Title: COMBINED EPITOPES, USE OF EPITOPES OR THEIR COMBINATION, COMPOSITION, USES OF THE COMPOSITION, ANTI-HIV-1 PROPHYLACTIC VACCINES, THERAPEUTIC VACCINES, METHOD FOR THE IDENTIFICATION OF EPITOPES AND METHODS FOR TREATMENT AND PREVENTION

Abstract: The present invention refers to new epitopes recognized by CD4+ T-lymphocytes. In addition, the present invention refers to the uses of such epitopes and their combinations, particularly in the treatment or prevention of disorders caused by the HIV-1 virus. The present invention also refers to a composition comprising said epitopes and the uses of said composition, particularly in the treatment or prevention of disorders caused by the HIV-1 virus. The present invention also refers to anti-HTV-1 prophylactic vaccines and therapeutic vaccines. Furthermore, the present invention refers to a method for the identification of epitopes and methods for treating or preventing an infection caused by the HIV-1 virus.
"EPITOPES, COMBINED EPITOPES, USE OF EPITOPES OR THEIR COMBINATION, COMPOSITION, USES OF THE COMPOSITION, ANTI-HIV-I PROPHYLACTIC VACCINES, THERAPEUTIC VACCINES, METHOD FOR THE IDENTIFICATION OF EPITOPES AND METHODS FOR TREATMENT AND PREVENTION"

FIELD OF THE INVENTION

The present invention refers to one or more epitopes recognized by CD4+ T-lymphocytes, as well as the use of said epitopes and their combinations, particularly in the treatment or prevention of disorders caused by the HIV-I virus.

The present invention also refers to a composition comprising said epitopes and to an use of said composition.

In addition, the present invention refers to anti-HIV-1 prophylactic and therapeutic vaccines.

Furthermore, the present invention refers to a method for the identification of epitopes and methods for treating or preventing infections caused by the HIV-I virus.

BACKGROUND OF THE INVENTION

Although the development of an effective vaccine against HIV-I is urgent, in view of the 40 million people currently infected with the virus, no effective vaccine for this purpose has been developed yet.

In 2003, at least 18 experimental prophylactic vaccines were known to be undergoing clinical trials.
(Phases 1 to 3) (McMichael AJ & Hanke T, Nature Medicine, 2003) and an even greater number of pre-clinical approaches attempting the preparation of an effective vaccine. Almost all of the products well known in the art use similar common approaches for preparing an immunogen. These methods include the use, in the vaccine, of integral sequences HIV-1 proteins, as recombinant proteins, recombinant DNA, or inserts in recombinant virus vectors.

Results of several vaccine trials that passed the pre-clinical stage were reported between 2003 and 2004. The Phase III trial (efficacy) of the REMUNE vaccine (combination of depleted whole inactive HIV-I of the envelope protein), tested on over 5,000 individuals at risk of infection, failed to demonstrate protection and the trials were ended (McCarthy M, HIV Vaccine fails vaccine trial, THE LANCET, 361:755-756, 2003). In 2004, a promising candidate vaccine developed by a team from University of Oxford and University of Kenya reached the stage of Phase I/II trials but demonstrated immunogenicity in no more than one-third of tested individuals (International Conference Aids Vaccine 2004, Lausanne, Switzerland).

One of the disadvantages of the DNA vaccines, whether recombinant or from viral vectors encoding genes or whole proteins of the HIV-I, such as those already known in the state of the art, is the principle used by them, which facilitates the development of molecular escape mechanisms by the HIV-I in response to immune and other pressures. Such a disadvantage may be one of the reasons for the failure of such vaccines. Furthermore, variation in the
sequence may lead to several immunological escape mechanisms, while the immunization with T-cell epitopes from multiple viral gene products out of the context of native HIV-I proteins can avoid recognition escape mechanisms, maintaining the induction of significative cellular immune responses (Newman MJ et al, 2002).

