example known epitopes may be represented to a greater extent in a peptide mixture.

In one embodiment the peptides in the mix are from one or more relevant viral antigens, for example one, two, three, four or more.



In one embodiment the peptide mix comprises or consists of sequences from at least LMP1 and LMP2. In addition in one embodiment EBNA1 and/or BARP1 are added to the antigen mixture to reduce the chances of immune escape by mutation or down-regulation of viral antigens.

Peptide as employed herein intended to refer to short polymers of amino acids linked by peptide bonds, wherein the peptides contain at least 2 but generally not more than 50 amino acids.

The peptides employed are sufficiently long to present one or more linear epitopes, for example are on average 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20 amino acids long.

In one embodiment some of the peptides of the mixture overlap (in relation to the sequence of a single antigen), that is to say that they are from a single antigen and are arranged such that portions of the fragments and certain sequence of amino acids from the parent sequence occur in more than one peptide fragment of the mix. The overlap of the peptides means that there is redundancy in the amino acid sequence. However, this method maximises the opportunity to present epitopes from the parent antigen in an appropriate manner, particularly when epitope mapping information is not available for the parent antigen.

In one embodiment 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 amino acids overlap in each peptide.

In one embodiment the peptides in the libraries for each protein are 15 amino acids long and overlap by 11 amino acids so that all potential HLA class I epitopes can be presented from a protein. The peptides can be longer, for example 20 amino acids overlapping by 15 or 30 amino acids overlapping by 25.

Examples of suitable peptides sequences include in the sequence listing filed herewith.

In one embodiment the peptide mix comprises or consists of 2-1000 peptides, more specifically 2-500, for example 2-400, 2-300, 2-200 or 2-100 such as 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 132, 133, 134, 135, 136, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 152, 153, 154, 155, 156, 157, 158, 159, 160, 161, 162, 163, 164, 165, 166, 167, 168, 169, 170, 171, 172, 173, 174, 175, 176, 177, 178, 179, 180, 181, 182, 183, 184, 185, 186, 187, 188, 189, 190, 191, 192, 193, 194, 195, 196, 197, 198, 199, 200 or more peptides.

The peptides of step a) or the peptides employed to create antigen specific dendritic cells in step a) or b) or employed to prepare the antigen presenting cells of step b) of the process are independently, selected based on the criteria above. However, the process is most efficient where the some or all of the peptides employed or the materials such as dendritic cells and/or antigen presenting cells employed are pulsed with some or all of the same epitopes. In this situation step b) then reinforces and augments the responses generated in step a).

The peptide mixes described above may also be used to generate antigen presenting T cells (T-APCs) which are employed in some embodiments in step b) of the process. To prepare the antigen presenting T cells, they are generated from PBMCs as described below and are pulsed with a relevant peptide mix, for example peptides are added at 1 to 100 ng peptide/15x10<sup>6</sup> T-APCs for each peptide library.

Each library as employed herein may refer to the peptides made for each antigen.

In one embodiment the antigen presenting cells are autologous.

Activated T cells express HLA class I and upregulate class II antigens as well as CD80 and CD86 (transiently).

In one embodiment after pulsing with peptides to provide the specific T-APCs the latter are irradiated to ensure that they don't expand any further when they are employed in step b).

5 Irradiation may be effected employing a source of gamma radiation or a source of X-rays.

In one embodiment on day 9 to 14, about  $5 \times 10^7$  responder T cells from step a) are stimulated with  $5 \times 10^7$  irradiated T-APCs and  $5 \times 10^7$  aK562s in a GRex 500 in 400  $\mu$ l of medium containing IL-15 for up to 14 days.

US 2003/0147869 discloses that the aK562 cells described therein may be engineered to render them  
10 antigen presenting cells. These cells in theory could be expressed in significant numbers. One may think that these could be employed as an alternative to T-APCs of step b). However these aK562 antigen presenting cells do not express HLA and if they did the HLA phenotype would not match the effector T cell restriction pattern and they would activate alloantigen-specific T cells. The present inventors have found that in the absence of HLA-expressing cells there is poor activation of the relevant  
15 target T cell population. Whilst in theory these aK562 cells could be engineered to express HLA this would be need to be matched to the patient in each case thereby making the process unnecessarily complicated and expensive.

There are 7,196 HLA alleles. These can be divided into 6 HLA class I and 6 HLA class II alleles for each individual (on 2 chromosomes). They can be mixed and matched in any way and therefore introducing  
20 the appropriate combination of HLA alleles into aK562 cells to (1) reactivate all potential T cells and (b) induce no allo-reactivity would be impossible at this moment in time.

The T-APCs according to the present invention present on average at least one epitope from a target antigen and for example may present 2, 3, 4, 5, 6 or more epitopes.

In one embodiment the T-APCs present epitopes from more than one target antigen, for example 2, 3,  
25 4 or more target antigens.

In one embodiment the ratio of cells (or CTLs) obtained from step a) to T-APCs is in the range 4:1 to 1:2, for example 1:1. A high proportion of T-APCs maximises the efficiency of the expansion. Usually it is difficult to generate dendritic cells in such high proportions, which means the time taken for expansion of the relevant T cell population may be longer with dendritic cells than the time taken for expansion  
30 with antigen presenting cells. In some instances where the numbers of the dendritic cells are very low the expansion may not occur at all. Thus the process employing antigen presenting cells in step b) may be advantageous in that the periods taken for expansion are shorter and thus provide a more efficient process.

Thus in one embodiment T-APCs are employed in step b) of the present method.

35 The use of T-APCs in the present process replaces the use LCLs in the prior art process. LCL cell lines are immortalised by infection with live EBV virus. Avoiding the use of LCLs in the present process is a huge advantage.

LCLs engineered to present other viral antigens through infection with adenovirus vectors, or pulsed with peptides can provide an effective second stimulation, but for weak antigens, both adenoviral and  
40 EBV proteins from the vector and the LCL respectively produce strong competition, so that the major component of the final CTL product is often specific for adenovirus or dominant EBV antigens expressed in LCLs but not expressed in type 2 latency tumors (lymphoma and NPC).

The use of simple peptides in step b) does not seem viable because the inventors' experience is that presenting peptide mixes to the CTLs resulting from step a) simply "confuses" the cells and results in them presenting peptides on their surface. This then results in the CTLs being targets for each other and they start to destroy themselves. This causes depletion of the cells which is clearly undesirable and contrary to the purpose of performing the process.

In one embodiment step b) is performed more than once, for example 2, 3, 4, 5 or more times until sufficient numbers of the relevant antigen specific T cell population are obtained.

Sufficient numbers may, for example be sufficient cells to continue expanding *in vivo* and stimulating the patient's immune response to the target virus infection and/or target associated cancer, such as 1 to 90 x10<sup>3</sup>, 1 to 90 x10<sup>4</sup>, 1 to 90 x10<sup>5</sup>, 1 to 90 x10<sup>6</sup>, 1 to 90 x10<sup>7</sup>, 1 to 90 x10<sup>8</sup> or more cells, such as 80x10<sup>7</sup> cells.

In one embodiment there is provided a process wherein if cells do not expand sufficiently after step b) they may receive additional stimulation with:

(1) peptide-pulsed, irradiated autologous activated T cells and irradiated co-stimulatory aK562 cells,

(2) irradiated PBMCs from blood bank approved allogeneic donors and/or

(3) submitogenic doses of anti-CD3; 1 to 100 ng per mL for example 50 ng per mL.

T-APCs are not professional antigen presenting cells. Therefore, in addition to the antigen presentation provided by the T-APCs a co-stimulator factor is required to stimulate T cell expansion and differentiation.

An artificial co-stimulatory factor as employed herein is an exogenous factor which is added to the culture to provide a T cell activation signal to complement the T-APC antigen presentation signal, for example where together the co-stimulatory factor and the T-APC antigen presentation signal stimulates or facilitates the autologous T cells expanding in a specific manner i.e. that stimulates the target population of cells as they expand to be specific for the target antigen, wherein the specificity aspect is elicited by the T-APC. An exogenous factor is one that is not present in the culture of PBMCs without addition or where the naturally occurring amounts present in the cell culture are low and are augmented by addition of exogenous amounts of the factor.

Beads bearing CD80/86 may be employed as a co-factor. Beads with anti-CD28 (ligand for CD80/86) and anti-4-1BB are available but they also contain anti-CD3 which eliminates the desired antigen mediated specificity of the expansion. Thus beads with anti-CD28 (ligand for CD80/86) and anti-4-1BB in the absence of anti-CD3 form an aspect of the present disclosure.

In one embodiment the artificial co-stimulatory factor is a cell or cell-line engineered to express particular protein(s) on its surface or associated with its surface (see US 2003/0147869 for association of certain antibodies on the surface of the cell), for example a HLA negative cell line and which has been genetically modified to express co-stimulatory molecules, such as the aK562 cell-line as disclosed in US 7,745,140. Cell lines such as the latter may be employed in the process of the invention to provide a prolonged co-stimulatory signal.

Thus in one embodiment the cell line such as aK562 does not express HLA.

Thus in one embodiment the cell line such as aK562 does not express antigen.

aK562 cell as employed herein may refer to the original cell line described in US 7,745,140. However, for the purposes of the present process generally an anti-CD3 antibody will not usually be loaded onto the Fcγ receptor on the surface thereof. Preferably the aK562 cell as referred to herein is a derivative

of the original aK562 cell line comprising at least one, such as one, two, three or four co-stimulatory factors on the surface thereof, in particular independently selected from the group comprising an anti-CD28 antibody, an anti-CD80 antibody, an anti-CD86 antibody and 4-1BBL.

One or more co-factors may have a role to play in reducing apoptosis of T cells and inducing a proliferative cycle of, for example about 7 to 10 cell divisions.

The cell is engineered to express co-stimulatory factors on its surface that provide signals that are important in the stimulation (activation or differentiation) or survival of T cells and complement the signal generated by the presentation of antigens on the surface of antigen presenting cells. Examples of the molecules that may be expressed on the cell line surface selected from the group comprising CD80, CD86, CD83, 4-1BB-ligand and a combination thereof, for example 1, 2, 3, or 4 thereof. The four elements together provide a powerful co-stimulatory signal.

In one embodiment the aK562 cell further expresses OX-40 ligand on its surface.

These cells act together with the T-APCs, described above, to provide all the signals necessary for T cell activation.

CD80 and CD86 bind CD28, a surface glycoprotein present on about 80% of peripheral T cells in humans. In combination with the activation of the T cell receptor, this binding provides potent stimulation of T cells. What is more CD28 (on T cells) binding to its ligand in conjunction with T cell receptor engagement induces the production of IL-2.

In one embodiment the culture in step b) comprises only endogenous IL-2.

Alternatively the artificial co-stimulatory factory may be agonistic antibody for a relevant receptor such as antibodies that target for example CD28. These antibodies may be provided directly into the media, or attached to beads, or may be attached to the surface of the cell line (such as aK562). This is described in detail in US 2003/0147869, incorporated herein by reference.

Providing the co-factors on beads does not lead to the most efficient antigen specific expansion

possible. Thus in one embodiment the co-factors are expressed on or associated with a cell such as aK562 cell.

This artificial co-stimulatory cell employed in step b) of the present invention (such as the aK562) cell seems to be acting by a new and surprising mechanism. The process of the present disclosure is evidence for the first time that antigen presentation can be provided on one cell and the co-stimulation can be provided by a different cell to stimulate and activate T cells. In the context of generating antigen specific T cells this is very surprising because in practice in the past generally aK562 cells have been engineered to express HLA molecules or an antibody that targets the T cell receptor and model the *in vivo* systems where the T cell receptor mediated signal and the co-stimulatory signal is provided by one cell (i.e. the contact is between two cells and for example CD28 ligation on T cells in conjunction with TCR engagement induces the production of IL-2 which triggers continued proliferation; June et al 1994, Jenkins et al 1993 and Schwartz 1992). Thus certain aK562 cell lines have been established which express MHC class 1 A2 and A3Britten, CM.; Meyer, R.G.; Kreer, T.; Drexler, I.; Wolfel, T.; Herr, W. (2002), "The use of HLA-A\*0201-transfected aK562 as standard antigen-presenting cells for CD8(+) T lymphocytes in IFN-gamma ELISPOT assays", *Journal of Immunological Methods* **259** (1-2): 95-110, and Clark, R.E.; Dodi, I.A.; Hill, S.C.; Lill, J.R.; Aubert, G.; Macintyre, A.R.; Rojas, J.; Bourdon, A. et al. (2001), "Direct evidence that leukemic cells present HLA-associated immunogenic peptides derived from the BCR-ABL b3a2 fusion protein", *Blood* **98** (10): 2887-93. Thus in certain embodiments the co-

- stimulatory cell such the aK562 cell is effectively an autonomous co-stimulatory factor (Third Party Co-stimulatory factor) which together with the T-APCs stimulates the antigen specific expansion of the target T cell population. Whilst in theory a aK562 cell that expresses the Fc receptor and that can be loaded with anti-CD3 antibody (such OKT3 antibody), to provide an antigen independent activation signal, may be employed in the present processes in at least one embodiment the aK562 (or engineered cell) is not loaded with an anti-CD3 antibody because generally the non-specific signal is not desirable. The present disclosure also extends to the use of an engineered cells line, such as aK562 cell, for providing an artificial co-stimulatory signal independent of antigen presentation in the expansion of T cells, such as autologous T cells.
- Thus in one embodiment the process is characterised further in that the process of expanding the population of cells in step b) is surprising in the context of relying upon only (1) T-APCs (2) peptides to pulse/load the latter and (3) aK562 cells transduced to express only co-stimulatory molecules without antigen-specific presentation qualities. This process as described achieves expansion on an antigen-specific basis (as measured by an increasing percentage of antigen-specific T cells during the expansion process) despite the fact that the aK562 cells have not been engineered to present specific antigens. Without the addition of other APCs in this step (e.g. DCs), the expansion process is therefore relying upon T-APCs to achieve the first signal activation with respect to presentation of the target antigen. This is unusual since T-APCs are considered sub-optimal with respect to both antigen presentation and co-stimulation. The role of aK562 cells to provide co-stimulation and to enhance expansion has been described before, but the previous demonstration showed general CD3+ T cell expansion at the cost of reduced antigen specificity since the aK562 cell were not engineered to present specific antigens. In the present invention, the antigen-specificity is increased with expansion which indicates that the first signal antigen recognition step is being achieved by the T-APC which is acting synergistically with the second signal co-stimulation from the aK562 cell. This bifurcation of the antigen presentation and co-stimulation signals contravenes the currently embraced paradigm for antigen-specific T cell expansion wherein the first signal and second signal are delivered from the same APC.
- In one embodiment the invention provides a method of stimulating and activating antigen specific T cell expansion employing antigen presentation on a first cell (or population of cells) and an artificial co-stimulatory factor which is a second distinct cell.
- In one embodiment the engineered cell line, such as the aK562 cell line is irradiated before use in the method of the present disclosure, for example with a gamma radiation source or an X-ray source. The engineered cell line, such as the aK562 cell line may be provided in a frozen form in which case irradiation of the cells may be performed after freezing.
- In one embodiment the ratio of CTLs to co-stimulatory factor (for example where the co-factor is a cell line) is in the ratio of 2:1 to 1:10 respectively, such as 1:5.
- Suitable ratios of CTLs: antibody: co-stimulatory cells are in the range 1:1:0.2 to 1:1:10, such as 1:1:5. Advantageously, the method employing T-APCs and artificial co-stimulatory factors, such as engineered cells, may provide for improved levels of antigen specific T cell expansion in comparison to prior art methods.
- The cytokine or cytokines employed in step a) and optionally step b) must be appropriate for stimulating and/or activating T cell growth or differentiation or perform some other useful function such as promotes T cell survival or the like.

Cytokines that may be employed in the process of the current disclosure include IL-1, IL-2, IL-4, IL-6 IL-7, IL-12 and IL-15.

In one embodiment the cytokines employed in the process according to the present disclosure are independently selected from IL-4, IL-7 and IL-15, such as a combination thereof.

- 5 In one embodiment in step a) the cytokines employed are IL-4 and/or IL-7. Whilst not wishing to be bound by theory the inventors believe that these cytokines have a role to play in regards to frequency, repertoire and expansion of viral antigen specific cells.

In one embodiment if IL-2 is employed in step a) then it is added at about day 3 or 4 of the culture and not at the outset.

- 10 The repertoire of T cells may be determined by ELISPOT analysis after stimulation with peptide libraries aliquotted into pools such that each peptide is uniquely represented in two pools (Kern, F., N. Faulhaber, C. Frommel, E. Khatamzas, S. Prosch, C. Schonemann, I. Kretzschmar, R. Volkmer-Engert, H. D. Volk, and P. Reinke. 2000. Analysis of CD8 T cell reactivity to cytomegalovirus using protein- spanning pools of overlapping pentadecapeptides. Eur J Immunol. 30:1676-1682 and Straathof, K. C, A. M. Leen, E. L. Buza, G. Taylor, M. H. Huls, H. E. Heslop, C. M. Rooney, and C. M. Bollard. 2005. Characterization of latent membrane protein 2 specificity in CTL lines from patients with EBV-positive nasopharyngeal carcinoma and lymphoma. J. Immunol. 175:4137-4147).

In one embodiment in step b) the cytokine employed is IL-15. Whilst not wishing to be bound by theory it is believed by the inventors that the IL-15 has a pro-survival effect on the relevant T cell population.

- 20 In one embodiment the cytokine employed in step b) is IL-15.

Cytokines such as IL-15 may be replaced during culture e.g. twice weekly.

In one embodiment only the particular cytokines described in any one of the embodiments herein are employed.

IL-12 has a role in Th1 focussing and exogenous IL-12 may be omitted if a balanced Th1/Th2 is desired.

- 25 In one embodiment the process of the present disclosure does not employ exogenous IL-12. However, in the context of the present T cell product a Th1 response in the CD4+ population is thought to be desirable.

The exogenous cytokines may be added at any stage of the process as appropriate, including concomitant addition when the cells are transferred into the culture system or at the start of the given step. The latter applies to step a) and/or step b).

- 30 The presence of exogenous cytokines in step a) and/or step b) may alternatively be added part way through the step, for example 1, 2, 3, 4 days or more after the step is initiated.

In one embodiment the process of the present disclosure is employed to provide cell population comprising a CD4+ T cell population, for example a Th1 population. A Th1 population as employed herein is intended to refer to a CD4+ population wherein 5% of the cells or more, such as 10, 20, 30, 40, 50, 60, 70, 80, 90% or more are classified as Th1.

- 35 Th1 cells, amongst other functions, generally maximise the killing efficacy of macrophages and the proliferation of cytotoxic CD8+ cells.

Memory T cells are a sub-category of Th1 cells.

- 40 In one embodiment the population of cells obtained from the process comprise a sub-population of memory T cells. Whilst not wishing to be bound by theory we believe that a substantial portion of the T cells obtained from the process will be derived from the memory portion of the starting population.

In one embodiment IL-2 is not employed in step (a) because the expansion promoted by this cytokine may be too non-specific and produce expansion of NK cells, T cells of unwanted function/specificity and T regulatory cells that may produce a certain amount of anergy in the cells. Thus in one embodiment the only IL-2 present in the culture is endogenous IL-2, i.e. it is secreted by the cells.

- 5 In one embodiment IL-2 is not employed in step (b) because it may promote a more differentiated phenotype than IL-15

Anergy as employed herein is intended to refer to a lack of responsiveness of the cells to antigen stimulation. Anergy can be measured using a functional assay, for example interferon gamma secretion by the relevant cell population. A lower level of interferon gamma secretion in comparison to full  
10 functional (non-anergic) cells may be indicative of a degree of anergy. Of course the greater the degree of anergy in the cells the lower the particular (marker) functionality will be.

As described above anergy in the context of the expanded product is intended to be a generic term that refers to reduced cell function in one or more relevant ways. The term includes cell exhaustion, for example where the cells are no longer able to divide. The cell is then referred to as senescent. Cells  
15 stop dividing because the telomeres, protective bits of DNA on the end of a chromosome required for replication, shorten with each copy, eventually being consumed.

It is thought that that PD-1 (programmed cell death protein 1; Uniprot Q15116) which is expressed on the surface of cells, is a marker of anergy. In one embodiment less than 10%, for example 9, 8, 7, 6, 5,  
4, 3, 2, 1, 0.5% or less of the relevant expanded T cell population express PD-1, such as about 1%.

- 20 In patients with high levels of anergy in the T cells such as patient's with relapsed NPC the frequency of LM P/EBNA1-specific T cells may be increased in the presence of blocking antibodies to PD-L1 or PD-1, i.e. which block signalling from the ligand binding to the receptor.

This phenomenon was not found in healthy donors, suggesting that type 2 latency antigen specific T cells might be anergic in NPC patients.

- 25 Target antigen as employed herein is intended to refer to the antigen which is employed to generate specificity in the T cells to the therapeutic target, such as a particular virus, for example EBV. Thus the cells infected by target virus or cancer cells will usually express the target antigen and hence will themselves become a target for clearance by the immune system.

- 30 In one embodiment the population of T cells expanded are a balance CD4+ and CD8+ population, that is to say the cell population comprises both CD4+ and CD8+ cells but not necessarily in equal amounts.

Balanced in the context of the present specification is employed to infer that one of the populations CD4+ or CD8+ is not depleted during expansion.

- 35 The cell populations expanded using the process of the present disclosure comprises the desired T cell population and generally will not consist only of the desired population. The final product administered to the patient may include a number of other cells that the process did not target the expansion of. In one embodiment the desired population of CD4+ and CD8+ cells comprises about 60% or less of the total population of cells. Frequency of the cell populations may be measured employing a gamma-IFN ELISPOT assay or employing Multimer (e.g. Tetramer) staining both of which are known to persons skilled in the art.

- 40 In one embodiment the T cells population obtained from the process are diverse when analysed by spectratyping, but without the emergence of dominant clone. That is to say the T cell diversity in the

starting sample is substantially represented in the expanded T cells, i.e. the expansion is not generally providing a monoclonal or oligo clonal target cell population.

A significant proportion for example 30% to 60% of the expanded cells will generally express effector memory markers including CD27, CD28, CD62L and/or CD45RO.

5 It is expected that 50% or more, such as, 60, 70, 80, 90 or 95% of the antigen specific T cells according to the present disclosure may be capable of killing target cells expressing antigens that were used in the expansion process.

We believe sufficient cells even for the highest cell doses required for the treatment of patients can be prepared employing two stimulations employing methods of the present disclosure taking in the range  
10 of 16 to 24 days of T cell culture compared to 28 to 63 days of LMP-T cell culture using Ad vectors and LCLs.

What is more the cells produced by the process of the present invention may be advantageous in that there are more specific, produce higher levels of gamma interferon and/or express lower levels of anergy markers compared to cells prepared by the prior art processes.

15 In one embodiment, anergic T cells from cancer patients may grow poorly and require three stimulations. The third stimulation might include additional stimulatory components. Since at this time the majority of cells should be specific, there is not a concern about expansion of unwanted cells.

In one embodiment the third stimulation culture will comprise (1) peptide-pulsed, irradiated autologous activated T cells, (2) irradiated costimulatory cell line such as aK562 cells, (3) irradiated  
20 PBMCs from blood bank approved allogeneic donors and (4) submitogenic doses of anti-CD3; 1 to 100 ng per mL for example 50 ng per ml.

In one embodiment one or more tests are performed to establish the suitability of the cells for use in therapy, for example an interferon gamma ICS (intracellular cytokine staining) and/or an interferon gamma ELISPOT assay.

25 In one embodiment the suitability testing is combined in a single flow cytometry assay.

In one embodiment a <sup>51</sup>Cr release assay for non-specific killing is performed, but this will generally not form part of the suitability assays.

In one embodiment batch release testing is set out in **Table 1**:

TEST	Specification
1. <b>Viability</b> Dye exclusion or flow cytometry	> 70% Viable
2. <b>Purity</b> Flow cytometry	> 80% CD3+
<b>3. Safety</b>	
3.1 Endotoxin (LAL assay)	≤ 5.0 EU/ml
3.2 Bacterial sterility	
Aerobic Bacteria- Bactec	Negative
Anaerobic Bacteria - Bactec	Negative
Fungal contaminants - Bactec	Negative
3.3 Mycoplasma PCR Assay	Negative
3.4 Non-specific killing*	<10% killing target cells at 20:1 E:T ratio
4. Identity	HLA class 1 antigen identical with patient/donor



<b>5. Potency</b>	
5.1 IFN- V+ cells	> 3 standard deviations greater response to stimulation with specific peptides than to control stimulation
<b>6. Dosing</b>	
CD3+ number	1-10 x 10 <sup>7</sup> cells per m <sup>2</sup> or 2-20 x 10 <sup>5</sup> per kg BW

The process of the present disclosure may be performed in a well or containers but most suitably is performed in a gas-permeable system. Vessel as employed herein is intended to refer to any type of container suitable for retaining the cells, media etc, for example a bag, such as an infusion type bag (provided with a gas permeable portion) or a rigid vessel, such as the GRex™ system. The gas permeable layer facilitates rapid expansion of cells and minimizes the number of media changes required.

In one embodiment step a) is performed in a GRex™ 10 system. The GRex™ 10 system is suitable for culturing up to 1x10<sup>8</sup> T cells.

In one embodiment step b) is performed in a GRex™ 100 system.

WO 2005/035728 incorporated herein by reference describes how to prepare a gas permeable vessel. In one embodiment silicone gas permeable material is employed.

In one embodiment the system employed is a GRex™ system from Wilson Woolf. In one embodiment the system is adapted to provide aseptic preparation as described in US provisional application serial number 61/550,246 incorporated herein by reference.

Generally the system is seeded with about 0.5 million cells per cm<sup>2</sup> of surface area. In a GRex-10 with a surface area of 10 cm<sup>2</sup>, a minimum of 5 million and up to 20 million cells would be seeded.

In one embodiment in step (a) 20 million cells might be seeded in a GRex-10. Within PBMC, less than 1% of cells are specific for the peptides, so at most 0.2 million specific cells are seeded. The remaining PBMCs act as feeder cells.

In one embodiment in step (b) 50 million irradiated aK562 cells plus 10 million peptide-pulsed activated and irradiated T cells and 10 million effector T cells would be seeded into a GRex-100 (100 cm<sup>2</sup>). In this case only the effector T cells will proliferate.

In one embodiment the present disclosure extends to the cell composition obtained directly from the process.

In one embodiment the process according to the present disclosure generates sufficient CD3+ cells to provide at least two individual doses for a patient.

The invention also extends to compositions with the same desirable characteristics of the cell populations prepared by the methods disclosed herein.

Thus in one aspect the present disclosure provides an autologous T cell population expanded *in vitro* to contain a population of T cells specific to a target antigen, such as a virus, wherein the population is substantially free of target virus contamination and responses to viral vectors.

The present process also relates to the preparation of dendritic cells pulsed with peptide mixes and the cell populations obtained therefrom, for use in the expansion of antigen specific T cell populations, particularly as described herein.

The present process also relates to the process of preparing T-APCs pulsed with peptide mixes and the cell populations obtained therefrom.

The present disclosure also relates to autologous T cell populations described herein, for example comprising a population of CD4+ and CD8+ antigen specific T cells, wherein the antigen is associated with virus infected cells or cancer cells.

In one or more embodiments the cell populations according to the present disclosure have one or more advantageous properties in comparison to cells prepared by the prior art method.

In one embodiment the average cell diameter of cells in the relevant T cell population is 10 to 14 $\mu$ M.

Advantageously the cells cultures of the present invention produce generally low toxicity after infusion, for example are associated with few toxicity intolerance responses, for example inflammatory responses, cell damage, flu like symptom, nausea, hair loss or the like.

The cell populations according to the present disclosure also provide advantageous properties, for example high levels of interferon gamma expression.

High levels of interferon gamma expression as employed herein is intended to refer to the fact that on average cell populations prepared by the current method may express higher levels of interferon gamma than cells prepared by prior art methods and certainly express higher levels of interferon gamma than the original anergic cells obtained from the patient.

In one embodiment the cells of the present disclosure show enhanced antigen specificity, for example in an assay disclosed herein, for example the cells of the present disclosure contain a higher frequency of cells that secrete cytokines in response to stimulation with the antigens in comparison to cells prepared by a prior art method. This figure will usually be a mean which is a per cell value derived from the values obtained for the population and divided by the number of cells present.

In one embodiment the cell populations of the present disclosure show comparable avidity (not significantly different) to cell populations prepared by a prior art method.

To determine avidity, autologous activated T cells may be pulsed with dilutions of peptide, labelled with <sup>51</sup>chromium and used as targets in a standard cytotoxicity assay. The most avid T cells are those that kill target cells pulsed with the lowest concentration of peptide. Alternatively IFN gamma production can be measured using an ELISPOT assay with dilutions of peptide.

In one embodiment the cell populations of the present disclosure show increased ability to kill target cells in comparison to cell populations prepared by a prior art method.

In one embodiment the T cell populations provided by the present disclosure are effective in expanding *in vivo* to provide an appropriate immune response to cells infected by a target virus and/or cancer cells associated with a target virus.

The present invention also extends to compositions comprising the autologous T cell populations according to the invention. These compositions may comprise a diluent, carrier, stabilizer, surfactant, pH adjustment or any other pharmaceutically acceptable excipient added to the cell population after the main process steps. An excipient will generally have a function of stabilizing the formulation, prolonging half-life, rendering the composition more compatible with the *in vivo* system of the patient or the like.

In one embodiment a protein stabilizing agent is added to the cell culture after manufacturing, for example albumin, in particular human serum albumin, which may act as a stabilizing agent. The amounts albumin employed in the formulation may be 0 to 50% w/w, such as about 12.5% w/w.

In one embodiment the formulation also contains a cryopreservative, such as DMSO. The quantity of DMSO is generally 20% or less such as about 12% in particular 10%w/w.

5 In embodiment the process of the present invention comprises the further step of preparing a pharmaceutical formulation by adding a pharmaceutically acceptable excipient, in particular an excipient as described herein, for example diluent, stabilizer and/or preservative.

Excipient as employed herein is a generic term to cover all ingredients added to the T cell population that do not have a biological or physiological function.

Once the final formulation has been prepared it will be filled into a suitable container, for example an infusion bag or cryovial.

10 In one embodiment the process according to the present disclosure comprises the further step of filling the T cell population or pharmaceutical formulation thereof into a suitable container, such as an infusion bag and sealing the same.

In one embodiment the container filled with the T cell population of the present disclosure or a pharmaceutical composition comprising the same is frozen for storage and transport, for example is  
15 store at about -135°C, for example in the vapor phase of liquid nitrogen.

In one embodiment the process of the present disclosure comprises the further step of freezing the T cell population of the present disclosure or a pharmaceutical composition comprising the same. In one embodiment the "product" is frozen by a controlled rate freezing process, for example reducing the temperature by 1 °C per minute to ensure the crystals formed are small and do not disrupt the cell  
20 structure. This process may be continued until the sample has reached about -100°C.

A product according to the present disclosure is intended to refer to a cultured cell population of the present disclosure or a pharmaceutical composition comprising the same.

In one embodiment the product is transferred, shipped, transported in a frozen form to the patient's location.

25 In one embodiment the product according to the present disclosure is provided in a form suitable for parenteral administration, for example infusion, slow injection or bolus injection. In one embodiment the formulation is provided in a form suitable for intravenous infusion.

In one aspect the present disclosure provides a method of transporting a product according to the present disclosure, from the place of manufacture, or a convenient collection point to the vicinity of the  
30 intended patient, for example where the T cell product is stored below 0°C, such as -135°C during transit.

In one embodiment the temperature fluctuations of the T cell product are monitored during storage and/or transport.

35 In one embodiment there is provided a product of the present disclosure for use in treatment, for example in the treatment of a viral associated disease or malignancy, such as EBV infection, CMV infection, adenovirus infection, HIV infection, hepatitis C or B infection, parvovirus infection, influenza virus infection, or a cancer from viral origin, for example EBV-associated lymphoma or carcinoma, HHV8-associated sarcoma, papillomavirus-associated carcinoma or SV40-associated cancers.

40 Other viral morbidities include varicella zoster virus infection, vaccinia virus infection and complications of either of the same.

In one embodiment the treatment is of an immunosuppressed patient.

In one embodiment, the patient is not immune-compromised.

In one embodiment there is provided a method of treating a patient with a product according to the present disclosure comprising the step of administering a therapeutically effective amount of product defined herein.

5 Therapeutically effective amount, does not necessarily mean an amount that is immediately therapeutically effective but includes a dose which is capable for expansion *in vivo* (after administration) to provide a therapeutic effect.

Thus there is provided a method of administering to a patient a therapeutically effective amount which is a sub-therapeutic dose of expanded T cells which are capable for expansion *in vivo* to provide the desired therapeutic effect, for example.

10 In one embodiment the antigen specific T cell population produced is specific to EBV virus and prevents, ameliorates or eliminates cells infected with EBV and/or clinical pathologies associated therewith, for example EBV associated cancers.

Symptoms of infection include fever, sore throat, and swollen lymph glands. Sometimes, a swollen spleen or liver involvement may develop. Heart problems or involvement of the central nervous system occurs only rarely. EBV remains dormant or latent in a few cells in blood for the rest of the person's life and can be reactivated, for example in immunosuppressed patients when immune controls are reduced or absent. Whilst the infection is not fatal in individuals with a healthy immune system, the infection can lead to severe complications and death in immunosuppressed patients.

EBV is best known as the cause of infectious mononucleosis. It is also associated with particular forms of cancer, particularly Hodgkin's lymphoma, non-Hodgkin's lymphoma, Burkitt's lymphoma, nasopharyngeal carcinoma, gastric carcinoma and central nervous system lymphomas associated with HIV. Finally, there is evidence that infection with the virus is associated with a higher risk of certain autoimmune diseases, especially dermatomyositis, systemic lupus erythematosus, rheumatoid arthritis, Sjogren's syndrome, and multiple sclerosis.

25 In one embodiment the target patient population for treatment has nasopharyngeal carcinoma.

In one embodiment the target patient population for treatment has gastric carcinoma.

Thus T cell populations specific to EBV according to the present disclosure can be employed in the treatment of prophylaxis of one or more of the above conditions.

30 In one embodiment there is provided a method of treatment or prophylaxis of a varicella zoster virus infection and/or complications associated therewith, including encephalitis, pneumonia, postherpetic neuralgia, shingles, zoster multiplex, myelitis, herpes ophthalmicus and zoster sine herpette, comprising administering a therapeutically effective amount of a T cell population comprising T cells specific to VZV, according to the present disclosure.

35 In one embodiment there is provided a method of treatment or prophylaxis of a vaccinia virus infection and/or complications associated therewith.

In one embodiment a dose of at least  $1 \times 10^7$  cells per  $m^2$  or  $2 \times 10^5$  per kg is employed.

In one embodiment a first dose of about  $2 \times 10^7$  per  $m^2$  and a second dose of  $2 \times 10^7$  or  $1 \times 10^8$  T cells per  $m^2$  is employed.

40 In one embodiment the present disclosure provides use of a peptide mix covering a viral or cancer antigen(s) for generating autologous dendritic cells suitable for use in expansion of autologous antigen specific T cells.

In one embodiment the present disclosure provides use of a peptide mix covering a viral or cancer antigen(s) for generating autologous T-APCs suitable for use in the expansion of antigen specific T cells.

In one embodiment the present disclosure provides use of aK562 cells or other artificial co-stimulatory factory for use as a co-stimulatory factor without concomitantly presenting antigen thereon in the expansion of antigen specific T cells (in particular autologous T cell expansion), for example employed in conjunction with T-APCs for expansion of antigen specific T cells, in particular antigen specific T cells, such as autologous antigen specific T cells.

In one embodiment there is provided a kit comprising a peptide mix covering a viral or cancer antigen(s) and an artificial co-stimulatory factor such as an aK562 cell, in particular for use in T cell expansion, such as autologous antigen specific T cell expansion. In one embodiment the kit further comprises IL-15.

In one embodiment the kit comprising a peptide mix covering a viral or cancer antigen(s) and IL-4 and IL-7 for use the expansion of antigen specific T cells, for example autologous antigen specific T cells.

**PROTOCOLS AND EXAMPLES**

**Abbreviations**

<b>APCs</b>	Antigen presenting cells
<b>ATC</b>	Activated T cell
<b>CTLs</b>	Cytolytic T lymphocytes
<b>DCs</b>	dendritic cells
<b>EBV</b>	Epstein-Barr Virus (a virus from the Herpes family of viruses)
<b>LCL</b>	lymphoblastoid cell line
<b>EBV-LCL</b>	lymphoblastoid cell line infected with EBV
<b>LMP1</b>	Latent Membrane Protein 1 Uniprot number P03230
<b>LMP2</b>	Latent Membrane Protein 2 Uniprot number Q1HVJ2
<b>BARF1</b>	protein encoded by the BamHI rightward reading frame uniprot number P03228
<b>EBNA1</b>	EBV nuclear antigen 1 Uniprot number P03211
<b>PBMC</b>	peripheral blood mononuclear cell
<b>CMV</b>	Cytomegalovirus

**PROTOCOL 1: PROCESSING OF SAMPLE TO OBTAINPBMCs**

The steps below should be performed in a certified biological safety cabinet, using aseptic techniques and following universal precautions.

A blood sample or an apheresis sample or a buffy coat sample (which is the fraction of an anti-coagulated blood sample after density gradient centrifugation that contains most of the white blood cells and platelets) is diluted with an equal volume of Dulbecco's Phosphate Buffered Saline or RPMI medium (for example RPMI 1640 available from Life Technologies), at ambient (room) temperature.

In a 50ml centrifuge tube, 15ml Lympho-prep is carefully overlaid with approximately 30ml of diluted blood. This step can be adjusted to utilize all the available cells.

The material is centrifuged at 400 x G for about 40 minutes at ambient temperature.

Aliquots, for example 3 x 1 ml plasma aliquots may be stored at -80°C.

The PBMCs interface is harvested into an equal volume of Dulbecco's Phosphate Buffered Saline or RPMI medium (for example RPMI 1640 available from Life Technologies). Centrifuge at about 450 x G

for about 10 minutes at room temperature and then aspirate supernatant. The pellet obtained should be loosened and re-suspended in about 20ml Dulbecco's Phosphate Buffered Saline or RPMI medium (for example RPMI 1640 available from Life Technologies).

If the process is performed multiple times then the cells may be combined in a single centrifuge tube and thoroughly re-suspended. The cells will generally be counted by removing 20 $\mu$ l of cells and adding 20 $\mu$ l of 50% red cell lysis buffer and counted using a hemacytometer in accordance with the manufactures instructions.

The PBMCs may be used in the expansion antigen specific T cells, expansion of CD3 and CD28 activated T cells and in the preparation of dendritic cells.

10 NOTES: Since the PBMCs cells prepared are ultimately intended for infusion into patients it is essential to adhere to procedures for the identification and handling of patient samples. Only one patient's sample should be handled at a time. Generally the number of PBMCs recovered will be in the range 0.5 to 2.0 x 10<sup>6</sup> PBMCs per ml of blood. Up to 1 x 10<sup>9</sup> may be recovered from a buffy coat.

**References:**

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**PROTOCOL 2: GENERATION OF DENDRITIC CELLS**

Dendritic cells can be differentiated from adherent of CD14-selected PB mononuclear cells (PBMC) by culture in GM-CSF and IL-4. The dendritic cells can then be matured using GM-CSF, IL-4, IL-1 $\beta$ , IL-6, TNF- $\alpha$  and PGE-1 or PGE-2 (PGE= Prostaglandin E). The dendritic cells loaded with peptides can present the peptides on HLA class I and class II molecules to antigen-specific T cells via the TCR.

Preparation of Adherent PBMCs- Dilute heparinized peripheral blood, for example 30ml, in an equal volume of Dulbecco's Phosphate Buffered Saline or RPMI medium (for example RPMI 1640 available from Life Technologies), at ambient (room) temperature.

Alternatively thaw previously frozen PBMC, wash twice in CellGenix DC medium, and count the cells.

In a 50ml centrifuge tube, 15ml Lymphoprep is carefully overlaid with approximately 30ml of diluted blood. This step can be adjusted to utilize all the available cells.

The material is centrifuged at 400 x G for about 40 minutes at ambient temperature.

35 The PBMCs interface is harvested into an equal volume of Dulbecco's Phosphate Buffered Saline or RPMI medium (for example RPMI 1640 available from Life Technologies). Centrifuge at about 450 x G for about 10 minutes at room temperature and then aspirate supernatant. The pellet obtained should be loosened and re-suspended in about 20ml Dulbecco's Phosphate Buffered Solution.

The above steps are the same as Example and then the material is then centrifuged at 400 x G for about 40 5 minutes at room temperature. The supernatant is then removed and the pellet is re-suspended in 20ml of DC medium.

Count cells as defined in Example 1. If the concentration is greater than  $5 \times 10^6$  per ml adjust to  $5 \times 10^6$  per ml by adding CellGenix DC medium. If the concentration is less than  $5 \times 10^6$  per ml the pellet and re-suspend at  $5 \times 10^6$  per ml in CellGenix DC medium.

Transfer 10ml of cells per 75cm<sup>2</sup> flask or 2ml of cells/well of a 6 well plate.

- 5 Transfer to 37°C and 5% carbon dioxide for about 2 hours. Rinsing the flask three times 10mls of Dulbecco's Phosphate Buffered Saline or RPMI medium combining the supernatants containing the PBMC non-adherent fraction.

For a T-75 culture flask (available from Falcon) add 10ml of DC culture medium containing 1000 units per ml of IL-4 and 800 units per ml of GM-CSF to the adherent cells.

- 10 For a 6-well plate add 2mls of DC culture medium containing 1000 units per ml of IL-4 and 800 units per ml of GM-CSF to the adherent cells.

Transfer the flasks or plated to an incubator at 37°C and 5% carbon dioxide.

If not previously cryopreserved, non-adherent cells may be cryopreserved for future use in the preparation of responder T cells.

- 15 On day 3 or 4 add 1000 units per ml of IL-4 and 800 units per ml of GM-CSF.

A summary of the cytokines added are provided in **Table 1**:

Cytokine	Final Concentration	Stock	Volume to be Added per ml
GM-CSF	800U/ml	2,800U/ml	3.8µl
IL-4	1000U/ml	400U/µl	33µl
TNF-α	10ng/ml	1µl/ml Dilute 1:100 (10ng/ml)	13µl Diluted solution
PGE-1	1 µg/ml	0.5µg/ml	2.7µl
IL-1β	10ng/ml	10ng/µl	10µl
IL-6	100ng/ml	10ng/µl	10 µl

The culture is then incubated for about 2 days at 37°C and 5% carbon dioxide after which the dendritic cells are ready of preparation as stimulators for autologous PBMC.