In the specific and direct area of the development of a product, De Groot AS et al in Engineering immunogenic consensus T helper epitopes for a cross-clade HIV vaccine (2004) and in HIV vaccine development by computer assisted design: the GAIA vaccine (2005) identified conserved epitopes of the recognized HIV-I, over a large proportion of patients, by using computer-aided algorithms. The documents mentioned above also reported the insertion of said epitopes in vaccines undergoing testing at the pre-clinical stage with experimental animals. Previous reports also showed the selection of epitopes based on their ability to bind to multiple HLA molecules, with inferior results to those of De Groot AS et al, since HIV-1+ patients did not recognize said epitopes (Van den Burg SH et al, Journal of Immunology (1999), Identification of a conserved universal Th epitope in HIV-I). Wilson CC et al (Journal of Virology 2001. Identification and antigenicity of broadly cross-reactive...) also use another search algorithm for sequences binding multiple HLA class II molecules, identifying a group of peptides frequently recognized by mononuclear blood cells of HIV-1+ patients.

Thus the development and testing of new formulations and combinations of epitopes remains a top priority in
health research worldwide.

Among the broadly recognized challenges in obtaining effective vaccines against infections by HIV-I (McMichael AJ & Hanke T, Nature Medicine (2003)), the following are cited:

a) the construction of a vaccine capable of inducing high levels of neutralizing antibodies;
b) the construction of a vaccine capable of generating cellular immune responses (from T-lymphocyte) of greater intensity;
c) initiating new Phase III trials sooner, with an eventual combination of vaccines.

Although anti-HIV-1 cytotoxic CD8+ T-lymphocytes effectively destroy the cells infected by the virus, their activity is fundamentally dependent on the presence of anti-HIV-1 CD4+ T-cells (Rosenberg et al, 1997; Kalams et al, 1998; Heeney et al, 2002). Therefore, the incorporation of appropriate HIV-I epitopes recognized by the CD4+ T-cells may be essential to the success of an anti-HIV-1 vaccine candidate. However, there are only a few epitopes known for HIV-I CD4+ T-lymphocytes, when compared to the CD8+ epitopes, which have been much more searched for.

Therefore, it is necessary to identify new epitopes of HIV-I that are recognized by the CD4+ T-cells in the majority of individuals. These epitopes may be incorporated in an candidate vaccine based on epitopes that generate immune responses in a significant proportion of the population exposed to the virus. It is expected that this strategy, together with new strategies for the formulation
of immunogens, may lead to a protective vaccine.

However, the identification of such peptides is
difficult and very expensive when using the traditional
methodology of overlapping peptides: hundreds or thousands
of peptides synthesized step by step, with the difference
of a few amino acid residues, have to be tested.
Furthermore, said approach may not identify some of the
epitopes in the regions between adjacent peptides, thus
being cumbersome and not totally effective.

Recent literature reports have identified HIV-I
epitopes by using algorithms, as mentioned above. However,
the test results have not demonstrated the expected
recognition potential of the epitopes, i.e., recognized by
the greatest majority of HIV-1+ patients.

**BRIEF DESCRIPTION OF THE SEQUENCE LISTING**

The present invention will be more clearly understood
upon reading the following non-restrictive detailed
description and the Sequence Listing.

The Sequence Listing contains the 3-letter code for
amino acids, as defined in accordance with the IUPAC-IUBMB
standards described in *Nucleic Acids Res.*, 13:3,021-3,030

The Sequence Listing presented herein has the
objective of defining the amino acid sequences of 16
epitopes specific to HIV-I, for the first time identified
as being widely recognized by CD4+ cells of patients with
HIV-I, which are an object of the present invention.
BRIEF DESCRIPTION OF THE INVENTION

The present invention refers to conserved and new, previously unknown, multiple epitopes of the HIV-I, which are widely recognized by CD4+ T-lymphocytes.

The present invention also refers to the use of said epitopes.

Other embodiments of the present invention are combinations comprising 2 or more of the new epitopes and the use of said combinations.

In addition, the present invention refers to compositions comprising at least one of the new epitopes and the use of said compositions.

Further objects of the present invention are prophylactic anti-HIV-1 vaccines and therapeutic vaccines, which contain one or more of said epitopes.

Moreover, the present invention refers to a method for the identification of new epitopes, which comprises the steps of:

a) selection of sequences conserved -from the consensus sequence of HIV-I of subtype B, having at least 15 amino acid residues.

b) selection, among the sequences selected in step a), of those that bind promiscuously multiple HLA-DR molecules;

c) expansion of the sequences of said peptides selected at the N-terminal and C-terminal ends;

d) confirmation of recognition of the selected
peptides by T-lymphocytes of HIV-1+ patients.

Another embodiment of the present invention is a method for treating or preventing infections caused by the HIV-I virus.