- 20 The dendritic cells are harvested and counted. Note that cells are lost when washed several times, therefore the wash steps can be omitted when dendritic cell numbers are limiting. Dendritic cells do not divide and the omission of irradiation is not important.

If greater than  $5 \times 10^5$  transduced dendritic cells are recovered send approximately  $10^5$  for phenotyping along with non-transduced dendritic cells if the latter are available.

- 25 If less than  $5 \times 10^5$  transduced dendritic cells are recovered then proceed to peptide loading.

The supernatant is aspirated after centrifugation and the 5µl (5ng) of each peptide library (i.e. for each antigen) per  $1 \times 10^6$  dendritic cells is added. The mixture is then incubated for 30 to 60 minutes in 5% carbon dioxide incubator and then irradiated at 30 Grays.

- 30 The cells are re-suspended in 20mls of medium and centrifuged for about 5 minutes at 400 x G. Re-suspend the cells at  $10^5$  per ml with CTL culture medium. The dendritic cells are now ready for use as stimulators for PBMCs.

As with Example 1 because these cells are to be infused into patients appropriate procedures must be followed.

Recovery of total dendritic cells should be 1 to 8% of the starting population.

**References:**

- Gahn Bei *et al*/ Adenoviral Gene Transfer into Dendritic Cells Efficiently Amplifies the Immune Response to the LMP2A-Antigen; a potential treatment strategy for EBV virus-positive Hodgkin's lymphoma. *Int. J. Cancer* 2001 Sep 1; 93(5): 706-13
- 5 Bollard CM *et al*/The generation and characterisation of LMP2-specific CTLs for use in adoptive transfer from patients with relapsed EBV-positive Hodgkin disease *J. Immunother* (1997). 2004 Jul-Aug; 27(4):317-327
- Gottschalk S *et al*/ Generating CTLs against subdominant Epstein-Barr virus LMPI antigen for the Blood 2003 Mar 1; 101(5): 1905-12.

**PROTOCOL 3: GENERATION OF T-APCs**

1. Coat non tissue culture treated 24 well plates or T-75 non tissue culture treated flasks with CD3 and CD28 antibodies (Miltenyi)
  - 1.1. Calculate the number of wells/flasks to be coated based on the PBMC number to be plated at  $1 \times 10^6$  cells per well or 30-50  $\times 10^6$  cells per T-75 flask
  - 1.2. For wells, plate 0.5 ml of H<sub>2</sub>O containing 1 ng CD3 (5  $\mu$ l/ml of H<sub>2</sub>O of 0.2mg/ml stock) and 1  $\mu$ g CD28 (2L/ml of H<sub>2</sub>O of 0.5mg/ml stock) perwell.
  - 1.3. For T-75 flasks, after prewashing with 18 ml of H<sub>2</sub>O; use 90  $\mu$ l CD3 (stock 0.2mg/ml) and 36  $\mu$ l CD28 (stock 0.5mg/ml) in 18 ml H<sub>2</sub>O per flask.
  - 1.4. Incubate for at least 3 hours in 37°C incubator.
  - 1.5. Optional: CD3-CD28 coated plate can be incubated overnight at 4°C
2. Wash wells or flasks
  - 2.1. Remove CD3/28 solution and rinse once with 1 mL of PBS or medium per well or 10mls per T-75 flask
3. Re-suspend PBMCs at  $2 \times 10^6$  in 30 mL of T cell medium (10% human AB serum, 45% RPMI 1640 (or advanced RPMI), 45% EHA and 200 mM L-glutamine)
  - 3.1. Aliquot 2mls per CD3/28-coated well or 15 to 25 ml to a T-75 flask
  - 3.2. Transfer to incubator for 2 days with flask lying flat.
4. On day 2 add 100 units per mL of IL-2.
5. On day 4 to 5 cells re-suspend cells and count.
6. Split cells according to the cell number obtained
  - 6.1.  $5 \times 10^6$  cells per GRex 100
  - 6.2.  $5 \times 10^7$  per GRex 100.
7. 3 to 4 days later, harvest cells and count
  - 7.1. Either further expand as in step 4
  - 7.2. Or cryopreserve
8. Cryopreservation
  - 8.1. Centrifuge cells and aspirate supernatant
  - 8.2. Flick pellet to re-suspend and transfer to ice for 10 to 30 minutes
  - 8.3. Re-suspend at  $5 \times 10^6$  to  $10 \times 10^6$  cells per ml of ice cold cryopreservation medium (10% DMSO, 50% human AB serum, 40% RPMI 1640)
  - 8.4. Transfer to cryocontainers (cryobags or cryovials)



- 8.5. Freeze in freezing containers at  $\sim -1^{\circ}\text{C}$  per minute for at least 90 minutes, then transfer to liquid nitrogen
9. Prior to use at antigen-presenting cells, T cells must be re-stimulated on CD3/28 antibody-coated plates as in step 3, for 2 to 4 days to upregulate co-stimulatory molecules
  - 9.1. This step is critical as if T cells have a resting phenotype, they may induce T regulatory cells

**PROTOCOL 4: EXPANSION OF AUTOLOGOUS ANTIGEN SPECIFIC T CELLS USING A PREFERRED EMBODIMENT OF THE INVENTION**

1. Re-suspend PBMCs at  $2 \times 10^6$  per mL of T cells medium containing IL-4 and IL-7.  
T cell medium is 10% human AB serum, 45% Advanced RPMI, 45% EHAA and 200mM L-glutamine. IL-4 is added at 20ng per mL (10ng/ml final dilution) and IL7 at 3332 units per mL (1666U/ml final dilution)
2. Re-suspend peptide-coated dendritic cells at  $1 \times 10^5$  to  $2 \times 10^5$  per ml
3. Mix PBMCs and DCs at a 1 : 1 ratio
4. Transfer 20 ml per GRex-10 ( $20 \times 10^6$  PBMCs) and add 10mls additional medium containing IL-4 (1666 units per mL) and IL-7 (10ng per ml). Transfer to incubator for 7 days
5. On day 7 remove 20 ml medium  
Re-suspend cells and count.  
If  $>50 \times 10^6$  cells, split between two GRex-10s. If  $< 50 \times 10^6$  leave in one GRex-10.  
Add medium containing IL-4 and IL-7 to 30mls in each GRex-10 (final concentration of IL-4 is 1666 units per mL and of IL-7 is 10ng per ml)  
Transfer to incubator for 2 to 3 more days
6. On Day 9 or 10 re-stimulate with peptide-coated T-APCs and aK562 cells  
Harvest responder T cells from GRex and count  
Re-suspend at  $1 \times 10^6$  cells per ml  
Transfer to incubator
  - 6.1 Prepare aK562-CS cells to provide a 5:1 aK562-CS to T cell ratio
    - 6.1.1. Irradiate aK562-CS cells with 100 GY (Grays)
    - 6.1.2. Centrifuge for 5 minutes and 400 G
    - 6.1.3. Re-suspend in complete T cell medium
    - 6.1.4. Count and estimate the number of aK562-CS cells required
      - 6.1.4.1. The number of responder T cells x 5
  - 6.2. Prepare peptide-pulsed, irradiated, activated autologous T cells (T-APCs)
    - 6.2.1. Autologous T cells (ATCs) should have been stimulated or re-stimulated with CD3/28 two to 4 days before use
    - 6.2.2. Harvest sufficient number of ATCs for stimulation plus  $\sim 30\%$  to account for loss during processing
    - 6.2.3. Centrifuge cells at 400G for 5 minutes and aspirate supernatant
    - 6.2.4. Loosen cell pellet by finger-flicking
    - 6.2.5. Add 10ng of each peptide per  $10 \times 10^6$  ATCs
    - 6.2.6. Incubate at  $37^{\circ}\text{C}$  in  $5\% \text{CO}_2$  in air for 30 to 90 minutes

- 6.2.7. Re-suspend in ~20 mL medium
- 6.2.8. Irradiate 30 GY
- 6.2.9. Centrifuge 400G for 5 minutes and aspirate supernatant
- 6.2.10. Re-suspend at  $10^5$  cell per mL
- 6.3. Combine  $1 \times 10^7$  responder T cells with  $1 \times 10^7$ T-APCs and  $5 \times 10^7$  irradiated aK562-cs cells per GRex-100
  - 6.3.1. Add medium to 400mL
  - 6.3.2. Add 10ng per mL IL-7 (2mg) and  $6.6 \times 10^5$  units of IL-4
  - 6.3.3. Transfer to incubator for 3 to 4 days
- 7. Add IL-2 (50 to 100 units per mL) or IL-15 10 ng per mL
  - 7.1. Return to culture for 3 to 4 days
  - 7.2. Add cytokines every 3 to 4 days
- 8. Measure glucose from day 7
  - 8.1. Remove 1 drop of medium and test on standard hand held glucometer
  - 8.2. Glucose levels of less than 100 should trigger a change of medium/cytokines
  - 8.3. Responder T cells should be cryopreserved when sufficient number have been obtained
- 9. A third stimulation (steps 7,8 and 9) can be performed if insufficient cells are obtained

**Example 1: Generation of EBNA1, LMP1 & LMP2 specific T cells for healthy donors and patients with nasopharyngeal carcinoma and lymphoma.**

**Sub experiment: Generation of EBNA1, LMP1 & LMP2 specific T cells for lymphoma patient 1.**

**D1** Coating OKT3/CD28 plate

Prepared OKT3 and CD28 antibodies by adding 5ug each of OKT3 and CD28 antibodies to 5mL of sterile water. Added 0.5 mL of this mixture into each well of a non-tissue culture treated 24 well plate (24-w-p). Incubated at 4°C overnight. Note: this is culture day (-)8.

**D2** Generation of dendritic cells from frozen PBMCs by adherence & generation of OKT3 blasts from non-adherent PBMC population - culture day (-)7

Lymphoma patient 1 peripheral blood mononuclear cells (PBMCs) frozen stock taken out from GMP bank and follow up samples with permission of PI (Cath Bollard): 4 vials of PBMCs, all frozen between Feb and March 2010, 5 million each vial, total 20 million. PBMCs were thawed out in 2 vials of 40mL warm CellGenix media and spun down. PBMC count of patient 1: 22.5 million Spun down patient 1 PBMCs and re-suspended in a total of 6mL of warm CellGenix media, then plated PBMC out in 3 wells of a 24-w-p with 2mL per well. After 3 hours the non-adherent portion was gently washed off with sterile PBS twice and media was replaced with CellGenix with IL4 and GM-CSF. Incubated at 37°C. The non-adherent portion was used to generate OKT3 blasts.

**OKT3 blast generation:**

A plate with OKT3 and CD28 coated the day before and stored at 4°C was washed once with 0.5mL of T cell media per well. Non-adherent portion of PBMCs was plated out into ~10 wells. Incubated at 37°C

- D3** Fed OKT3 blasts with IL2 by replacing ½ of the media in the well with CTL Media with 100units of IL2/mL to make final culture concentration of 50units/mL
- D4** Fed DCs with IL4 & GMP-CSF by replacing 1mL of media per well with fresh media with IL4 & GMP-CSF  
 Moved OKT3 blasts into a new tissue culture treated 24-w-p. Incubated at 37°C.
- D7** Transduced 1 well of each patient DC with Ad-LMP1-LMP2 by harvesting cells with scraping, counted, spun down then added Ad-LMP1-LMP2 at MOI of 5000.  
 Patient 1: 0.5 million DC, thus added 0.5ul of virus at 5x10<sup>8</sup> 12vp/mL concentration  
 Flicked tubes every ~15 minutes while incubating at 37°C for 1.5 hrs. Resuspended cells back in media containing GM-CSF, IL4, IL1b, IL6, TNFa, and PGE2 and plated out back into 1 well each of a 24-2-p with 2mL of media.  
 For the condition without Adenoviral vector, ½ media was taken off, and then CellGenix media with 2x of GMP-CSF, IL4, IL1b, IL6, TNFa and PGE2 was added back into the wells. Incubated at 37°C  
 Continued feeding OKT3 blasts by replacing media or splitting cells into extra well.
- D9** DO Setting up first stimulation.  
 Harvested and counted dendritic cells:  
 Patient 1: Non-transduced DCs: 0.2 million; transduced DC: 0.05 million  
 Removed a frozen vial of patient 1 PBMCs from GMP (cells frozen in 2009) to be stimulated, then thawed cells using warm CTL media with 30% FBS:  
 Patient 1: 1 vial of frozen PBMCs at 10 million per vial, recovered 8 million  
 Pulsing of DC and whole PBMCs with pepmixes:  
 - Diluted EBNA1, LMP1 and LMP2 pepmixes by adding 1ul of stock pepmix solution (in DMSO) each into 200ul of sterile PBS.  
 - Spun down all non-transduced DC from both donors and aspirated all but ~50ul of media. Loosened pellets by flicking the tubes. Added ~10ul of diluted pepmix to the pellets, incubated at 37°C for 30 min with occasional flicking.  
 - 3 million of PBMCs from donor 1 were taken to a new clean tube each, spun down, and all but ~50ul of media aspirated. Added ~3 ul of diluted pepmix into the pellet.  
 Incubated at 37°C for 30 min with occasional flicking.
- Setting up T cell culture Day 0 for patient 1:  
 Current GMP condition: Spun down and re-suspended DCs into 0.5mL of media after incubation; took out 1 million of PBMCs, spun down and re-suspended in 0.5 mL of complete CTL media (2 million per mL). Added both into a well of a 48-w-p. Added 5ug of IL15 to make final concentration of 5ng/mL. DC:PBMC ratio of this condition was 1:20. Incubated at 37°C  
 Condition DC(px): Washed DC pulsed with pepmixes with 20mL of PBS. Re-suspended with 2mL of complete CTL media, plated out 1mL per well of a 24-w-p. Took out 4 million of PBMCs, spun down and re-suspended at 2 million per mL. Added IL4 & IL7 at 2x concentration (20ng/mL) to PBMCs. Added 1mL into each well that already had DC to bring final concentration of IL4 and IL7 down to 10ng/mL. DC:PBMC ratio of this condition was 1:20. Incubated at 37°C  
 Condition Px: Re-suspended PBMC pellet in 2mL of complete CTL media, added IL4 and IL7 at 10ng/mL, plated out into 1 well of a 24-w-p. Incubated at 37°C

**D10** Patient 1 OKT3 blasts expanded to 2 full 24-w-p. Count: 65 million. Frozen down in 4 vials with each vial at ~15 million cells.

**D15** Coating OKT3/CD28 plate

Prepared OKT3 and CD28 antibodies by adding 5ug each of OKT3 and CD28 antibodies to 5mL of sterile water. Added 0.5 mL of this mixture into each well of a non-tissue culture treated 24 well plate (24-w-p). Incubated at 4°C overnight.

**D16** Transduced LCL with Ad-LMP1-LMP2

~3million of LCL from donor 1 culture was taken out, spun down, then added 3ul of virus at concentration of  $5 \times 10^{12}$  vp/mL for MOI of 5000. Incubated at 37°C for 1.5 hr with interval flicking of tube. After incubation re-suspended cells in ~5 mL of complete RPMI. Incubated at 37°C

Activated OKT3 blasts:

Thawed out a vial each of OKT3 blasts from donor 1 at 10 million per vial in warm CTL media with 30% FBS. Spun down cells and re-suspended in 20mL of CTL media. Washed a plate of OKT3 & CD28 coated with CTL media, then plated cells out into 10 wells of a 24-w-p. Incubated at 37°C.

**D18 Culture day 10**

Harvested and counted day 10 T cells from patient 1:

Condition GMP: 2.13 million

Condition DC(px): 2.4 million

Condition Px: 3.04 million

Harvested patient 1 OKT3 blasts and irradiated at 30Gy. Count: 13.4 million

Harvested aK562cs and irradiated at 100Gy. Count: 4.4 million. Washed, spun down and re-suspended in 11 mL for a cell concentration of 0.4 million per mL

Harvested patient 1 transduced LCL, irradiated at 40Gy. Count: 5.7 million

Because we didn't have enough cells for testing antigen specificity by ELISpot or phenotyping for lymphocyte subtypes, we only stimulated these cultures.

**Second stimulation:**

Pulsing of OKT3 blasts with pepmixes:

- Diluted EBNA1, LMP1 and LMP2 pepmixes by adding 1ul of stock pepmix solution (in DMSO) each into 200ul of sterile PBS.
- Spun down OKT3 blasts and aspirated all but ~50ul of media. Loosened pellets by flicking the tubes. Added ~20ul of diluted pepmix to the pellets, incubated at 37°C for 30 min with occasional flicking. Washed with 20mL of PBS, spun down.
- Re-suspended OKT3 blasts at concentration of 1million per mL

Condition GMP:

Spun down LCL and re-suspended in 14.25 mL of CTL media for concentration of 0.4 million per mL. Plated out 0.5 mL or 0.2 million LCL into 4 wells of a 24-w-p.

Spun down D9 T cells for GMP condition, re-suspended in 4mL for approximately 0.6 million of T cells per well and added 1mL to each of the 4 wells with LCL. T cell to LCL ratio was about 3:1.

Incubated at 37°C.

Condition DC(px):

Spun down DC(px) day 9 T cells, resuspended in 4mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs into each of the well to make a final volume of 2mL per well, with T cell:OKT3 blasts:aK562cs ratio of 1:1:1 (low ratio of aK562cs due to low total aK562cs available). Incubated at 37°C.

Condition Px:

Spun down Px day 9 T cells, re-suspended in 6mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs into each of the well to make a final volume of 2mL per well, with T cell:OKT3 blasts:aK562cs ratio of 1:1:1 (low ratio of aK562cs due to low total aK562cs available). Incubated at 37°C.

**D21** Added IL2 to culture day 14 by replacing 1mL of media in each well with 1 mL of media with 100 units of IL2 per mL to make final IL2 concentration 50 units/mL.

**D22** Coating OKT3/CD28 plate

Prepared OKT3 and CD28 antibodies by adding 5ug each of OKT3 and CD28 antibodies to 5mL of sterile water. Added 0.5 mL of this mixture into each well of a non-tissue culture treated 24 well plate (24-w-p). Incubated at 4°C overnight.

**D23** Activated patient 1 OKT3 blasts by thawing out a frozen vial with 20 million cells, then plated them on OKT3/CD28 coated plate.

Transduced patient 1 LCL by spinning down 2 million of LCL leaving ~50ul of media left, then added 2ul of Ad-LMP1-LMP2 at  $5 \times 10^{12}$  vp/mL for MOI of 5000. Flicked tube every ~15 min, and after 1.5 hrs re-suspended in 5mL of complete RPMI media.

**D24** Culture day 16: replaced media with fresh media without any cytokines. Incubated at 37°C.

Coated plate for ELISpot assay on D17:

- Prewetted a 96-w immobilon-P membrane plate with 50ul of 35% EtOH per well. Washed with PBS.

- Made IFN $\gamma$  solution by adding 100ug purified mouse anti-human IFN $\gamma$  1-DIK antibody to 10mL of coating buffer. Added 100ul per well to coat plate. Store at 4°C overnight.

**D25 Culture Day 17**

Harvested and counted patient 1 day 17 T cells:

Condition GMP: 8.2 million

Condition DC(px): 5.5 million

Condition Px: 15.6 million

Harvested, irradiated at 30Gy and counted OKT3 blasts: 13.6 million

Harvested, irradiated at 40Gy and counted patient 1 LCL: 1.2 million

Harvested, irradiated at 100Gy and counted aK562cs: 22.5 million.

**Setting up ELISpot assay to detect IFN $\gamma$  release:**

- Blocked a 96-w-plate (with membrane coated with IFN $\gamma$  primary antibody from the day before) with T cell media for 1 hr at 37°C.

- Prepared responder cells: Took out ~1.5 million of T cells from GMP condition, 0.8 million from DC(px) condition, and 0.8 million from Px condition, spun down, and re-suspended at 0.5 million per mL.

- Prepared pepmix solution by adding 4ul of pepmix stock in DMSO (0.2ug/ul) into 800ul of T cell media the following antigens: EBNA1, LMP1, LMP2, and added 1.5ul of pepmix stock in DMSO into 300ul of T cell media of the following antigens EBNA3a, EBNA3b, EBNA3c, Bzlf1, NY-ESO1 (irrelevant antigen for negative control), staph aureus super-antigen (positive control). Prepared patient 1 LCL by taking out ~1 million of cells, spun down, resuspended in 1 million per mL. Added these antigens and targets to the responder T cells as following with each space represents 2 wells (duplicates):

GMP condition	EBNA1 Bzlf1	LMP1 LCL	LMP2 (-)ve control	EBNA3a (+)ve control	EBNA3b	EBNA3c
DC(px)	EBNA1	LMP1	LMP2	LCL	(-)ve control	(+)ve control
Px	EBNA1	LMP1	LMP2	LCL	(-)ve control	(+)ve control

-Added 100ul or 50,000 cells into each well, with GMP 24 wells, DC(px) and Px each with 12 wells.

- Incubated overnight at 37°C.

**OPTIONAL STEPS**

**Third stimulation :**

Pulsing of OKT3 blasts with pepmixes:

- Diluted EBNA1, LMP1 and LMP2 pepmixes by adding 1ul of stock pepmix solution (in DMSO) each into 200ul of sterile PBS.
- Spun down OKT3 blasts and aspirated all but ~50ul of media. Loosened pellets by flicking the tubes. Added ~20ul of diluted pepmix to the pellets, incubated at 37°C for 30 min with occasional flicking. Washed with 20mL of PBS, spun down.
- Resuspended OKT3 blasts at concentration of 1million per mL, then further dilute out to 0.4 million per mL

Condition GMP:

Spun down irradiated LCL and re-suspended in 10 mL of CTL media for concentration of 0.12 million per mL. Plated out 0.1 million or 1mL each into 6 wells of a 24-w-p.

Spun down 3 million of D17 T cells for GMP condition, re-suspended in 6mL for and added 1mL to each of the 6 wells with LCL. T cell to LCL ratio was about 4:1. Incubated at 37°C. Froze down the rest of T cells

Condition DC(px):

Spun down 2 million of DC(px) day 17 T cells, re-suspended in 10mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs into each of the well to make a final volume of 2mL per well, with T cell : OKT3 blasts:aK562cs ratio of 2:2:5. Incubated at 37°C.

(increased T cell and OKT3 blasts concentration since the growth after second stimulation was not great). Froze down the rest of T cells

Condition Px:

Spun down 2 million of Px day 17 T cells, re-suspended in 10mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs into each of the well to make a final volume of 2mL per well, with T cell : OKT3 blasts:aK562cs ratio of 2:2:5. Incubated at 37°C. (Increased T

cell and OKT3 blasts concentration to match with DC(px) condition). Froze down the rest of T cells.

**D29** Developed ELISpot IFN $\gamma$  plate.

Prepared secondary IFN $\gamma$  antibody (7B6-1 biotin) by adding 100 $\mu$ L of antibody to 10mL of PBS+0.5% BSA, then filtered through a 0.2 $\mu$ m filter to get rid of clumps of conjugated antibodies to prevent non-specific binding. Washed plate 6x with 100 $\mu$ L of PBS + 0.05% Tween per well each time. Added 100 $\mu$ L of this secondary antibody solution to each well. Incubated at 37°C for 2 hours.

After 1.5 hours prepared 10mL of Avidin-Peroxidase complex solution in PBS+0.05% Tween, mixed, and incubated at room temperature. At the end of the 2hr, washed plate 6x with 100 $\mu$ L PBS + 0.05% Tween per well each time. Added 100 $\mu$ L Avidin-Peroxidase complex solution to each well, incubated at room temperature for 1 hour.

At the end of the incubation, prepared AEC substrate by first dissolving AEC tablet into 2.5mL Dimethylformamide, then added 47.5mL acetate buffer (4.6mL 0.1N acetic acid, 11mL sodium acetate, 47mL water), next added 25 $\mu$ L 30% hydrogen peroxide, mixed & filtered with a 0.45 $\mu$ m filter. Washed plates with PBS+0.05% Tween, repeated 3x, washed with PBS, repeated 3x, added AEC substrate and let develop for up to 4 minutes. Stopped reaction by rinsing with water. Peeled of the back and dry the membranes. Punch out results and sent for counting.

**D31** Added IL2 to culture day 20 by replacing 1mL of media in each well with 1 mL of media with 100 units of IL2 per mL to make final IL2 concentration 50 units/mL.

**D32** Coating OKT3/CD28 plate

Prepared OKT3 and CD28 antibodies by adding 5 $\mu$ g each of OKT3 and CD28 antibodies to 5mL of sterile water. Added 0.5 mL of this mixture into each well of a non-tissue culture treated 24 well plate (24-w-p). Incubated at 4°C overnight.

**D33** Activated patient 1 OKT3 blasts by plated on-going patient 1 OKT3 blast culture on OKT3/CD28 coated plate at ~1 million per well.

Transduced patient 1 LCL by spinning down 2 million of LCL leaving ~50 $\mu$ L of media left, then adding 2 $\mu$ L of Ad-LMP1-LMP2 at 5x10<sup>6</sup> 12vp/mL for MOI of 5000. Flicked tube every ~15 min, and after 1.5 hrs re-suspended in 5mL of complete RPMI media.

**D34** Culture day 23: replaced media with fresh media without any cytokines and split any wells that is over ~4 million cells. Incubated at 37°C.

Coated plate for ELISpot assay:

- Prewetted a 96-w immobilon-P membrane plate with 50 $\mu$ L of 35% EtOH per well. Washed with PBS.

- Prepared IFN $\gamma$  solution by adding 100 $\mu$ g purified mouse anti-human IFN $\gamma$  1-D1K antibody to 10mL of coating buffer. Added 100 $\mu$ L per well to coat plate. Stored at 4°C overnight.

**D35 Culture day 24**

Harvested patient 1 day 24 T cell culture and counted:

Condition GMP: 22.2 million

Condition DC(px): 39.8 million

Condition Px: 16.2 million

Harvested OKT3 blasts, irradiated at 30Gy and counted: 9.8 million

Harvested patient 1 LCL, irradiated at 40Gy and counted: 2.3 million

Harvested aK562cs, irradiated at 100Gy and counted: 20 million

**Setting up ELISpot assay to detect IFN $\gamma$  release:**

- Blocked a coated 96-w-plate with T cell media for 1 hr at 37°C.
  - Prepared responder cells: Took out ~2.5 million each of T cells from GMP condition, DC(px) condition, and Px condition, spun down, and resuspended at 1 million per mL.
  - Prepared pepmix solution by adding 4ul of pepmix stock in DMSO (0.2ug/ul) into 800ul of T cell media the following antigens: EBNA1, LMP1, LMP2, and adding 1.5ul of pepmix stock in DMSO into 300ul of T cell media of the following antigens: EBNA3a, EBNA3b, EBNA3c, Bzlf1, NY-ESO1 (irrelevant antigen for negative control), staph aureus super antigen (positive control).
- Prepared patient 1 LCL by taking out ~1 million of cells, spun down, re-suspended in 1 million per mL. Added these antigens and targets to the responder T cells as following with each space represents 2 wells (duplicates):

GMP condition	EBNA1	LMP1	LMP2	EBNA3a	EBNA3b	EBNA3c
	Bzlf1	LCL	(- )ve control	(+ )ve control		
DC(px)	EBNA1	LMP1	LMP2	LCL	(- )ve control	(+ )ve control
Px	EBNA1	LMP1	LMP2	LCL	(- )vecontrol	(+ )ve control

- Added 100ul or 50,000 cells into each well, with GMP 24 wells, DC(px) and Px each with 12 wells.

- Incubated overnight at 37°C.

**Setting up co-culture of lymphoma patient 1 T cells and autologous LCL**

For each condition, added T cell and LCL to a well of a 24-w-p to the ratios below:

T cell to LCL ratio	40:1	20:1	10:1	5:1	1:1	1:1 allo LCL
T cell count(million)	0.5	0.5	0.5	0.5	0.5	0.5
LCL cell count(million)	0.0125	0.025	0.05	0.1	0.5	0.5

Added IL2 to the culture for final concentration of 50u/mL and mixed gently with transfer pipet. Phenotyped day 0 by taking out 200ul of each culture, washed with 3mL of PBS+0.5% FBS, spun down, aspirated supernatant and added 5ul each of the following antibodies: CD19-PE, CD56-FITC, and CD3-PerCP. Incubated at 4°C for 1 hour. Washed with 3mL PBS, spun down, aspirated supernatant and added 250ul of cytofix per tube and 50ul of CountBright beads. Analyzed with a flow cytometer.

**Fourth stimulation:**

Pulsing of OKT3 blasts with pepmixes:

- Diluted EBNA1, LMP1 and LMP2 pepmixes by adding 1ul of stock pepmix solution (in DMSO) each into 200ul of sterile PBS.
- Spun down OKT3 blasts and aspirated all but ~50ul of media. Loosened pellets by flicking the tubes. Added ~20ul of diluted pepmix to the pellets, incubated at 37°C for 30 min with occasional flicking. Washed with 20mL of PBS, spun down.
- Re-suspended OKT3 blasts at concentration of 1million per mL, then further diluted out to 0.4 million per mL
- Resuspended aK562cs at 2 million per mL



Condition GMP:

Spun down irradiated LCL and re-suspended in 18.4 mL of CTL media for concentration of 0.12 million per mL. Plated out 0.12 million or 1mL each into 10 wells of a 24-w-p. Spun down 5 million of D24 T cells for GMP condition, re-suspended in 10mL for and added 1mL to each of the 10 wells with LCL. T cell to LCL ratio was about 4:1. Incubated at 37°C. Froze down the rest of T cells

Condition DC(px):

Spun down 2 million of DC(px) day 24 T cells, re-suspended in 10mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs at 2 million per mL into each of the well to make a final volume of 2mL per well, with T cell: OKT3 blasts:aK562cs ratio of 1:1:5. Incubated at 37°C. Froze down the rest of T cells

Condition Px:

Spun down 2 million of Px day 24 T cells, re-suspended in 10mL of T cell media with 2x IL4 and IL7 concentration at 20ng/mL. Plated out 1mL per well of a 24-w-p. Added 0.5 mL of pepmix pulsed OKT3 blasts, then added 0.5 mL of aK562cs into each of the well to make a final volume of 2mL per well, with T cell: OKT3 blasts:aK562cs ratio of 1:1:5. Incubated at 37°C. Frozen down the rest of T cells.

#### **D36 Developed ELISpot IFN $\gamma$ plate.**

Prepared secondary IFN $\gamma$  antibody (7B6-1 biotin) by adding 100ul of antibody to 10mL of PBS+0.5% BSA, then filtered through a 0.2ul filter to get rid of clumps of conjugated antibodies to prevent non-specific binding. Washed plate 6x with 100ul of PBS + 0.05% Tween per well each time. Added 100ul of this secondary antibody solution to each well. Incubated at 37°C for 2 hours.

After 1.5 hours prepared 10mL of Avidin-Peroxidase complex solution in PBS+0.05% Tween, mixed, and incubated at room temperature. At the end of the 2hr, washed plate 6x with 100ul PBS + 0.05% Tween per well each time. Added 100ul Avidin-Peroxidase complex solution to each well, incubated at room temperature for 1 hour.

At the end of the incubation, prepared AEC substrate by first dissolving AEC tablet into 2.5mL Dimethylformamide, then added 47.5mL acetate buffer (4.6mL 0.1N acetic acid, 11mL sodium acetate, 47mL water), next added 25ul 30% hydrogen peroxide, mixed & filtered with a 0.45ul filter. Washed plates with PBS+0.05% Tween, repeated 3x, washed with PBS, repeated 3x, added AEC substrate and let develop for up to 4 minutes. Stopped reaction by rinsing with water. Peeled of the back and dry the membranes. Punched out results and sent for counting.

#### **Co-culture day 2 phenotyping**

- Replaced 1mL of media with 1mL of fresh t cell media with 100units of IL2.
- Phenotyped day 2 by taking out 200ul of each condition, wash with 3mL of PBS+0.5% FBS, spun down, aspirated supernatant and added 5ul each of the following antibodies: CD19-PE, CD56-FITC, and CD3-PerCP. Incubated at 4°C for 1 hour. Wash with 3mL PBS, spun down, aspirated supernatant and added 250ul of cytofix per tube and 50ul of CountBright beads. Analyzed with a flow cytometer.

#### **D38 Co-culture day 4 phenotyping**

Took out 200ul of each condition, washed with 3mL of PBS+0.5% FBS, spun down, aspirated supernatant and added IOul each of the following antibodies: CD19-PE, CD56-FITC, and CD3-PerCP (increased antibody amount to match with increased cell counts). Incubated at 4°C for 1 hour. Washed with 3ml\_ PBS, spun down, aspirated supernatant and added 250ul of cytofix per tube and 50ul of CountBright beads. Analyzed with a flow cytometer.

Added IL2 to culture day 20 by replacing 1mL of media in each well with 1 mL of media with IOUnits of IL2 per mL to make final IL-2 concentration 50units/mL. Spit confluent wells.

**D39 Co-culture day 5 phenotyping**

Took out 200ul of each condition, washed with 3mL of PBS+0.5% FBS, spun down, aspirated supernatant and added IOul each of the following antibodies: CD19-PE, CD56-FITC, and CD3-PerCP. Incubated at 4°C for 1 hour. Washed with 3mL PBS, spun down, aspirated supernatant and added 250ul of cytofix per tube and 50ul of CountBright beads. Analyzed with a flow cytometer.

**D41** Culture day 30: replaced media with fresh media without any cytokines and split any wells that is over ~4 million cells. Incubated at 37°C.

Coated plate for ELISpot assay:

- Prewetted a 96-w immobilon-P membrane plate with 50ul of 35% EtOH per well. Washed with PBS.
- Prepared IFN $\gamma$  solution by adding IOOug purified mouse anti-human IFN $\gamma$  1-D1K antibody to IOmL of coating buffer. Added IOOul per well to coat plate. Stored at 4°C overnight.

**D42 Culture day 31**

Harvested patient 1 day 31 T cell culture and counted:

Condition GMP: 17.2 million; Condition DC(px): 20.6 million; Condition Px: 26.8 million

**RESULTS**

**Table 2: Whole culture expansion:** Compared to current standard protocol (GMP), cultures set up with KATpx expanded just as well, if not better, by the end of the 4<sup>th</sup> stimulation.

**Table 2: Growth in million of cells:**

	D0	D9	D16	D23	D30
<b>Ad-DC</b>	1	2.13	8.317143	61.54686	211.7212
<b>DC</b>	1	0.6	1.375	27.3625	281.8338
<b>Px</b>	1	1.013333	5.269333	42.6816	571.9334

We have optimized a novel antigen presentation complex, KATpx, which produced equally good or better expansion and higher T cell antigen specific frequencies against EBNA1, LMP1 and LMP2 in a lymphoma patient. These cells efficiently eliminate tumor cells in co-culture and this killing was HLA specific. Moreover, using this approach we eliminated the need for LCL and Adenoviral vector, thus reducing generation time as well as activation of T cells specific for bystander antigens expressed by LCL and Adenoviral vector.

It is envisaged that more than one embodiment described herein may be combined, as technically appropriate. In the context of this specification "comprising" is to be interpreted as "including".

Aspects of the disclosure comprising certain elements are also intended to extend to alternative embodiments "consisting" or "consisting essentially" of the relevant elements. All references referred to herein are specifically incorporated by reference.

**Claims:**

1. A process for *in vitro* expansion of antigen specific T cells, such as autologous antigen specific T cells comprising the steps:
  - a) culturing a population of autologous PBMC cells in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR a peptide/peptide mix relevant to a target antigen(s), and
    - ii) at least one cytokine, and
  - b) culturing a population of T cells from step a) in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) and an artificial co-stimulatory factor, and
    - ii) optionally a cytokine, andcharacterized in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoding antigens in the expansion of the relevant T cell population.
2. A process according to claim 1 wherein step b) is performed two or more times (such as 3, 4, 5 times or more) until sufficient quantities of the relevant T cell population are obtained.
3. A process according to claim 1 or 2 wherein the culturing of step a) is performed for 12 days or less.
4. A process according to any one of claims 1 to 3, wherein the culturing step of step b) is performed for 12 days or less.
5. A process according to any one of claims 1 to 4, wherein culturing is performed in a vessel comprising a gas permeable culture surface.
6. A process according to any one of claims 1 to 5, wherein the T cells expanded are specific to an Epstein-Barr Virus, Vaccinia Virus or Varicella Zoster Virus antigen or antigens, such as EBV.
7. A process according to any one of claims 1 to 6, wherein the peptides of step a) and/or b) comprise between 2 and 1000 peptides, for example 2 to 500 peptides, 2 to 300 peptides or 2 to 100 peptides.
8. A process according to any one of claims 1 to 7, wherein the peptides of step a) and/or b) overlap, for example by 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more amino acids.
9. A process according to any one of claims 1 to 8, wherein the peptides of step a) and/or b) are about 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20 amino acids in length, such as 15 amino acids in length.
10. A process according to any one of claims 1 to 9, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP1.
11. A process according to any one of claims 1 to 10, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP2.
12. A process according to any one of claims 1 to 11, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen EBNA1.
13. A process according to any one of claims 1 to 12, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen BARF1.

14. A process according to any one of claims 1 to 13, wherein the cytokine present in step a) is IL-4.
15. A process according to any one of claims 1 to 14, wherein the cytokine present in step a) is IL-7.
16. A process according to any one of claims 1 to 15, wherein the cytokine present in step b) is IL-15.
17. A process according to any one of claims 1 to 15, wherein the artificial co-stimulatory is an engineered cell line, such as an aK562 cell, with one or more relevant protein or protein fragments present on the cell surface.
18. A process according to claim 17, wherein the protein or protein fragments are independently selected from CD80, CD86, CD83, OX-40 ligand and 41BB-ligand.
19. A process according to claim 18, wherein all of the said protein or protein fragments are present on the surface of the co-stimulatory cell.
20. A process according to any one of claims 1 to 19, wherein the process is performed in a GRex™ system.
21. An expanded autologous or allogenic T cell product obtainable or obtained from a process as defined in any one of claims 1 to 20.
22. An expanded autologous or allogenic T cell product according to claim 21, wherein one or more target cell populations has comparable target antigen specificity compared to cells expanded employing viral vectors and re-stimulated with EBV-LCLs that over-express certain target antigens.
23. An expanded autologous or allogenic T cell product according to claim 21 or 22, wherein one or more of target cell populations has higher target-antigen-specificity at a given timepoint than cells expanded employing viral vectors and re-stimulated with EBV-LCLs that over-express certain target antigens.
24. A pharmaceutical composition comprising an expanded T cell product as defined in any one of claims 21 to 23, and an excipient, carrier, stabiliser, surfactant or other non-therapeutically active component.
25. A method of treating a patient, such as an immunosuppressed patient, for viral infection or cancer induced by a virus by administration of an appropriate dose of an expanded T cell product as defined in any one of claims 21 to 23 or pharmaceutical composition as defined in claim 24.
26. Use of an engineered cell line such as an aK562 cell as an artificial co-stimulatory factor without concomitant presentation of antigen thereon, in the antigen specific expansion of T cells, such as autologous T cells.
27. Use of dendritic cells pulsed with antigen-peptide/peptide mixes for the expansion of antigen specific T cells.
28. Use of T-APCs pulsed with antigen-peptide/peptide mixes for the expansion of antigen specific T cells.
29. Use of antigen-peptide mixes with PBMCs for the expansion of antigen specific T cells.

## AMENDED CLAIMS

received by the International Bureau on 05 November 2012 (05.11.2012)

## Claims:

1. A process for *in vitro* expansion of antigen specific T cells, such as autologous antigen specific T cells comprising the steps:
  - a) culturing a population of autologous PBMC cells in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR a peptide/peptide mix relevant to a target antigen(s), and
    - ii) at least one cytokine, and
  - b) culturing a population of T cells from step a) in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) and an artificial co-stimulatory factor, and
    - ii) optionally a cytokine, and

characterized in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoding antigens in the expansion of the relevant T cell population.
2. A process according to claim 1 wherein step b) is performed two or more times (such as 3, 4, 5 times or more) until sufficient quantities of the relevant T cell population are obtained.
3. A process according to claim 1 or 2 wherein the culturing of step a) is performed for 12 days or less.
4. A process according to any one of claims 1 to 3, wherein the culturing step of step b) is performed for 12 days or less.
5. A process according to any one of claims 1 to 4, wherein culturing is performed in a vessel comprising a gas permeable culture surface.
6. A process according to any one of claims 1 to 5, wherein the T cells expanded are specific to a viral antigen or antigens of Epstein-Barr Virus, Vaccinia Virus or Varicella Zoster Virus, such as EBV.
7. A process according to any one of claims 1 to 6, wherein the peptides of step a) and/or b) comprise between 2 and 1000 peptides, for example 2 to 500 peptides, 2 to 300 peptides or 2 to 100 peptides.
8. A process according to any one of claims 1 to 7, wherein the peptides of step a) and/or b) overlap, for example by 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more amino acids.
9. A process according to any one of claims 1 to 8, wherein the peptides of step a) and/or b) are about 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20 amino acids in length, such as 15 amino acids in length.
10. A process according to any one of claims 1 to 9, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP1.
11. A process according to any one of claims 1 to 10, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP2.
12. A process according to any one of claims 1 to 11, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen EBNA1.
13. A process according to any one of claims 1 to 12, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen BARF1.