**DETAILED DESCRIPTION OF THE INVENTION**

The present invention refers to one or more synthetic peptides, either simple or having covalent modifications, such as miristoylation and other forms of lipopeptides, referred to herein as epitopes, conserved and isolated from the HIV-I genome, which bind in a promiscuous manner multiple HLA-DR molecules and are recognized by CD4+ T-lymphocytes in patients infected by the HIV-I.

Said epitopes are selected from the group consisting of:

- EELRSLYNTVATLYCVH (SEQ ID NO.: 1);
- SPEVPMFSALSE (SEQ ID NO.: 2);
- KRWIILGLNKIVRMYSPTSI (SEQ ID NO.: 3);
- DKELYPLASLRSLFG (SEQ ID NO.: 4);
- QRPLVTIKIGGQLKE (SEQ ID NO.: 5);
- GKIILVAHVASGYI (SEQ ID NO.: 6);
- TMLLGMIMCSAA (SEQ ID NO.: 7);
- ALFYKLDVVPID (SEQ ID NO.: 8);
- NTSYRLISCNTSVI (SEQ ID NO.: 9);
- SELYLYKVKIEPLGVAP (SEQ ID NO.: 10);
- ELLKTVRLIKFLYQSNP (SEQ ID NO.: 11);
- EAIIRILQQLLFIHF (SEQ ID NO.: 12);
- QQLLFIHFRICRHSRIG (SEQ ID NO.: 13);
- SLQYLALVALVAPKK (SEQ ID NO.: 14);
- VLEWRFDSRLAFHHV (SEQ ID NO.: 15);
- VLAIVALVVATIIAI (SEQ ID NO.: 16).

The corresponding sequence listing is mentioned at the end of each epitope.

Particularly, the epitopes of the present invention are selected from the group consisting of:

- EELRSLYNTVATLYCVH (SEQ ID NO.: 1),
- KRWIILGLNKIVRMYSPTSI (SEQ ID NO.: 3),
- QRPLVTIKIGGQLKE (SEQ ID NO.: 5),
- TMLLGMLMICSAA (SEQ ID NO.: 7),
- ALFYKLDVVID (SEQ ID NO.: 8),
- ELLKTVRFLKFLYSNPS (SEQ ID NO.: 11) e VLAIVALVVATIIAI (SEQ ID NO.: 16).

The epitopes of the present invention are particularly derived from the gag, pol, env, rev, vpr, vif, nef and vpu proteins of HIV-I.

Said epitopes may be derived from any type of HIV-I subtype but, preferably, they are derived from subtype B.

One of the advantages of the present invention is the recognition of said epitopes by CD4+ T-cells of multiple HIV-I patients.

According to the present invention, "epitopes" mean the epitopes mentioned above, their functional equivalents and mimetic sequences thereof.

A "functional equivalent" refers to structurally distinct sequences, fragments, analogues, derivatives or associations, which perform the same function to achieve equal results. It is understood that any alterations made
by those skilled in the art, which lead in an obvious manner to equivalent effects, shall also be considered as a part of the invention. More particularly, functional equivalents are the sequences presenting homology of at least 12 amino acids to the epitopes described above and perform the same function of said epitopes, exhibiting equal or similar results.

In accordance with the present invention, "mimetic sequences" are understood as being non natural amino acid sequences with modified structures, so that they present functions and results equal or similar to the sequences of the epitopes of the present invention.

The present invention also refers to the use of said epitopes.

One of the uses is in the preparation of a composition for treating disorders caused by the HIV-I virus. Preferentially, said disorders are related to the immunologic system having been impaired, such as, for example, progressive destruction of lymphocytes.

Another use is as an additive in the preparation of anti-HIV-1 vaccine compositions already known in the state of the art, to provide T cell help and increase of the immunogenicity and protective properties thereof.

The epitopes of the present invention have further use in diagnostic methods and in trials for the evaluation of the immune response of CD4+ T-lymphocytes in patients with HIV-I.

The cellular immune response to the epitopes of HIV-I shows correlation with progression of the disease.
Therefore, the evaluation of immune responses against the epitopes of the present invention, in patients HIV-1+ patients during the infection, or in response to immunological interventions, is essential for studying the effectiveness of the vaccine or other intervention. Individually, the epitopes of the present invention have been recognized by between 18% and 44% of patients and not a single HIV-I seronegative control individual. The group of 16 epitopes of the present invention was recognized by 32 of the 34 HIV-1+ patients tested (94%). Furthermore, patients with advanced disease and immunodeficiency showed reduced responses when compared to long-term nonprogressor (LTNP) HIV-1+ patients, demonstrating the efficacy for the evaluation of the HIV-I specific immune response.