## Claims:

1. A process for *in vitro* expansion of antigen specific T cells, such as autologous antigen specific T cells comprising the steps:
  - a) culturing a population of autologous PBMC cells in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR a peptide/peptide mix relevant to a target antigen(s), and
    - ii) at least one cytokine, and
  - b) culturing a population of T cells from step a) in the presence of:
    - i) dendritic cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) OR autologous antigen presenting T cells (T-APC's) cells which have been pulsed with a peptide/peptide mix relevant to a target antigen(s) and an artificial co-stimulatory factor, and
    - ii) optionally a cytokine, and

characterized in that the process does not employ live virus and/or viral vectors or the use of DNA or RNA encoding antigens in the expansion of the relevant T cell population.
2. A process according to claim 1 wherein step b) is performed two or more times (such as 3, 4, 5 times or more) until sufficient quantities of the relevant T cell population are obtained.
3. A process according to claim 1 or 2 wherein the culturing of step a) is performed for 12 days or less.
4. A process according to any one of claims 1 to 3, wherein the culturing step of step b) is performed for 12 days or less.
5. A process according to any one of claims 1 to 4, wherein culturing is performed in a vessel comprising a gas permeable culture surface.
6. A process according to any one of claims 1 to 5, wherein the T cells expanded are specific to a «viral antigen or antigens of Epstein-Barr Virus, Vaccinia Virus or Varicella Zoster Virus-antigen»-or antigens, such as EBV.
7. A process according to any one of claims 1 to 6, wherein the peptides of step a) and/or b) comprise between 2 and 1000 peptides, for example 2 to 500 peptides, 2 to 300 peptides or 2 to 100 peptides.
8. A process according to any one of claims 1 to 7, wherein the peptides of step a) and/or b) overlap, for example by 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15 or more amino acids.
9. A process according to any one of claims 1 to 8, wherein the peptides of step a) and/or b) are about 10, 11, 12, 13, 14, 15, 16, 17, 18, 19 or 20 amino acids in length, such as 15 amino acids in length.
10. A process according to any one of claims 1 to 9, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP1.
11. A process according to any one of claims 1 to 10, wherein the peptides of part a) and/or b) cover part or the full length of the antigen LMP2.
12. A process according to any one of claims 1 to 11, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen EBNA1.

13. A process according to any one of claims 1 to 12, wherein the peptides of part a) and/or b) cover part of or the full length of the antigen BARFI.

FIGURE 1: The prior art process for generating EVB specific T cells

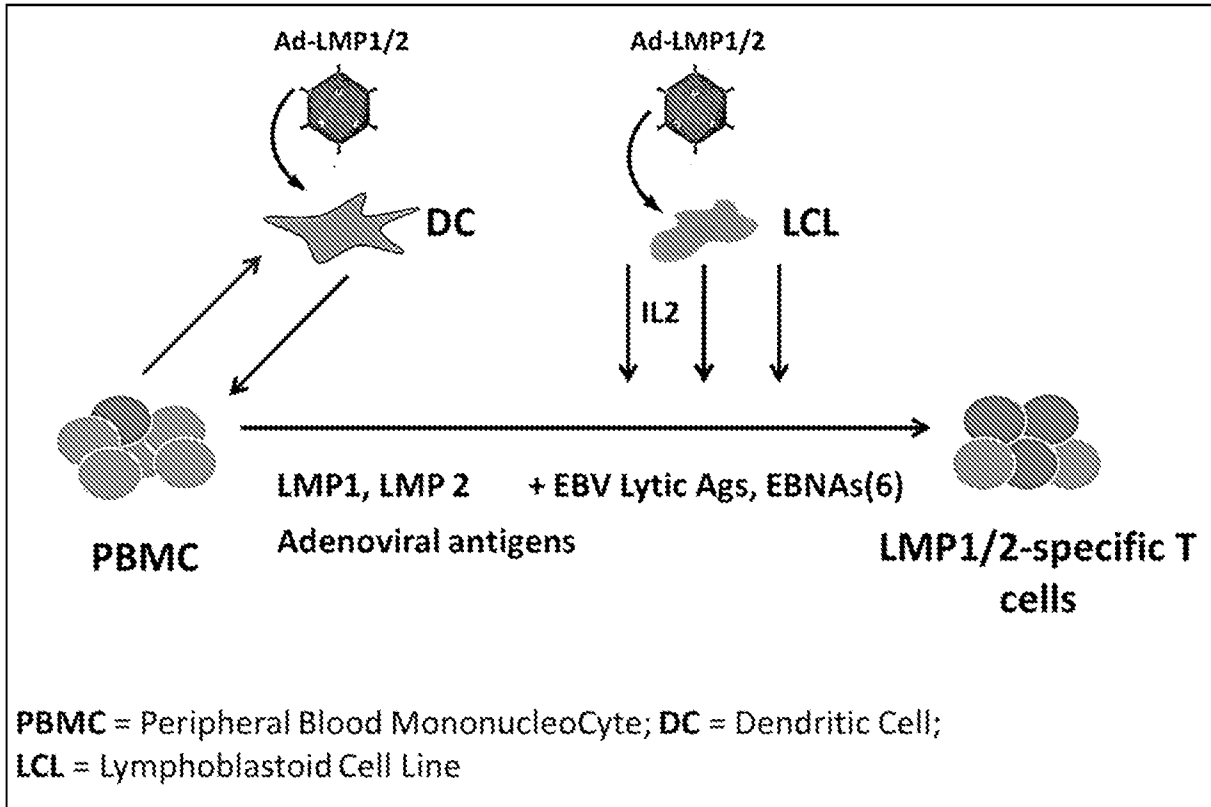
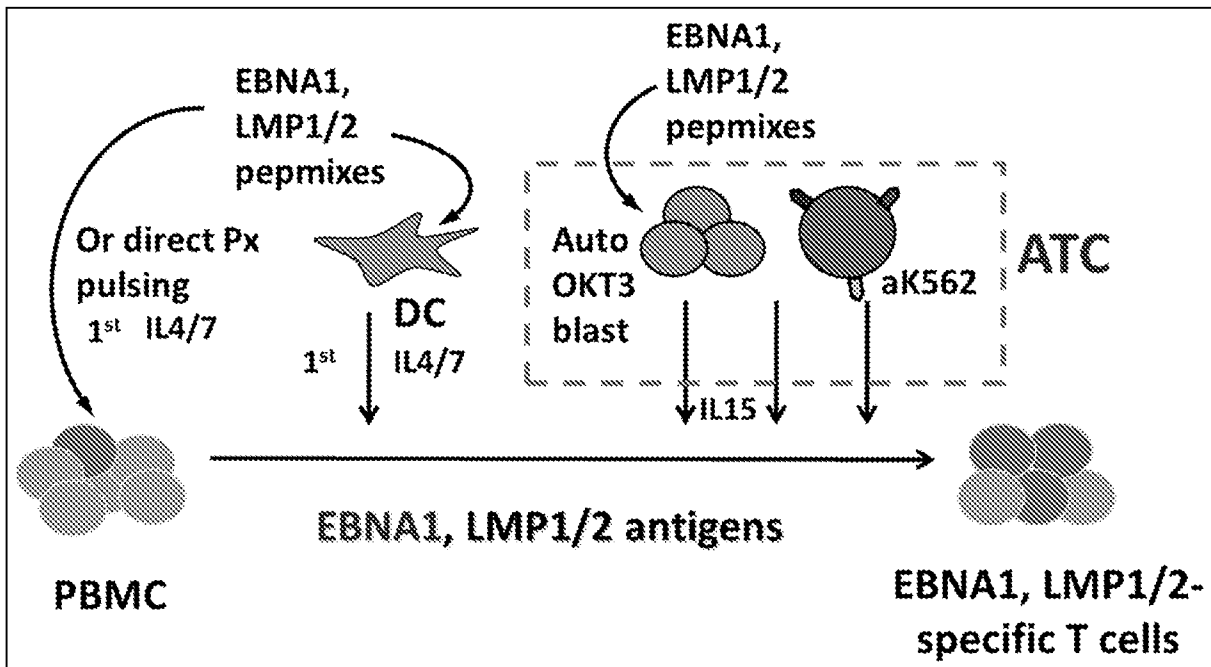
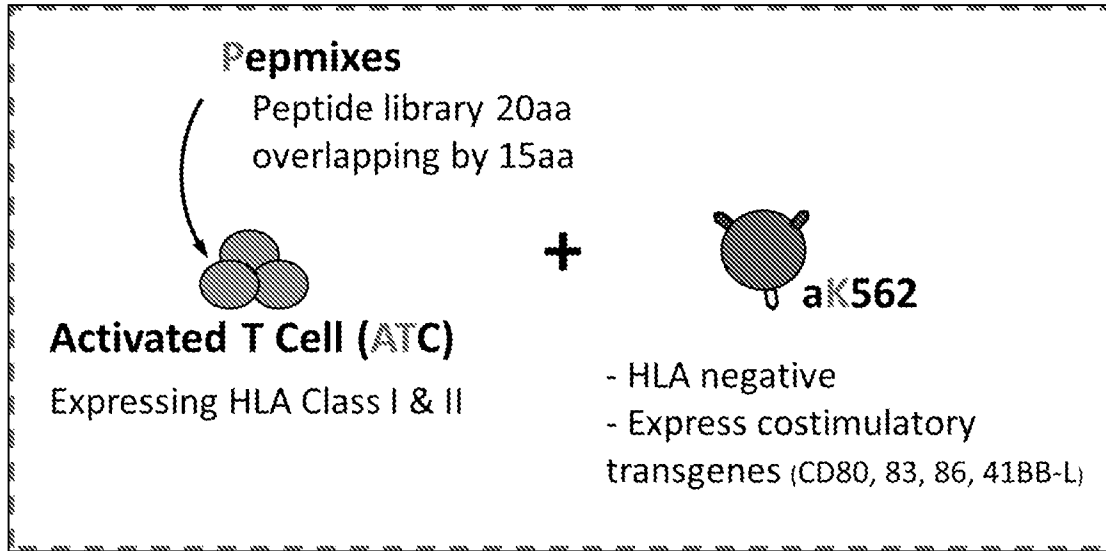


FIGURE 2: Diagram of the alternative methods demonstrated in this application of novel superior protocols for the generation of EBV specific T cells.





**FIGURE 3:** Diagram showing the combination of T-APC (activated T cells) and aK562 that is employed as the second stimulation in the invention.



**Figure 3A:** The expansion of specific T cells using aK562 and T-APC showing the different ratios used in step b) of the invention.

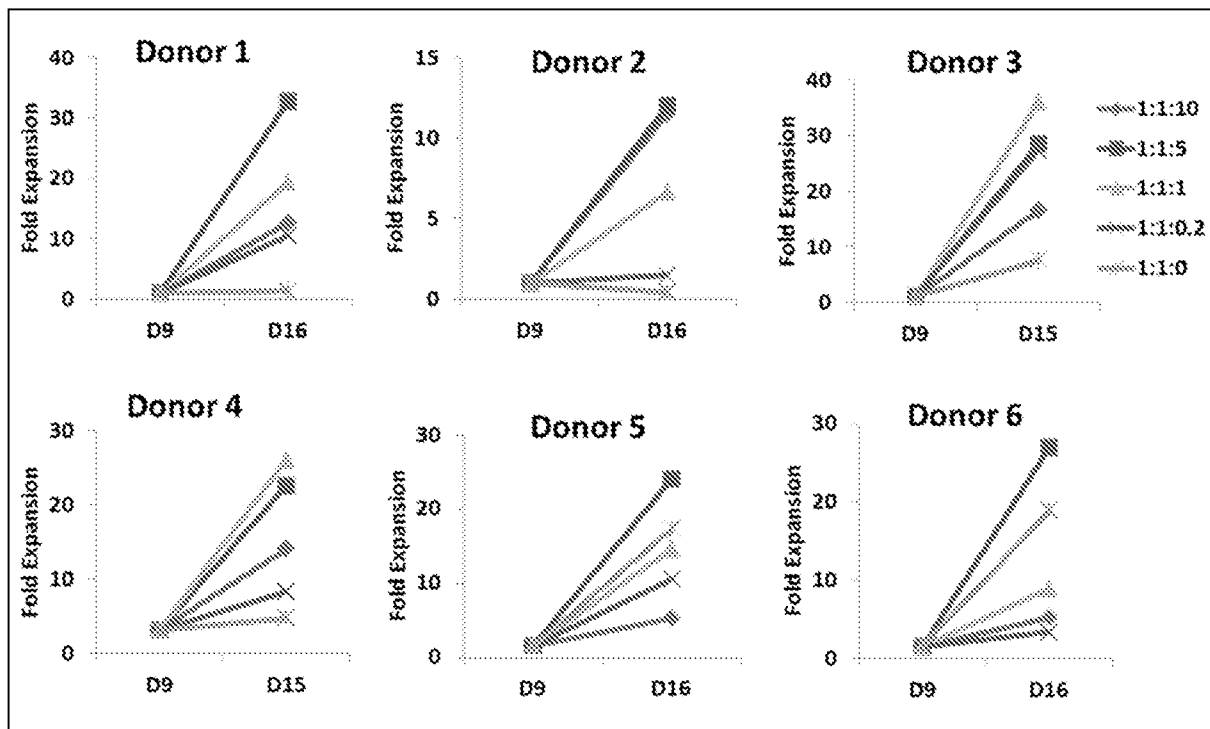


Figure 3B: The optimal CTL (T cell) to T-APC (OKT3 Blast) to aK562 ratio for T cell expansion in step b) of the invention.

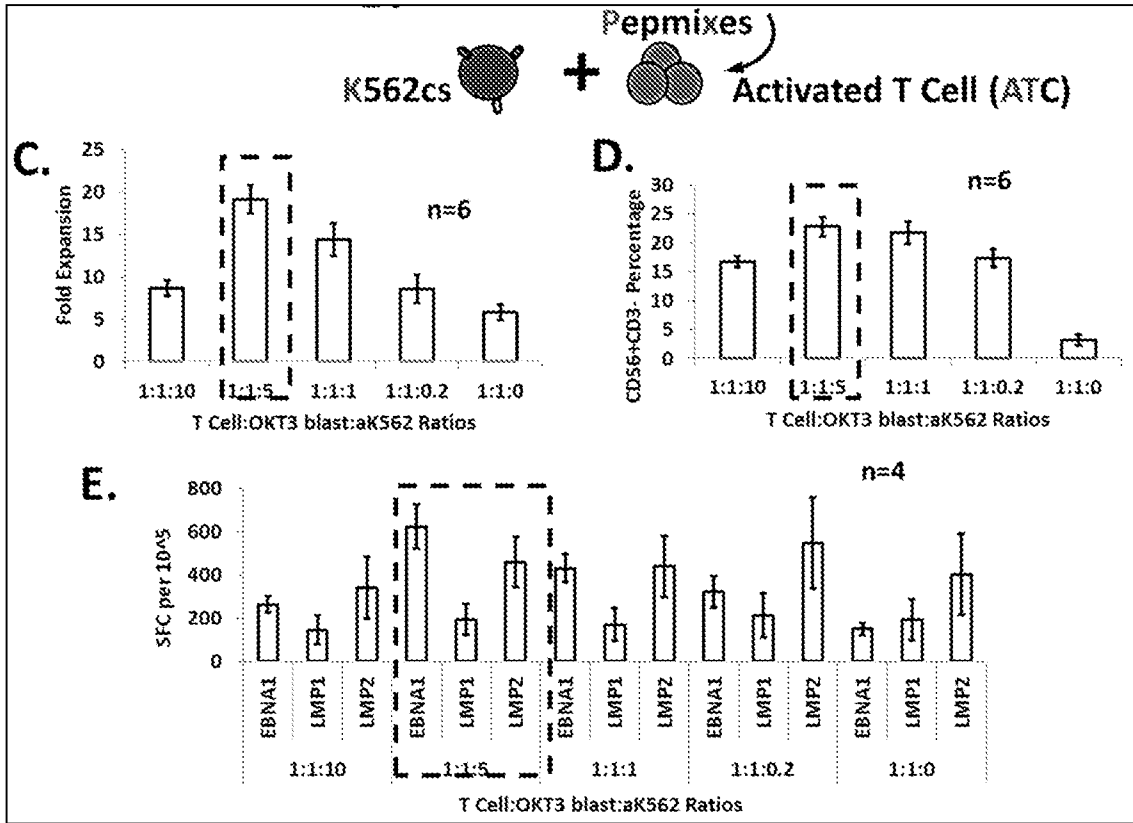


Figure 3C: A more detailed analysis of individual donors from figure 3B (E) showing specific IFN $\gamma$  production by expanded CTLs from 2 donors.

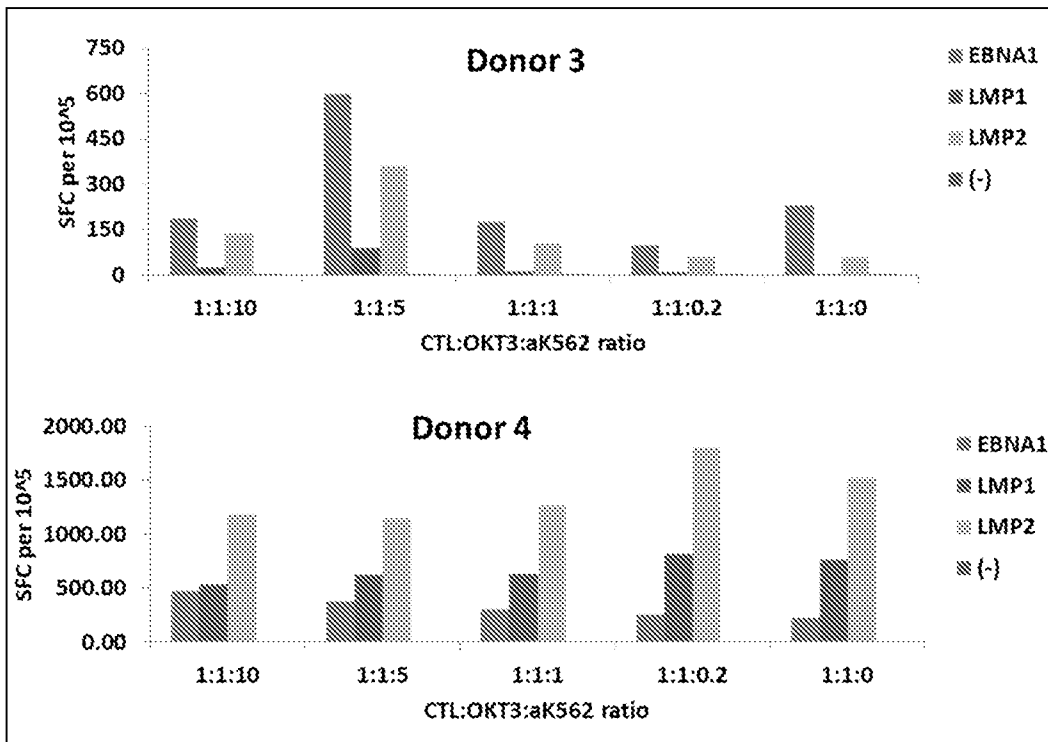


Figure 3D: T-APC CAN Act As Antigen Presenting Cells for Class I and II

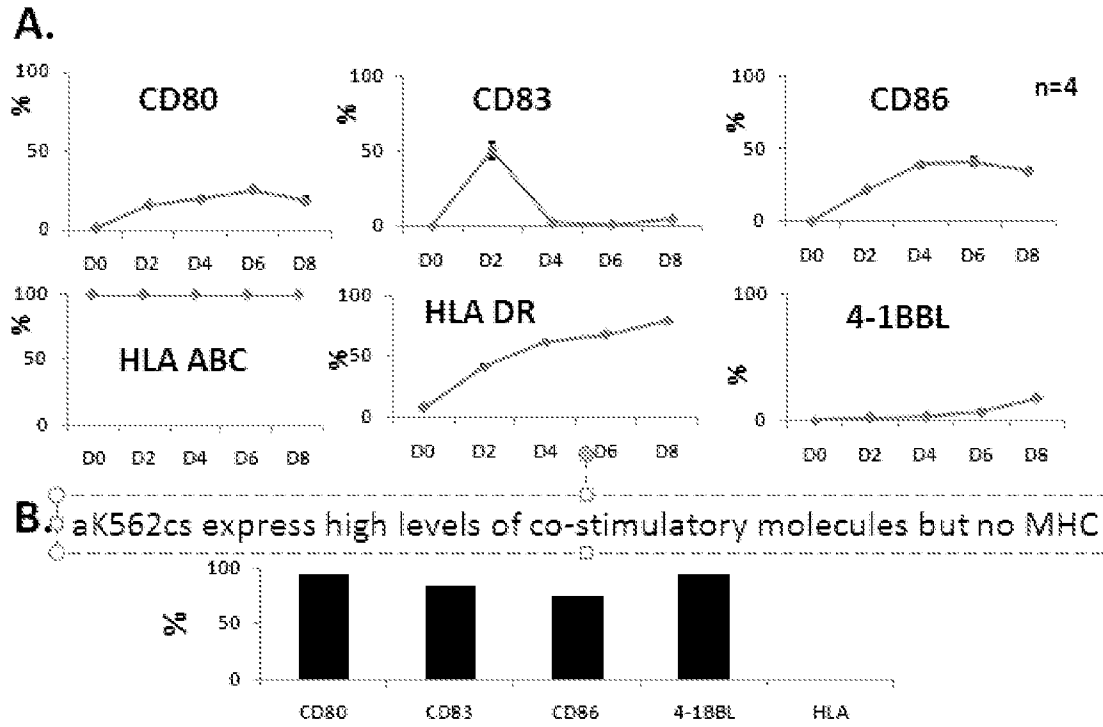


FIGURE 4A: A Schematic representation of the main stimulation steps used in the prior art process and the various embodiments of the invention.