Furthermore, the epitopes of the present invention are useful for the preparation of anti-HIV-1 prophylactic vaccines or therapeutic vaccines. Said vaccines may be more effective than those already known in the state of the art since the new epitopes of the present invention are recognized by the T-cells in a majority of individuals, thus covering a significant proportion of the population exposed to the virus.

Another object of the present invention is a combination comprising two or more of the new epitopes and the uses of said combinations.

The uses of the combination are in accordance with the above descriptions, namely: in the preparation of a composition for treating disorders caused by the HIV-I virus, as an additive in the preparation of anti-HIV-1
vaccine compositions already known in the state of the art, in diagnostic methods and tests for evaluating the immune response of CD4 T-lymphocytes of patients with HIV-I and in the preparation of anti-HIV-1 prophylactic vaccines or therapeutic vaccines.

It is also an object of the present invention a composition comprising at least one of the new epitopes. Said composition further comprises a pharmaceutically acceptable carrier or vehicle.

The compositions of the present invention may be in the solid or liquid form. Said compositions may be formulated for a rapid or prolonged release of their components and may further comprise compounds for stimulating and/or inhibiting the immunologic system. Said compositions may be prepared in accordance with conventional methods already known in the state of the art.

Further objects of the present invention are uses of the composition described above. This includes the use of said composition in the preparation of anti-HIV-1 prophylactic vaccines or therapeutic vaccines, as an additive in the preparation of compositions of existing experimental anti-HIV-1 prophylactic vaccines, in diagnostic methods and tests for evaluating the immune response of CD4+ T-lymphocytes of patients with HIV-I, as described above for the new epitopes.

The present invention also refers to an anti-HIV-1 prophylactic vaccine or a therapeutic vaccine. Said vaccines comprise at least one of the epitopes described above, in association with one or more pharmaceutically
acceptable adjuvants, vehicles, excipients, binding agents, carriers or preservatives.

Optionally, the vaccines of the present invention comprise a combination of epitopes or the composition of the present invention with one or more pharmaceutically acceptable adjuvants, vehicles, excipients, bonding agents, carriers or preservatives.

The vaccines in accordance with the present invention may be formulated according to the following forms, including one or more of the new epitopes:

a) combined with adjuvants or carriers as micro/nanoparticles/spheres;

b) a recombinant DNA construction with the sequences of one or more of the new epitopes, particularly containing other protein products, optionally inserted in carriers such as micro/nanoparticles/spheres;

c) a recombinant protein construction with sequences of the new epitopes, particularly combined with adjuvants, optionally inserted in carriers such as micro/nanoparticles/spheres;

d) a viral vector containing sequences of the new epitopes; and

e) a combination of the new epitopes with HIV-I immunogens already known in the state of the art.

Another object of the present invention is a method for the identification of the new epitopes, which comprises the steps of:

a) selection of the peptides conserved from the
consensus sequence of HIV-I of subtype B, with at least 15 amino acids;

b) selection, among the peptides selected in step a), of those binding promiscuously multiple HLA-DR molecules;

c) expansion of the sequences of said peptides selected at the N-terminal and C-terminal ends;

d) confirmation of recognition of the selected peptides by T-lymphocytes of HIV-1+ patients.

An algorithm is used to screen the whole HIV-I genome and to select sequences of HIV-I proteins, which may bind multiple HLA-DR molecules commonly found in the population. Preferentially, the selection of step b) is carried out by means of a TEPITOPE algorithm (Generation of tissue-specific and promiscuous HLA ligand databases using DNA chips and virtual HLA class LI matrices, Sturniolo, T et al, Nature Biotechnology 17, 555-562, 1999), which employs a matrix based on the results of real peptide binding tests for each one of the several HLA-DR molecules, and attributes scores to each sequence derived from the protein, selecting those having scores superior to the chosen threshold. In the case of the present invention, said selected peptides bind in a "promiscuous" manner to at least 18 HLA-DR molecules available on said algorithm, with a threshold of 3%.

After step b), 2 or 3 flanking residues are added to the N-terminal and C-terminal ends, in order to increase the frequency of peptide recognition.

Step d) is achieved by synthesizing the peptides
representing the epitopes of the present invention, followed by ELISPOT assay to detect IFN-γ-producing cells. For the purposes of the present invention, the positivity cutoff value used is ≥ 30 SFC IFN-γ /10^5 PBMC, calculated based on the levels of responses of HIV-I seronegative patients.

Further aims of the present invention are methods for treating or preventing an infection caused by the HIV-I virus. HIV-I infection is related to the impairment of the immunologic system of the patient.

In accordance with the present invention, the method for treating or preventing comprises administering a therapeutically effective amount of the composition described herein to the patient. Optionally, said method comprises administering a therapeutically effective amount of the combination of epitopes of the present invention or anti-HIV-1 prophylactic vaccines or therapeutic vaccines described herein to the patient.

A "therapeutically effective amount" designates an amount of epitopes, a combination thereof, a composition or vaccine effective for the "treatment" of a disease or a disorder in a patient. As defined in the present invention, this is considered to be an amount capable of avoiding contagion, reducing the number of HIV-I RNA copies/mm3 in the plasma, or increasing the number of CD4+ T-cells/mm3 in the peripheral blood of patients, or reducing the number of cells infected by the HIV-I virus and/or improving one or more of the symptoms associated to the infections caused by the HIV-I virus, as a result of inducing immune responses.
of the T-lymphocytes through the stimulation of the CD4+ T-cells by means of epitopes recognized by them.

According to the present invention, the administration to the patient can be made in any form, such as, for example, oral, intranasal, mucosal, local, transdermal and parenteral, (such as intravenous, intraperitoneal, intradermal, subcutaneous or intramuscular). Administration may also occur by means of the use of needle-less administration devices. Administration may further occur through the use of a combination of methods, such as being firstly administered parenterally with subsequent mucosal administration.

Preferably, administration is subcutaneous or intramuscular.

The following examples are illustrative and provide a clearer and more consistent understanding of the invention but are not intended to limit its scope.

EXAMPLES

Example 1

Identification of New Immunodominant Epitopes

To select the epitopes, sections of at least 15 consecutive conserved amino acids present in the consensus sequence (complete HIV-I genome) of the HIV-I subtype B (December 2002 version), available at the website: http://HIV-1-web.lanl.gov/content/index, were analyzed by using the TEPITOPE algorithm for the prediction of binding.
to HLA-DR molecules. The selected epitopes were those predicted to bind over the chosen threshold (3%) to the greatest possible number of HLA-DR molecules in a "promiscuous" manner. From the whole consensus viral genome of subtype B, epitopes of the proteins gag, env, pol, vpu, vpr, vif, vpu were selected capable of binding at least 18 of the 25 HLA-DR molecules available in the algorithm with a threshold of 3%, thus selecting epitopes with a high chance of binding HLA-DR molecules with great avidity. For each selected sequence (9-residue HLA-binding core), N- and C-terminal flanking residues were added, flanking so as to increase the percentage of patients recognizing each peptide.

Example 2

Evaluation of the Epitopes

In order to evaluate the capacity of the selected epitopes to be recognized by T-lymphocytes of HIV-1+ patients, said epitopes (described in Table 1 below) were synthesized in solid phase by using Fmoc chemistry and having a C-terminal amide.