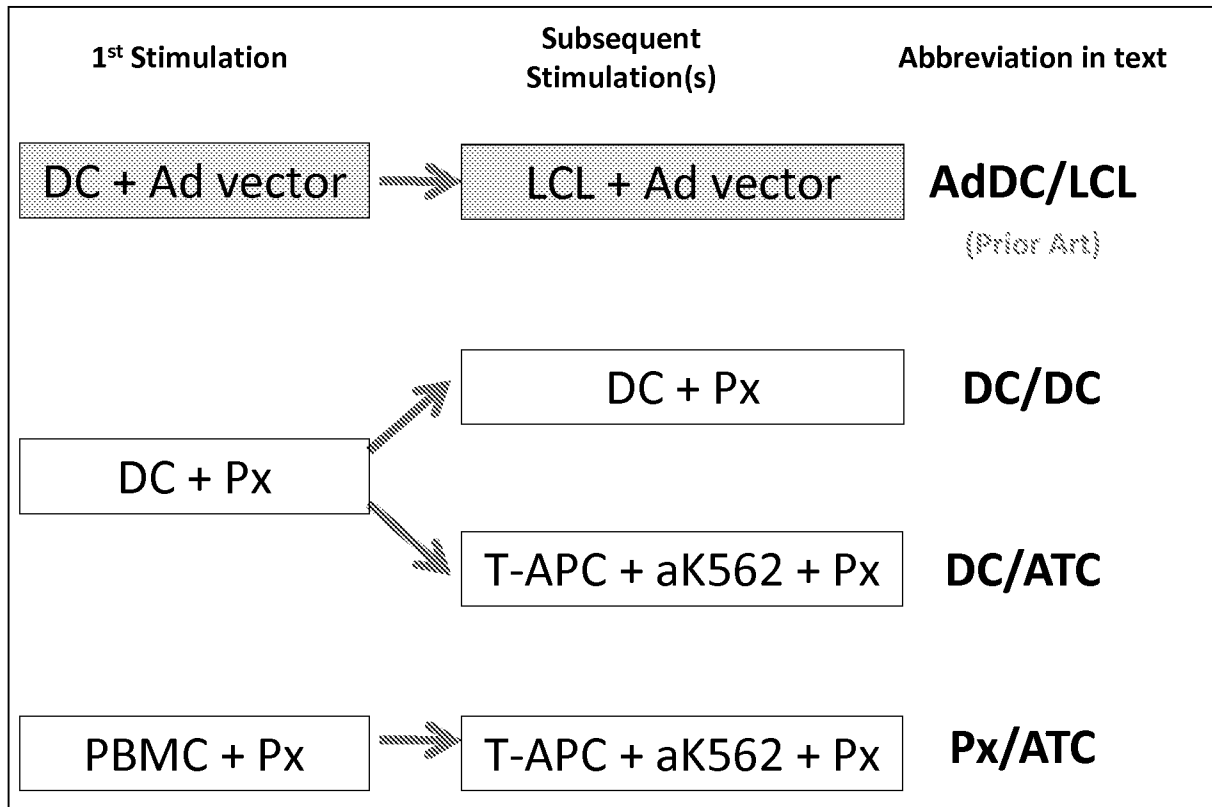
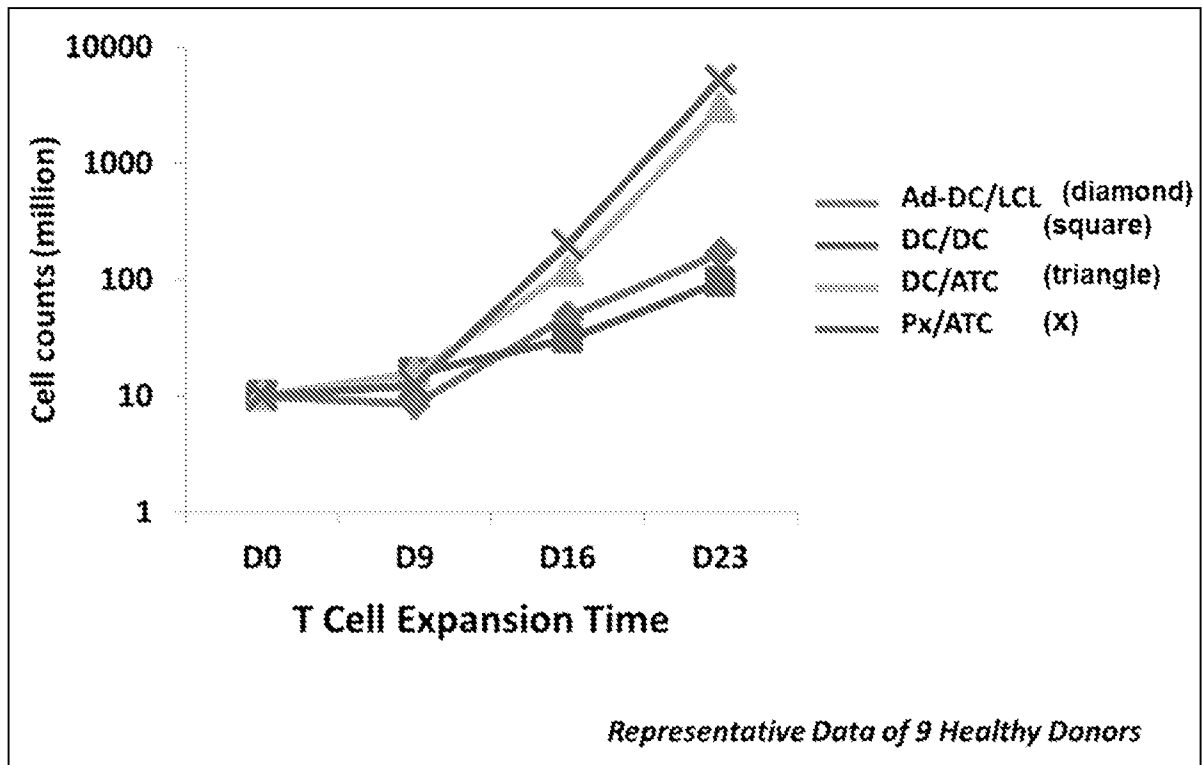
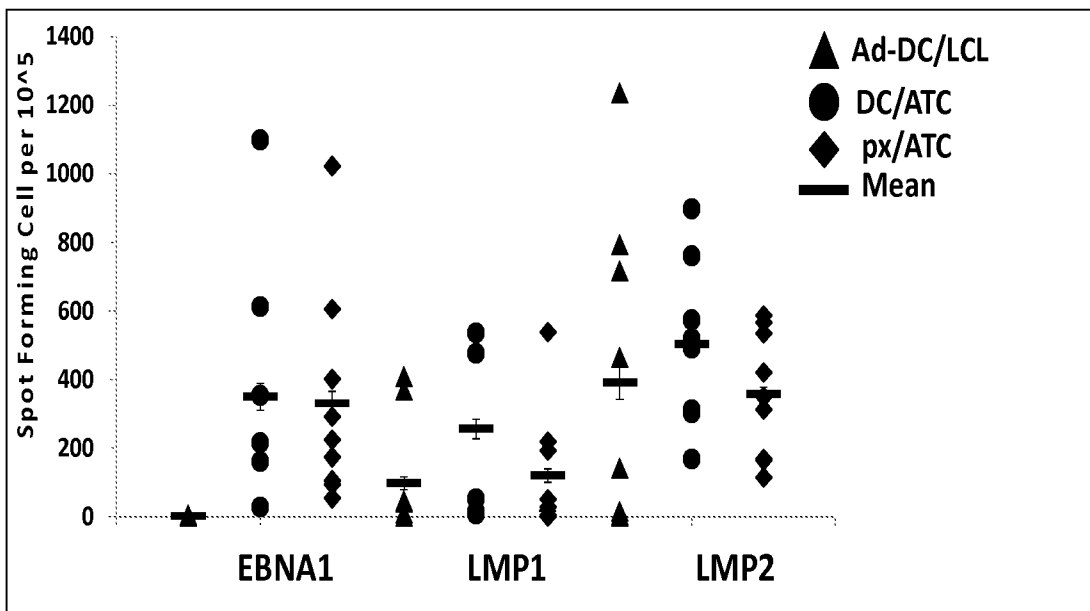
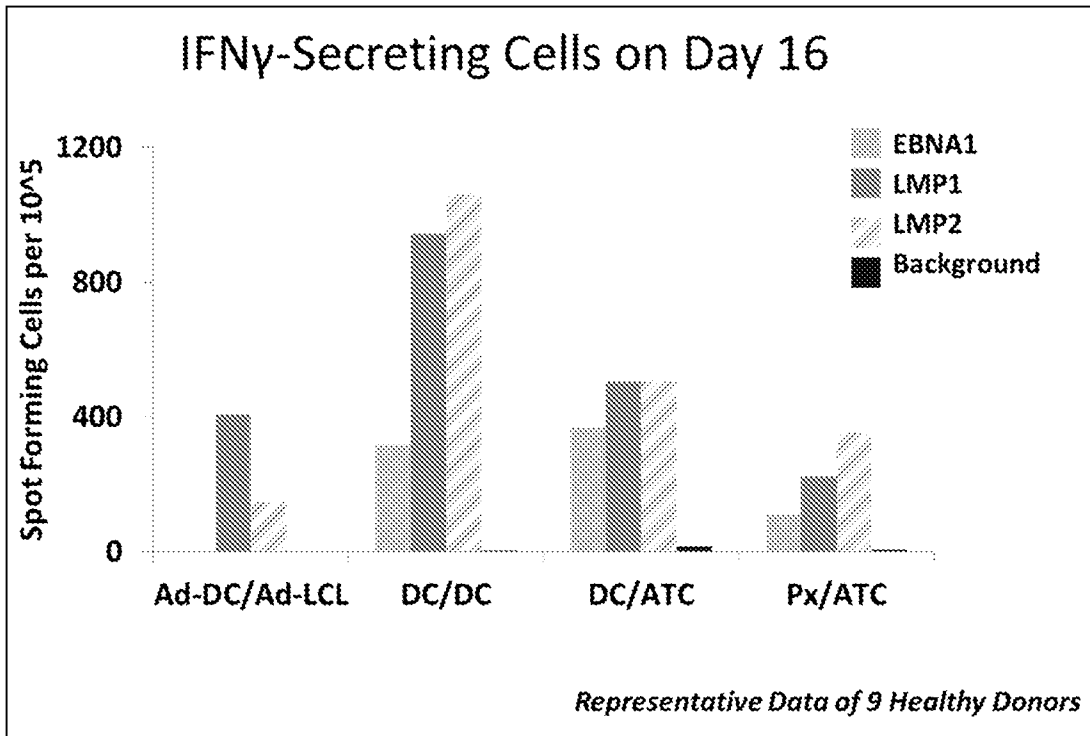


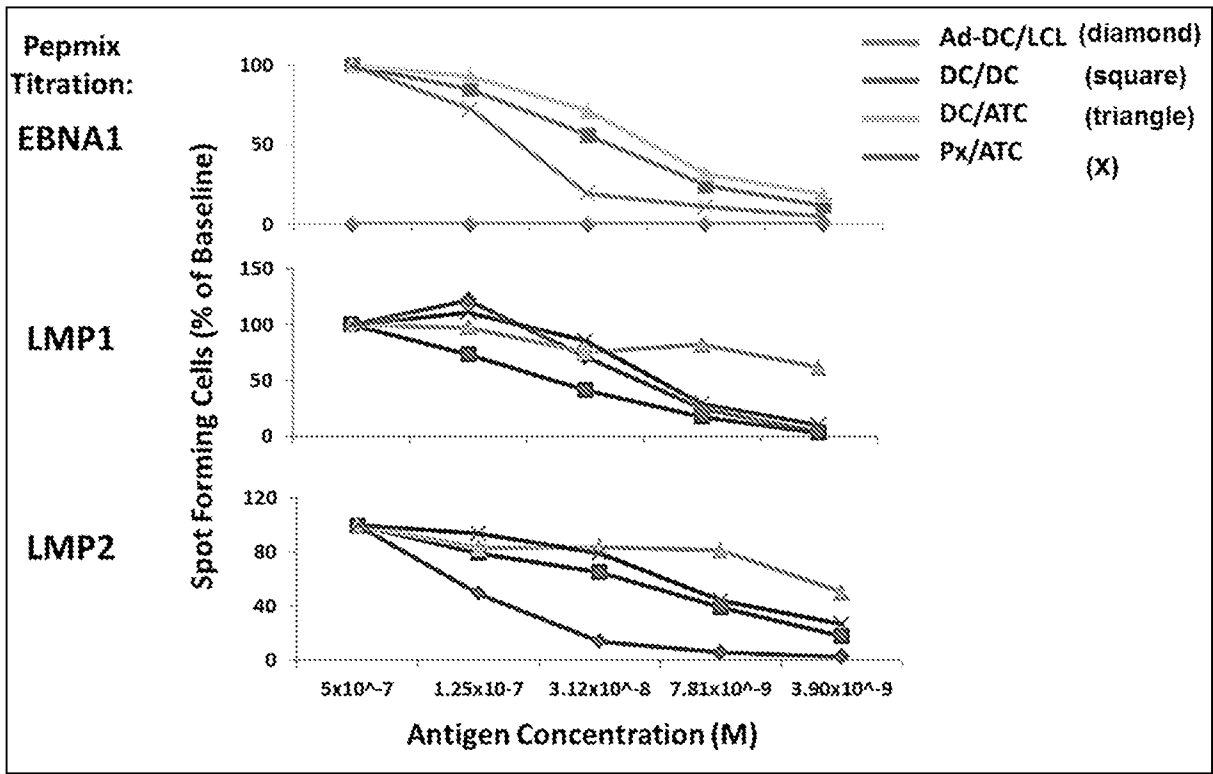
FIGURE 4B: Expansion of cells from a healthy donor in response to EBV antigens using the prior art and the new embodiments of the invention.



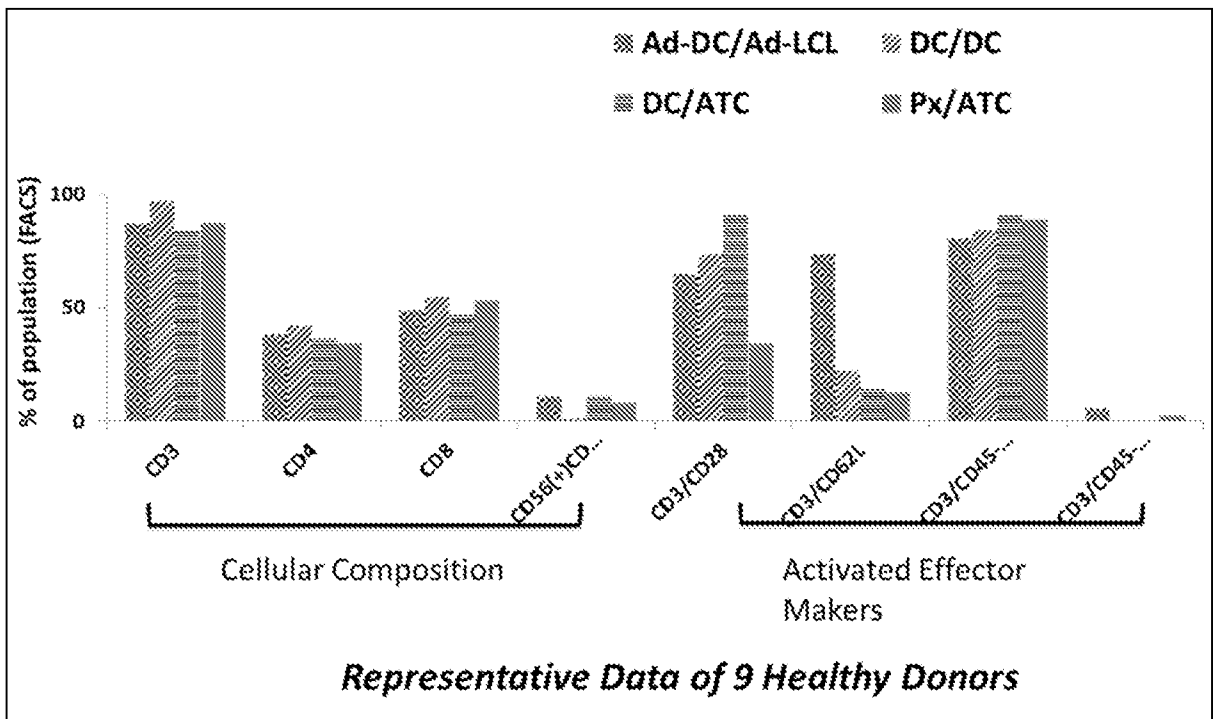
**FIGURE 4C:** The number of antigen specific, IFN $\gamma$  secreting T cells in the final cell population was increased in the embodiments of the invention compared to the prior art method



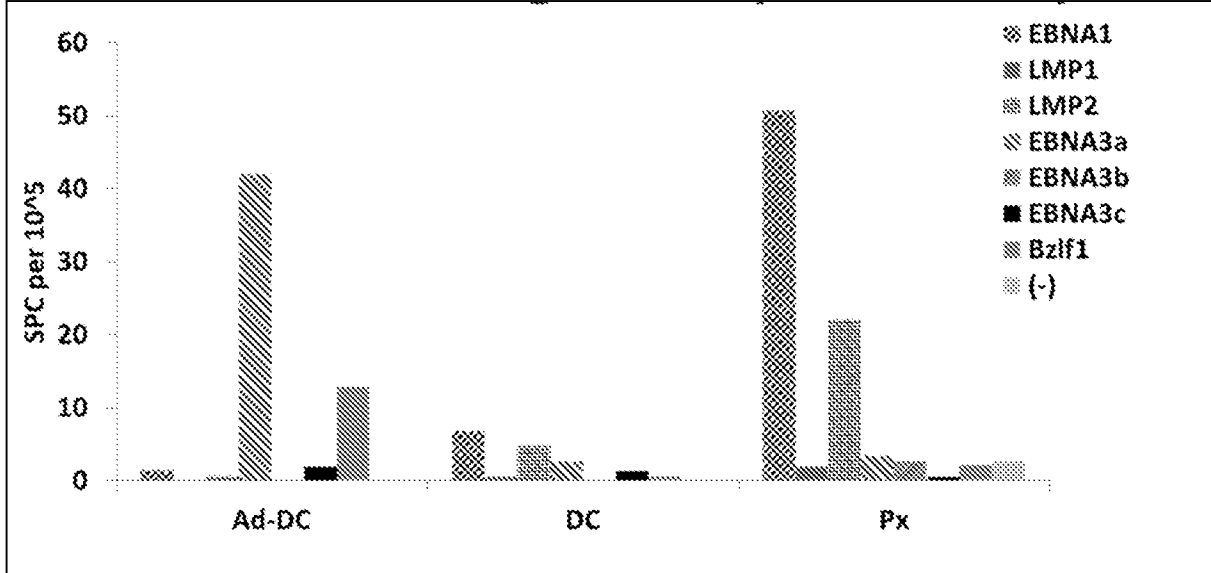
**FIGURE 4D:** The avidity of the T cell receptors for the different peptide antigens is not decreased by the use of the various embodiments when compared to the prior art process.



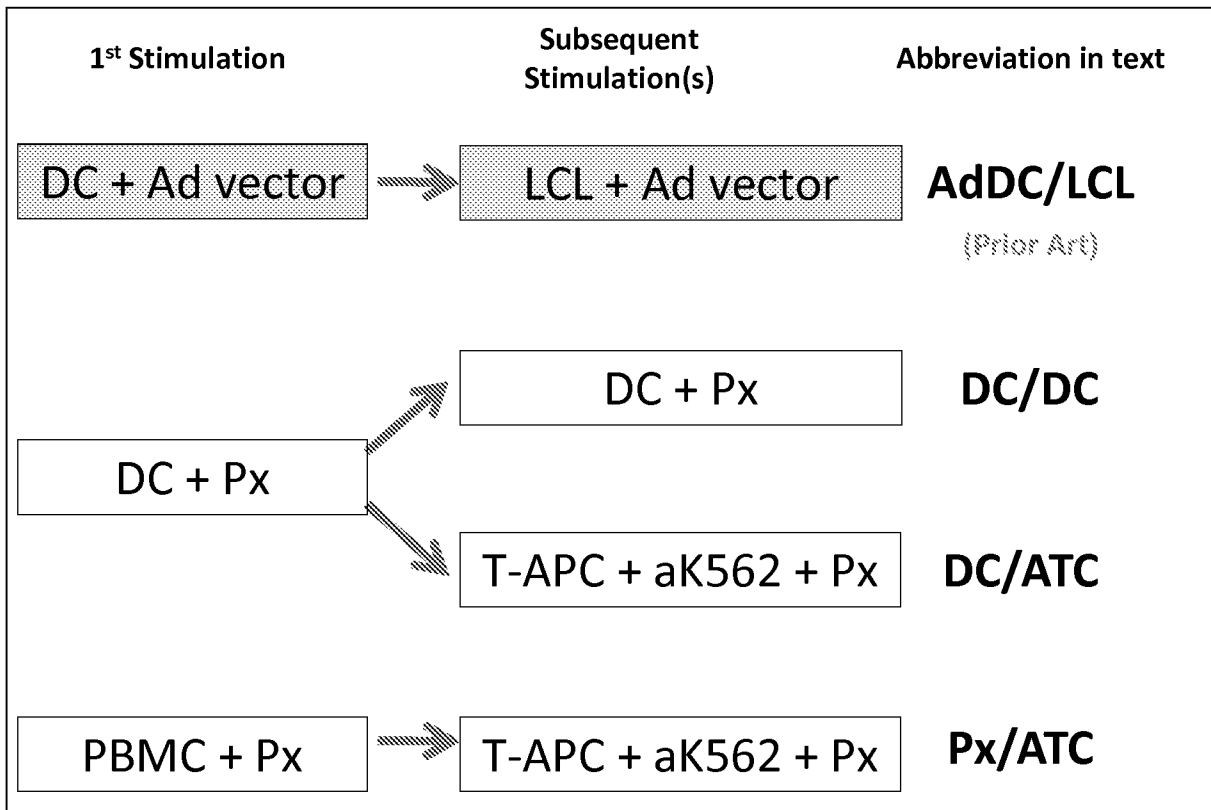
**FIGURE 4E:** T cell markers for the cellular composition and activation markers in the prior art method and the embodiments of the invention.