Table 1

Sequences of the synthetic peptides encoding Potentially Promiscuous Epitopes Derived from conserved regions of clade B HIV-I Consensus Sequence Selected by the TEPITOPE
<table>
<thead>
<tr>
<th>Identification</th>
<th>Peptides</th>
<th>Sequences</th>
<th>% of binding HLA molecules using TEPITOPE algorithm (3% threshold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>gag 1</td>
<td>Gag (73-89)</td>
<td>EELRSLYNTVATLYCVH 2</td>
<td>72</td>
</tr>
<tr>
<td>gag 2</td>
<td>Gag (165-177)</td>
<td>SPEVIPMFSALSE</td>
<td>88</td>
</tr>
<tr>
<td>gag 3</td>
<td>Gag (263-282)</td>
<td>KRWIILGLNKIVRMYVTSI</td>
<td>100</td>
</tr>
<tr>
<td>gag 4</td>
<td>Gag (480-494)</td>
<td>DKELEYPLASLRSLFG</td>
<td>76</td>
</tr>
<tr>
<td>pol 1</td>
<td>Pol (63-77)</td>
<td>QRPLVTIKIGGQLKE</td>
<td>64</td>
</tr>
<tr>
<td>pol 3</td>
<td>Pol (694-708)</td>
<td>GKIILVAVHVAGYI</td>
<td>72</td>
</tr>
<tr>
<td>env 2</td>
<td>Env(19-31)</td>
<td>TMLLGMLMICSA</td>
<td>80</td>
</tr>
<tr>
<td>env 3</td>
<td>Env(178-189)</td>
<td>ALFYKLDVVID</td>
<td>72</td>
</tr>
<tr>
<td>env 4</td>
<td>Env(198-211)</td>
<td>NTSYRLISCNTSVI</td>
<td>88</td>
</tr>
<tr>
<td>env 5</td>
<td>Env(487-504)</td>
<td>SELYLYKVKEPLGVAP</td>
<td>100</td>
</tr>
<tr>
<td>rev 1</td>
<td>rev(11-27)</td>
<td>ELLKTVRLIKFLYQSNP</td>
<td>88</td>
</tr>
<tr>
<td>vpr 2</td>
<td>vpr (58-72)</td>
<td>EAIIRILQOLLFIHF</td>
<td>96</td>
</tr>
<tr>
<td>vpr 3</td>
<td>vpr (65-82)</td>
<td>QOLLFIHFRIGCRHSRIG</td>
<td>72</td>
</tr>
<tr>
<td>vif 2</td>
<td>vif (144-158)</td>
<td>SLQYLALVALVAPKK</td>
<td>72</td>
</tr>
</tbody>
</table>
Identification | Peptides | Sequences | % of binding HLA molecules using TEPITOPE algorithm (3% threshold)
--- | --- | --- | ---
nef 1 | nef (180-194) | VLEWRFDSDLFH | 52 |
vpu 1 | vpu (6-20) | VLAIVALVATTIAI | 100 |

1. Position of the amino acid region selected from the HIV-1 protein: gag (73-89) : HIV-I gag protein, sequence 73-89.
2. Amino acid sequence of each selected peptide.

Synthetic peptide stocks were diluted to 25mg/mL in dimethyl sulphoxide, followed by ELISPOT assays for detection of IFN-γ-producing cells in response to the epitopes using peripheral blood mononuclear cells (PBMC) in 34 HIV-1+ patients of several clinical groups, long-term nonprogressors (LTNP), reconstituted, partial controllers, and progressors. Each epitope was tested in a concentration of 5µM. The cryopreserved cells were maintained in culture for 16 hours, placed on plates in the presence of the epitopes, incubated for a further 18 hours, washed and ELISPOT plates were developed for the identification of spot-forming cells/Interferon-γ-producing cells (IFN-γ SFC), which were then counted in an automated counter (Zeiss KS
ELISpot/Axioplan 2). Eight seronegative individuals were tested with all epitopes and all presented responses below 20 spots/10^6 PBMC; even the average plus 3 standard deviations was below 25 in all cases in the highest case (Table 2).

Table 2 below shows the frequency of the IFN-γ-producing cells in response to each of the HIV-I peptides in 8 healthy control individuals, seronegative for HIV-I. The absence of response for any peptide, shown in all of the 8 individuals tested, demonstrates the specificity of the reaction for patients with HIV-I. Based on these results, 30 SFC/10^6 PBMC IFN-γ was established as positive definition point.

Table 3 below shows individual recognition responses for each peptide by the 34 HIV-1+ patients tested. The present invention is capable of identifying previously unknown multiple conserved epitopes of HIV-I, recognized by CD4+ T-lymphocytes.

RESULTS

The present approach was capable of identifying previously unknown conserved multiple epitopes of HIV-I, recognized by CD4+ T-lymphocytes. The results herein indicate that 100% of the predicted epitopes were recognized by at least 6 of the 34 patients, and that 94% of the HIV-1+ patients tested recognized at least one of the 16 peptides tested. When compared to techniques well known in the state of the art, the results obtained by the
present invention are superior, the epitopes of the present invention being more efficient. DeGroot et al (2004) used another algorithm to identify 10 HIV-I consensus T cell epitopes, that were able to elicit IFN-γ Elispot responses in only 10 out of 34 (29%) HIV-1-infected LTNP patients. The increased frequency of peptide recognition observed in our study could be due the intrinsic immunogenicity of our selected peptides as compared to those of DeGroot et al. (2004) or to patient variability. Our results also differ from another study by De Groot et al. (2005), who identified 20 epitopes from HIV-I consensus sequences. Fifteen out of 20 (75%) such peptides were recognized by less than 18% of their patients, while all peptides in our study were individually recognized by 18% or more of tested patients.