**FIGURE 4F:** Culture of cells using the prior art skews responses towards antigens expressed in LCL cells rather than on the latency type 2 antigens present in EBV+ cancer cells from lymphoma and NPC patients.. The new methods therefore focus the specificity of the expanded CTLs towards clinically relevant antigens.



**FIGURE 5A:** A Schematic representation of the main stimulation steps used in the prior art process and the various embodiments of the invention.



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FIGURE 5B: Expansion of EBV specific T cells is enhanced in 2 of the 3 embodiments when compared to the prior art in NPC patients.

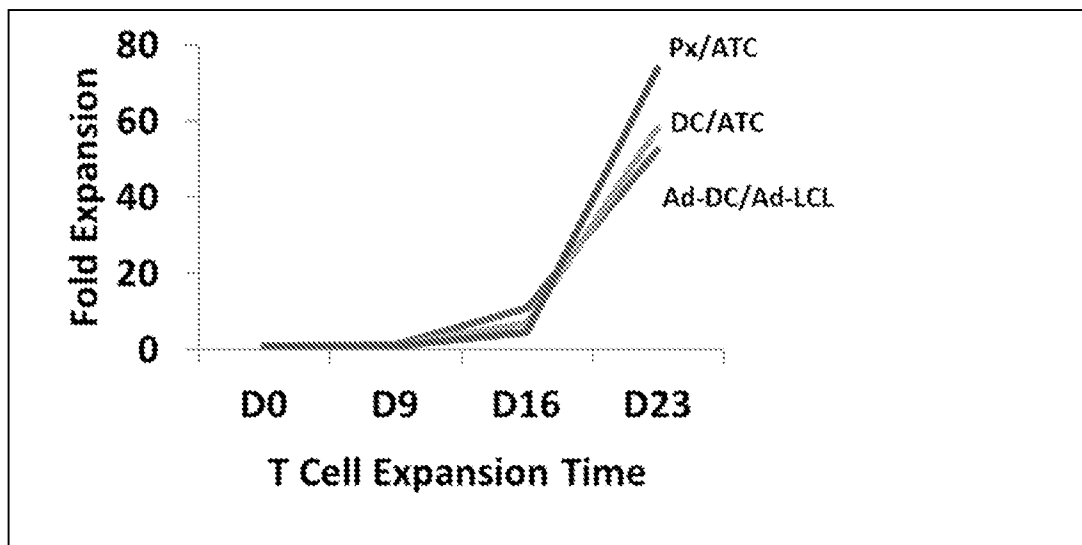
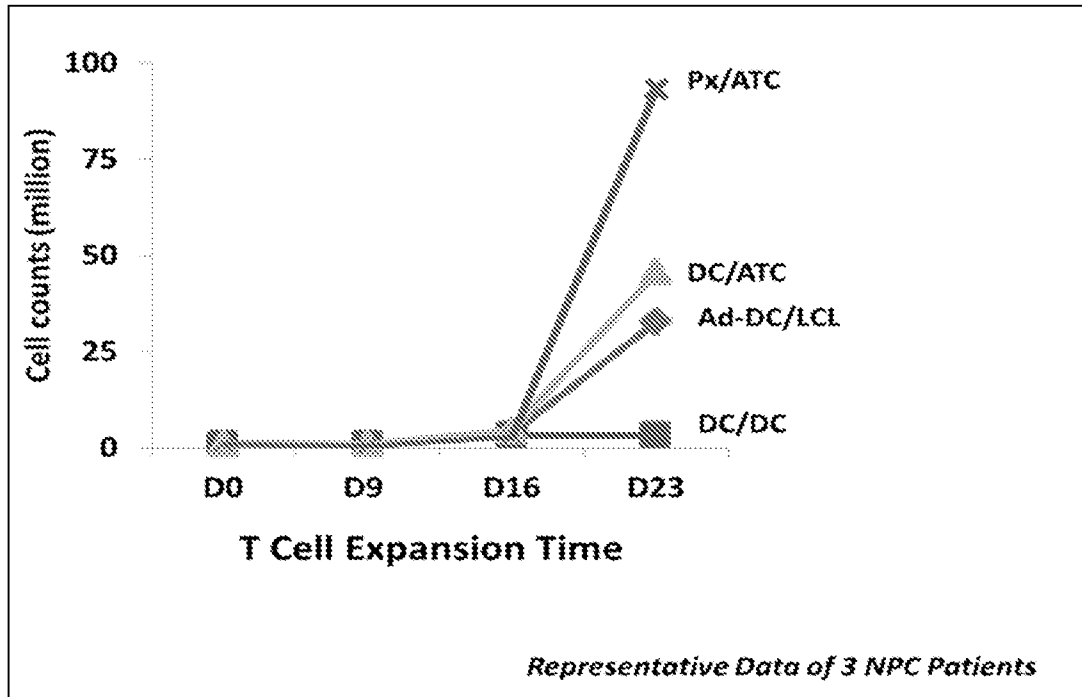




FIGURE 5C: EBV specific T cells expanded from NPC patients show enhanced specificity when using the embodiments of the invention compared to the prior art method.

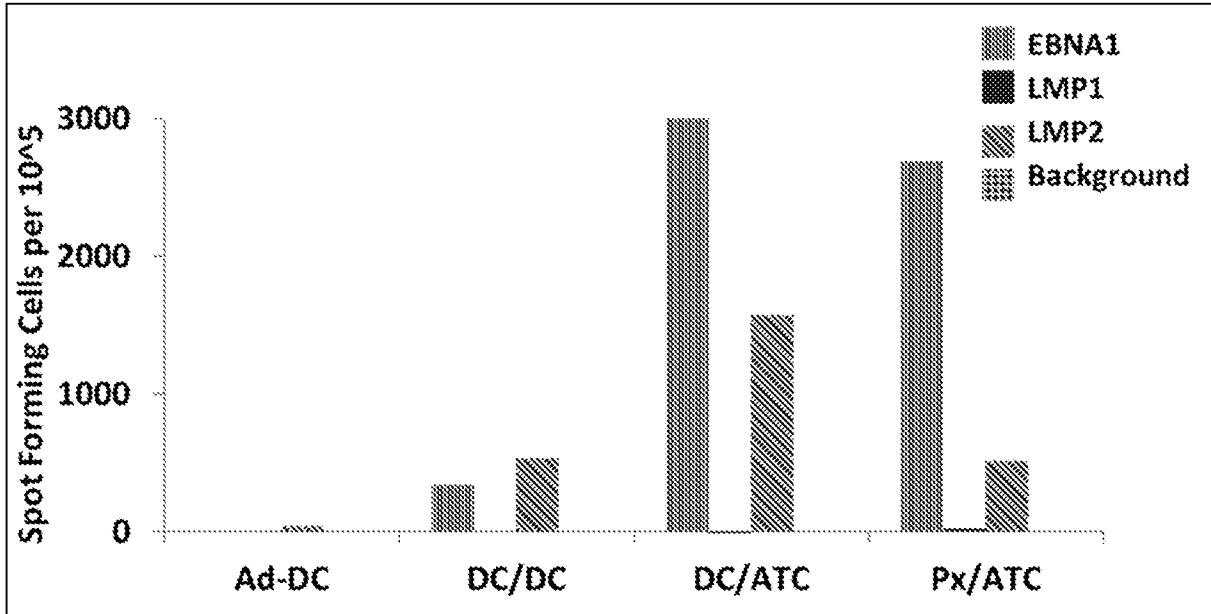


FIGURE 5D: The avidity of the T cell receptors for the different peptide antigens is not decreased by the use of the various embodiments when compared to the prior art process in NPC patients.

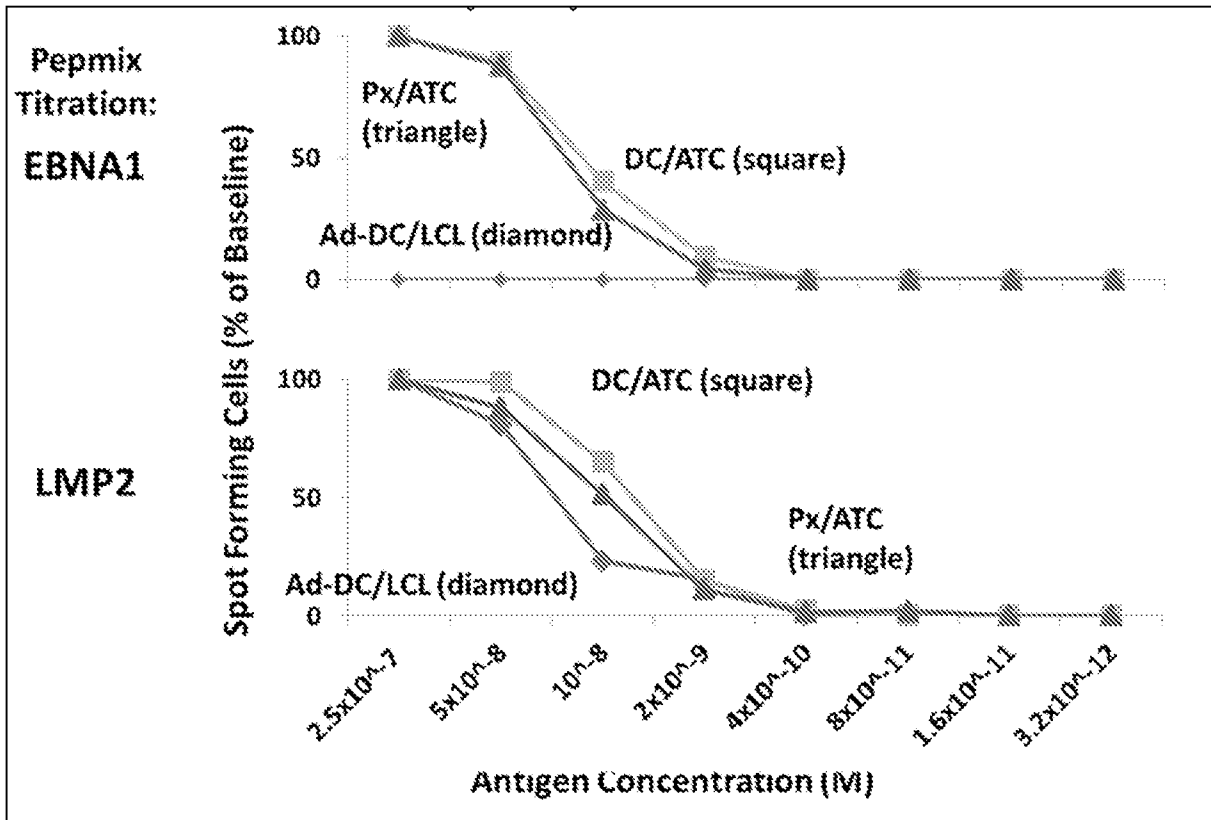


FIGURE 5E: T cell markers for the cellular composition and activation markers in the prior art method and the embodiments of the invention in NPC patients.

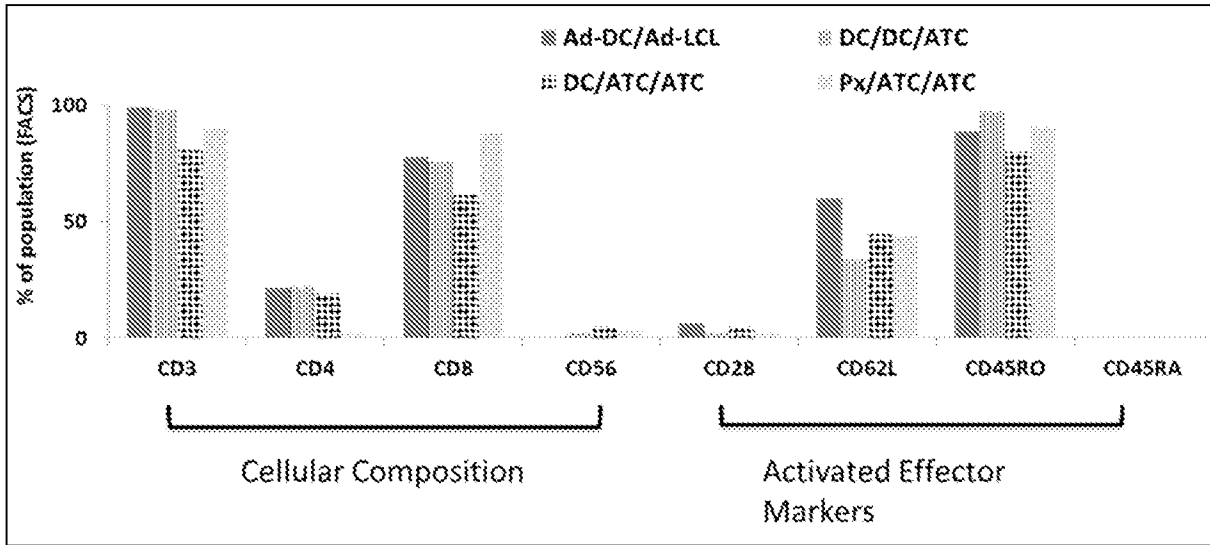


FIGURE 5F: T Cells from NPC patients expanded employing the new embodiments of the invention(CD3+/CD19-) kill LCL (EBV+ cancer cell-line, CD3-/CD19+) better in co-culture for 4 and 10 days as T cells expanded by the prior art.

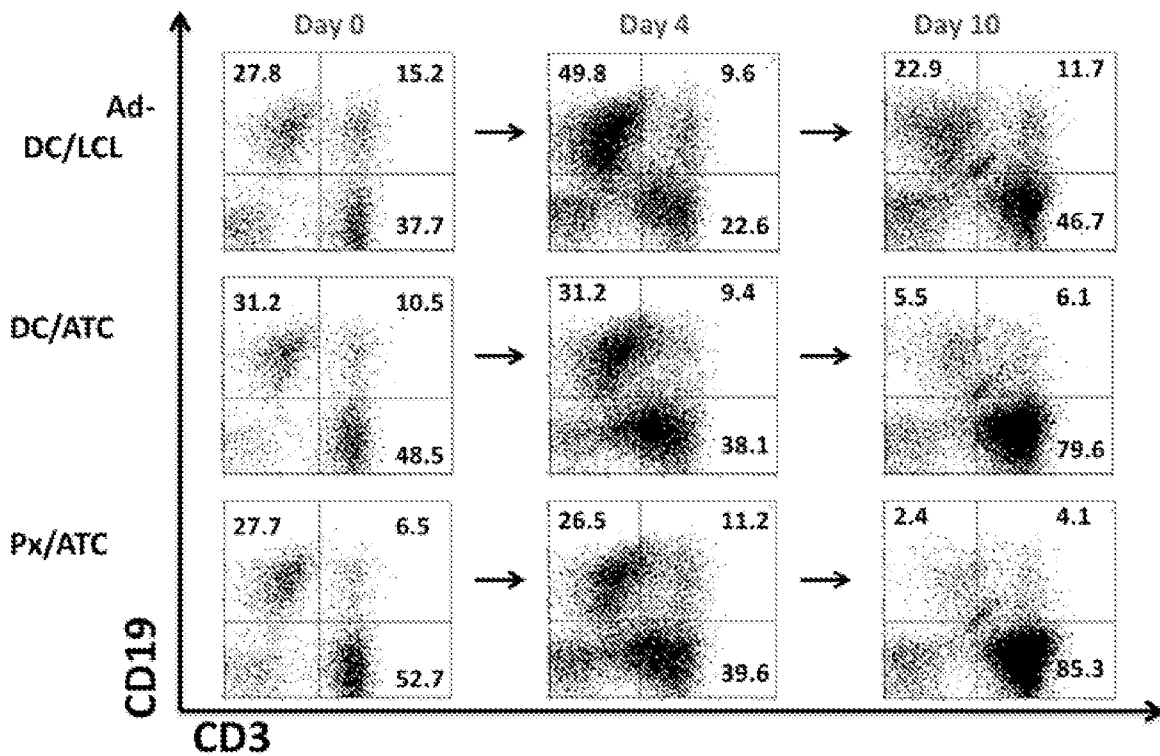


FIGURE 6A: A Schematic representation of the main stimulation steps used in the prior art process and the various embodiments of the invention.

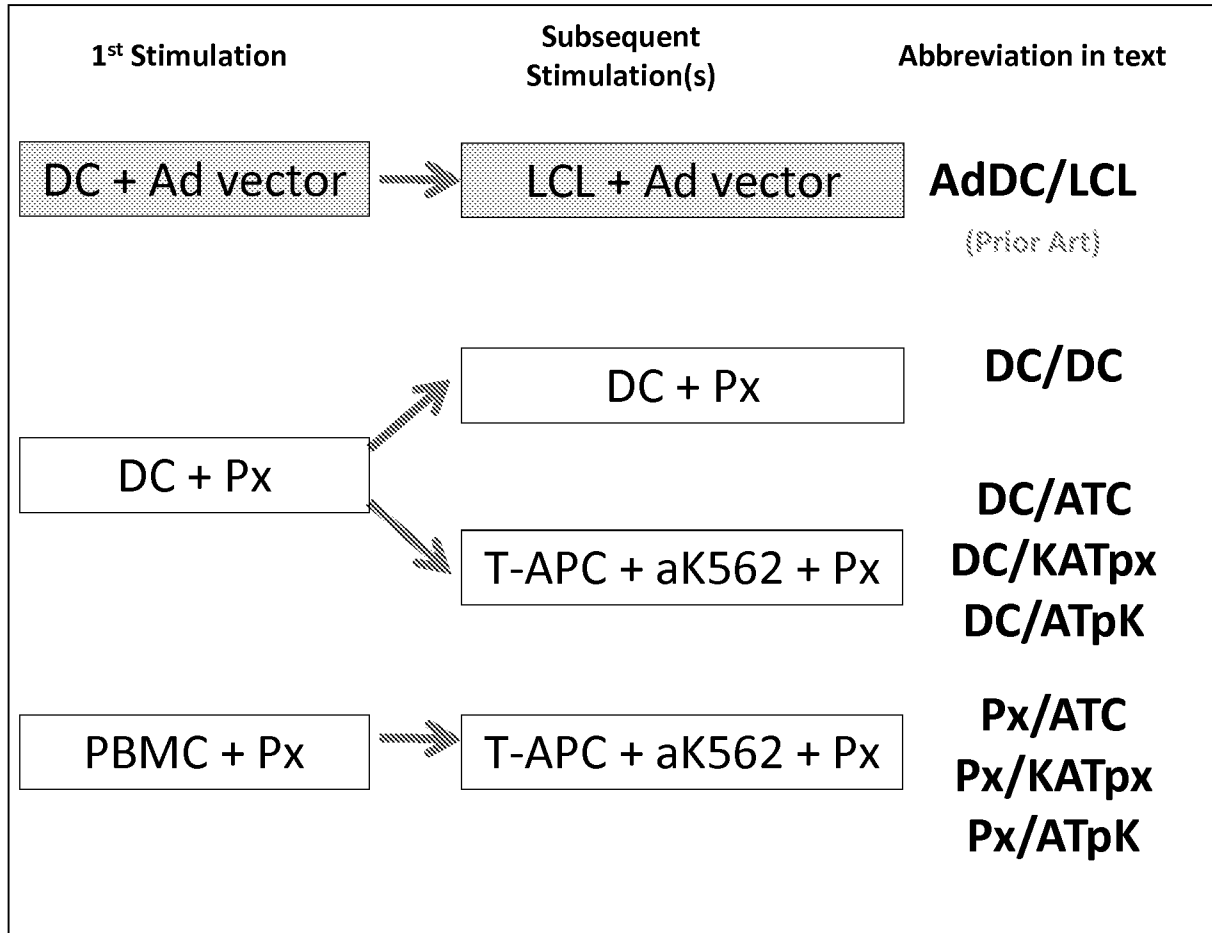


FIGURE 6B: Expansion of EBV specific T cells is enhanced in 2 of the 3 embodiments when compared to the prior art in lymphoma patients.