Furthermore, we have shown that HIV-1-infected patients recognize on average 5 peptides. More than 75% recognized more than 2 peptides approximately 40% of them responded to 5 or more epitopes, and 6 patients (19%) responded to 10 or more epitopes, corroborating their antigenicity and immunodominance. The increased proportion of peptides recognized, together with the increased proportion of patients recognizing each peptide, indicates that the potential for recognition of multiple epitopes in the context of the present invention is visibly superior to those of working in similar concepts.

It can be noted that the patients recognized an average of 5.5 epitopes (minimum of one, maximum of 16). The peptide poll was recognized by 44% of the tested individuals, while the peptides gag3, rev1 and gagl were
recognized, individually, by 31% to 41% of the patients. The combination of 3 most frequently recognized peptides (poll, gag3, rev1) was sufficient to elicit ELISPOT responses in 79% (27/34) of the tested HIV-1+ patients. It can also be noted that a combination of only 7 peptides (poll, gag3, rev1, env3, env2, gag1, vpul) elicited response from all 94% of patients displaying responses to at least one of the 16 peptides. Finally, it can be noted that the average response amplitude to the 16 peptides by each patient was of 722 IFN-γ/10^6-producing spots (minimum of 42, maximum of 4455). The data indicates that the tested peptides are frequently recognized by peripheral blood mononuclear cells in non-selected HIV-1+ patients, which indicates that they occur naturally during infection by HIV-I.

It should be noted that the sequences of the present invention are conserved in most major subtypes of HIV-I studied (A, B, C, D, F). These observations reinforce the findings that our selected epitopes - or sequences highly homologous to them - are broadly represented across several clades of HIV-I, raising the possibility that patients infected with other non-clade B HIV-I may also recognize epitopes originally identified in HIV-I clade B consensus. Such sequences demonstrate the same immunodominant properties relating to other subtypes of HIV-I and possess vaccine potential. The results obtained with the peptides of the present invention show that these peptides can be recognized by individuals with HIV-I that do not carry the
tested sequence, thus corroborating the concept of cross-reactivity and sustaining the possibility of crossclade immunization to afford protection against multiple subtypes of HIV-I (Table 4).
Table 2. Frequency of T cells producing IFN-gamma in response to selected HIV-I peptides among peripheral blood mononuclear cells from HIV-I seronegative individuals

<table>
<thead>
<tr>
<th>Peptides</th>
<th>IFN-T spots / 10^6 PBMC*</th>
<th>Average (AV)</th>
<th>Standard deviation (SD)</th>
<th>AV + 3SD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C#01  C#02  C#03  C#04  C#05  C#06  C#07  C#08</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gag (73-89)</td>
<td>0    7    0    2    0    0    0    5</td>
<td>1.7</td>
<td>2.8</td>
<td>10.0</td>
</tr>
<tr>
<td>Gag (161-177)</td>
<td>0    2    0    0    0    0    0    5</td>
<td>0.9</td>
<td>1.8</td>
<td>6.3</td>
</tr>
<tr>
<td>Gag (263-282)</td>
<td>0    12   5    12   0    0    0    0</td>
<td>3.6</td>
<td>5.4</td>
<td>20</td>
</tr>
<tr>
<td>Gag (480494)</td>
<td>0    0    5    12   8    0    0    5</td>
<td>3.7</td>
<td>4.6</td>
<td>17.4</td>
</tr>
<tr>
<td>Pol (63-77)</td>
<td>0    0    20   12   3    0    1    5</td>
<td>5.1</td>
<td>7.3</td>
<td>26.9</td>
</tr>
<tr>
<td>Pol (694-708)</td>
<td>0    2    0    0    0    0    0    0</td>
<td>0.2</td>
<td>0.7</td>
<td>2.4</td>
</tr>
<tr>
<td>Env(19-31)</td>
<td>0    7    5    0    0    0    0    15</td>
<td>3.4</td>
<td>5.4</td>
<td>19.7</td>
</tr>
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<td>Env(178-189)</td>
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SFC/10^6 PBMC were calculated according to the following equation: (average spots in duplicate peptide wells — average spots in duplicate control wells)*5. Positive responses (>30 SFC/10^6 PBMC).
Table 3. Recognition of the selected HIV-I clade B conserved consensus epitopes, by peripheral blood mononuclear cells in patients chronically infected by HIV-I in several phases of clinical progression, using the IFN-γ ELISPOT assay.

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<th>Peptides</th>
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<th>Progressors</th>
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<td>555</td>
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<td>Gag (165-177)</td>
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<td>35</td>
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<td>70</td>
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<td>vpu (6-20)</td>
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<td>nef (180-194)</td>
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Number of recognized epitopes: 5, 8, 12, 2, 16, 3, 2, 4, 2, 3, 6, 2, 4, 1, 1, 3, 1, 0, 1, 5, 0, 13.

SFC/10⁶ PBMC were calculated according to the following equation: (average spots in duplicate peptide wells - average spots in duplicate control wells)*5. Positive responses (>30 SFC/10⁶ PBMC) are highlighted in gray background.
Table 4: Percentage of isolates of each HIV-I subtype bearing sequences identical to the immunodominant epitopes of the HIV-I clade B consensus.

Consensus sequences which are present in 20% or more of the isolates in clade B HIV-I are underlined. Shaded areas represent non-B clade sequences with significant representation of sequences identical to immunodominant consensus B subtype epitopes. Numbers in parentheses represent the percentage of isolates with single amino acid divergences in relation to immunodominant consensus B subtype epitopes.

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<th>Identification</th>
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<th>C</th>
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<th>F</th>
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</table>
CLAIMS


2. EPITOPE, according to claim 1, wherein the epitope is selected from the group consisting of: SEQ ID NO.: 1, SEQ ID NO.: 3, SEQ ID NO.: 5, SEQ ID NO.: 7, SEQ ID NO.: 8, SEQ ID NO.: 11 and SEQ ID NO.: 16.

3. EPITOPE, according to claim 1 or 2, wherein the epitope is derived from the proteins gag, pol, env, rev, vpr, vif, nef and vpu.

4. EPITOPE, which is a functional equivalent or a mimetic sequence of one of the epitopes as defined in claim 1.

5. Use of the epitopes as defined in any one of claims 1 to 4, in the preparation of a composition for treating disorders caused by the HIV-I virus.

6. Use of epitopes as defined in any one of claims 1 to 4, as an additive in the preparation of anti-HIV-1 vaccine compositions.
7. Use of epitopes as defined in any one of claims 1 to 4, in a diagnostic method.

8. Use of epitopes as defined in any one of claims 1 to 4, in tests for the evaluation of the immune response of CD4+ T-lymphocytes in patients with HIV-I.

9. Use of epitopes as defined in one of the claims 1 to 4, in the preparation of anti-HIV-1 prophylactic vaccines or therapeutic vaccines.

10. Combination of epitopes comprising two or more of the epitopes as defined in any one of claims 1 to 4.

11. Use of the combination of epitopes as defined in claim 10, in the preparation of a composition for treating disorders caused by the HIV-I virus.

12. Use of the combination of epitopes as defined in claim 10, as an additive in the preparation of anti-HIV-1 vaccine compositions.

13. Use of the combination of epitopes as defined in claim 10, in a diagnostic method.

14. Use of the combination of epitopes as defined in claim 10, in tests for the evaluation of the immune response of CD4+ T-lymphocytes in patients with HIV-I.
15. Use of the combination of epitopes as defined in claim 10, in the preparation of anti-HIV-1 prophylactic vaccines or therapeutic vaccines.

16. Composition comprising at least one of the epitopes as defined in any one of claims 1 to 4 and a pharmaceutically acceptable carrier or vehicle.

17. Use of the composition as defined in claim 16, in the preparation of anti-HIV-1 prophylactic vaccine or therapeutic vaccine.

18. Use of the composition as defined in claim 16, as an additive in the preparation of anti-HIV-1 vaccine compositions.

19. Use of the composition as defined in claim 16, in a diagnostic method.

20. Use of the composition as defined in claim 16, in tests for the evaluation of the immune response of CD4+ T-lymphocytes in patients with HIV-I.

21. Anti-HIV-1