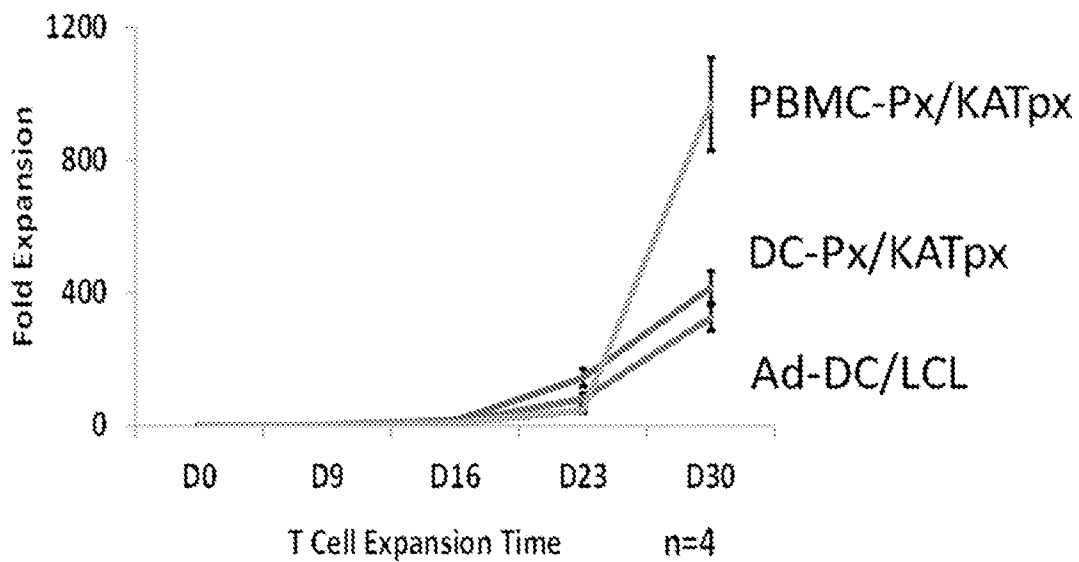
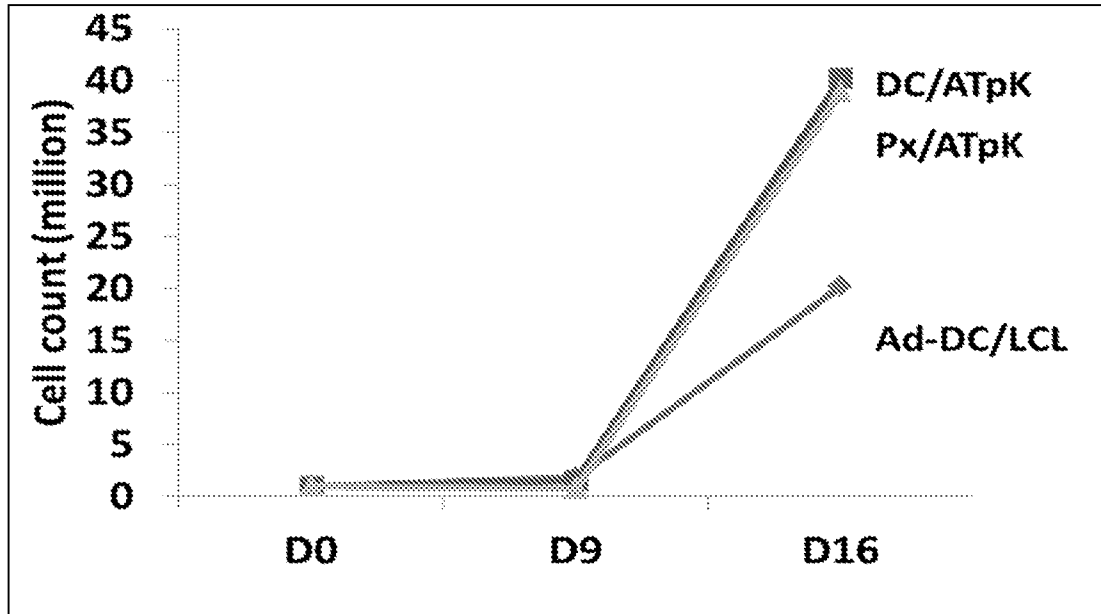


FIGURE 6C: EBV specific T cells expanded from lymphoma patients show enhanced specificity when using the embodiments of the invention compared to the prior art method.

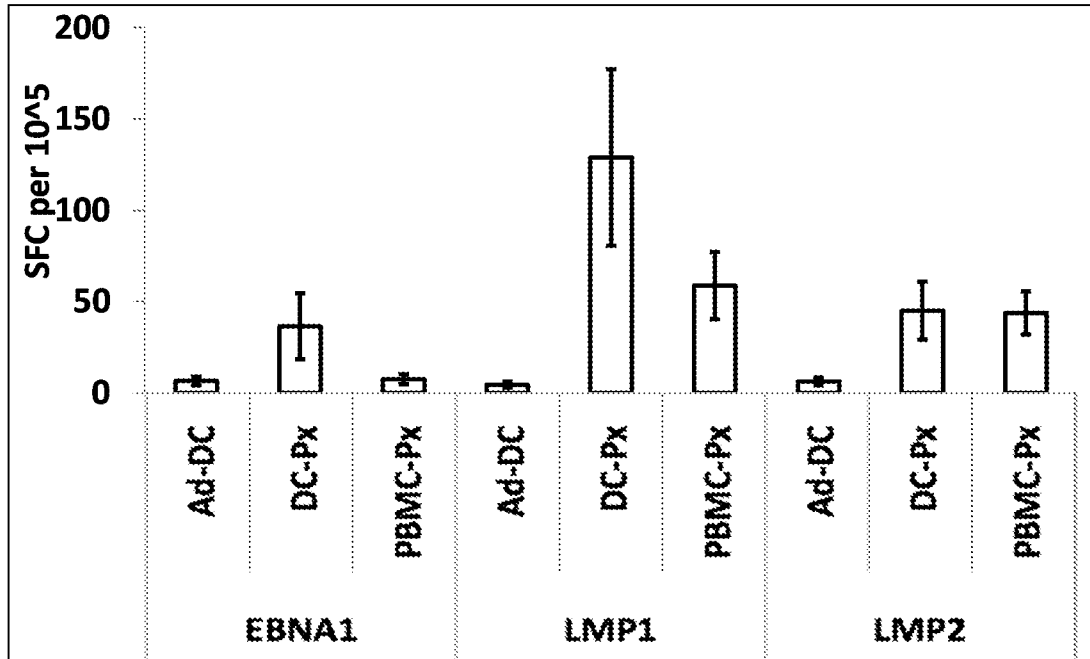
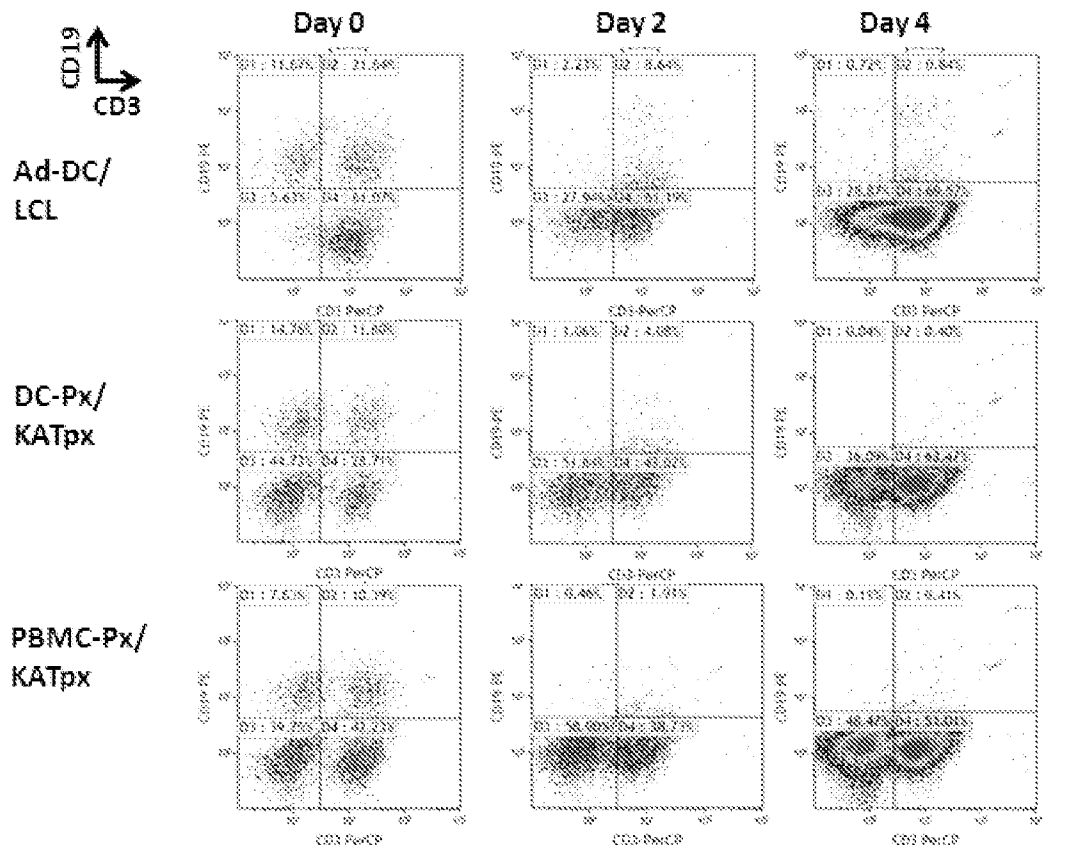
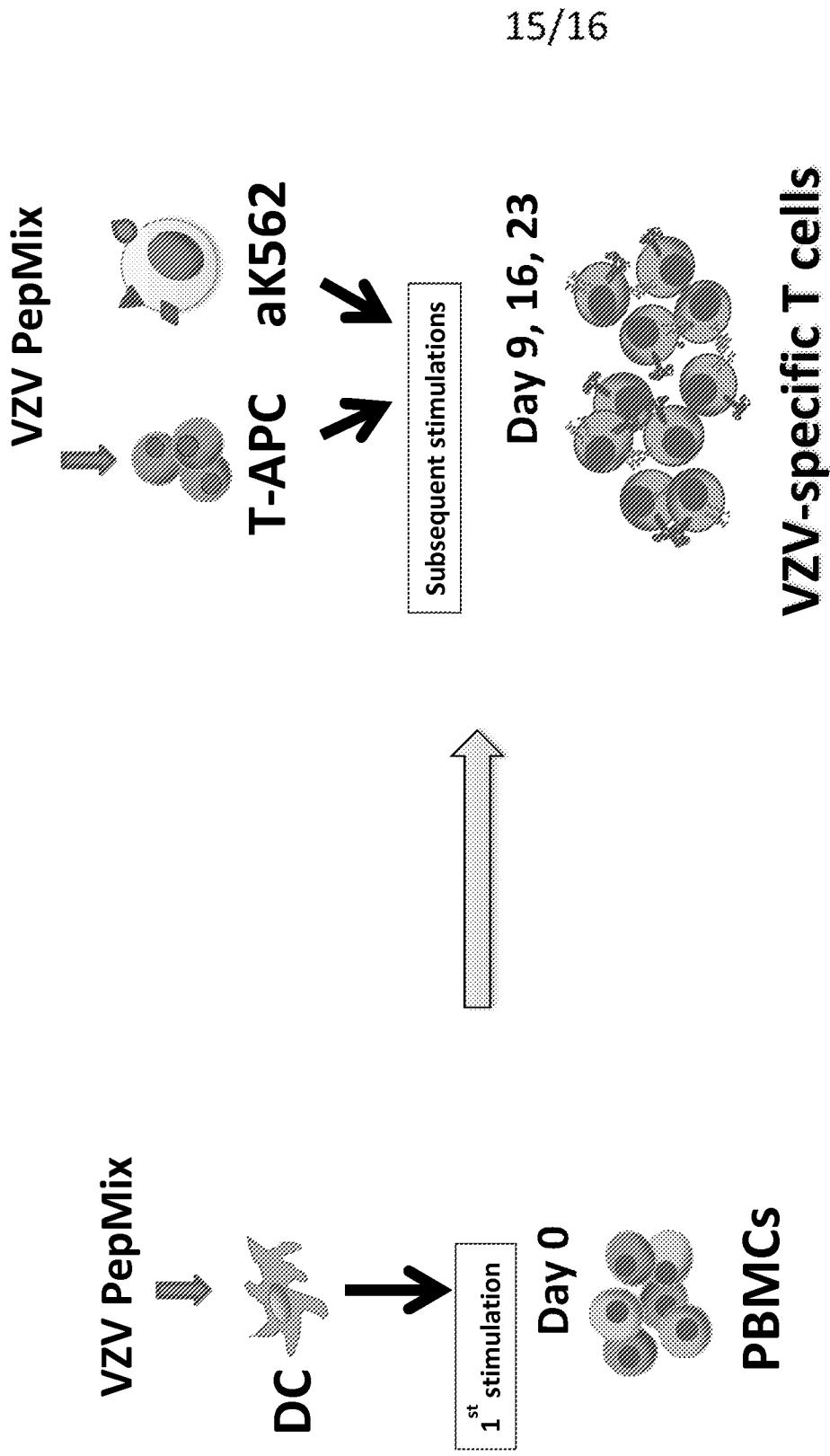


Figure 6D: T Cells from lymphoma patients expanded employing the new embodiments of the invention(CD3+/CD19-) kill LCL (EBV+ cancer cell-line, CD3-/CD19+) equally well or better in co-culture for 2 or 4 days as T cells expanded by the prior art.



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FIGURE 7A: is a schematic representation of the embodiment of the invention that was used to expand VZV specific T cells from the PBMC of a healthy donor



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FIGURE 7B: Expansion of VZV-specific T cells

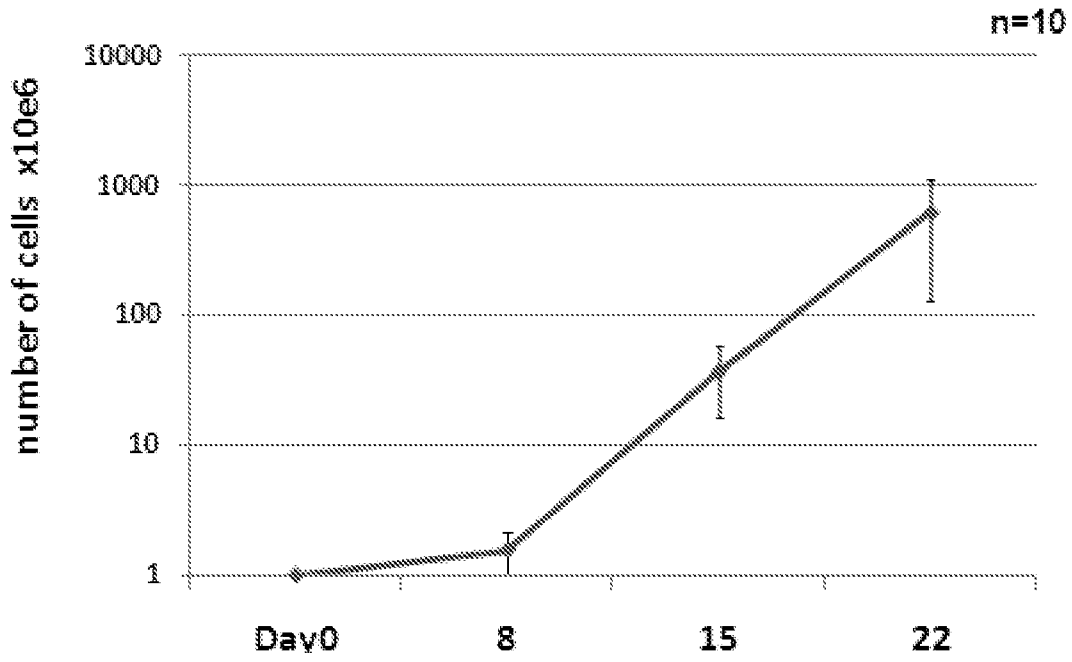
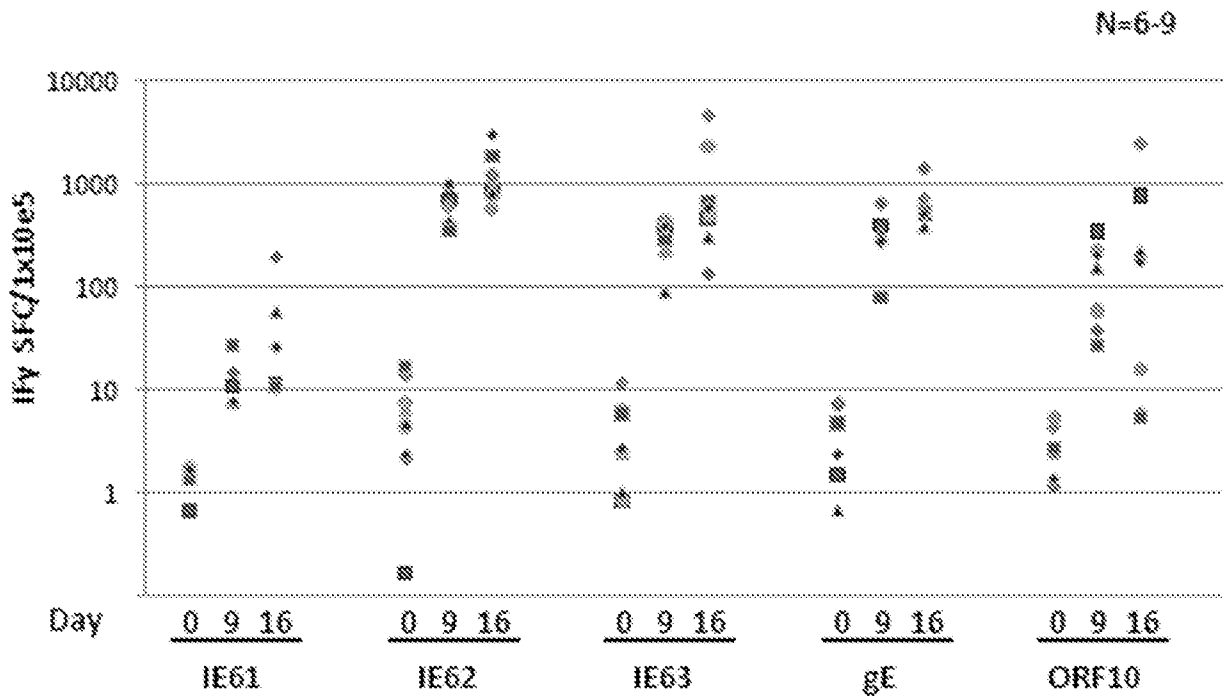


FIGURE 7C: IFN $\gamma$  secreting cells in cultures of VZV specific T cells



**INTERNATIONAL SEARCH REPORT**

International application No  
PCT/GB2012/05Q896

A. CLASSIFICATION OF SUBJECT MATTER  
INV. C12N5/Q783 A61K35/14  
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)  
C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal , BIOSIS, EMBASE, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>ANDO JUN ET AL: "Towards Phase 2/3 Trials for Epstein - Barr Virus (EBV)-Associated Malignancies" , BLOOD, vol . 118, no. 21, November 2011 (2011-11) , page 1727 , XP55034426, &amp; 53RD ANNUAL MEETING AND EXPOSITION OF THE AMEPsICAN -SOCI ETY-OF-HEMATOLOGY (ASH) ; SAN DI EGO, CA, USA; DECEMBER 10 -13, 2011 ISSN : 0006-4971</p>	<p>1-4, 6-13, 17-19 , 21-26, 28,29</p>
Y	<p>abstract ----- - / - -</p>	<p>5-16,20</p>

Further documents are listed in the continuation of Box C.

See patent family annex.

\* Special categories of cited documents :

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"E" earlier application or patent but published on or after the international filing date

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"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

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"&" document member of the same patent family

Date of the actual completion of the international search

3 August 2012

Date of mailing of the international search report

10/08/2012

Name and mailing address of the ISA/

European Patent Office, P.B. 5818 Patentlaan 2  
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Fax: (+31-70) 340-3016

Authorized officer

Manu, Domini que



## INTERNATIONAL SEARCH REPORT

International application No  
PCT/GB2012/05Q896

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>GERDEMANN ULRI KE ET AL: "Cytotoxi c T lymphocytes simul taneously targeti ng multiple tumor-associ ated anti gens to treat EBV negative lymphoma. ", MOLECULAR THERAPY : THE JOURNAL OF THE AMERICAN SOCI ETY OF GENE THERAPY DEC 2011 LNKD- doI :10.1038/MT .2011 .167 PUBMED:21915103, vol . 19, no. 12, 13 September 2011 (2011-09-13) , pages 2258-2268, XP008154173, ISSN : 1525-0024</p>	1-4, 14-16, 21-25 , 27,29
Y	the whole document	5-13, 16, 20
X	<p>-----</p> <p>GERDEMANN ULRI KE ET AL: "Generati on of multivirus-speci fic T cells to prevent/treat viral infections after allogenei c hematopoiети c stem cel l transplant. ", JOURNAL OF VISUALIZED EXPERIMENTS : JOVE 2011 LNKD- DOI :10.3791/2735 PUBMED:21654628, no. 51, May 2011 (2011-05) , XP55034564, ISSN : 1940-087X</p>	21-25
Y	the whole document	5, 14, 15, 20
T	<p>-----</p> <p>ULRI KE GERDEMANN ET AL: "Rapi dly Generated Mul tivi rus-speci fic Cytotoxi c T Lymphocytes for the Prophylaxi s and Treatment of Vi ral Infecti ons" , MOLECULAR THERAPY, vol . 20, no. 8, 17 July 2012 (2012-07-17) , XP55034394, ISSN : 1525-0015, DOI : 10.1038/mt .2012 .130 the whole document</p> <p>-----</p>	

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/ GB2 012/ 050896

### Box No. I Nucleotide and/or amino acid sequence(s) (Continuation of item 1.c of the first sheet)

1. With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, the international search was carried out on the basis of:
  - a. (means)
    - on paper
    - in electronic form
  - b. (time)
    - in the international application as filed
    - together with the international application in electronic form
    - subsequently to this Authority for the purpose of search
2.  In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
3. Additional comments: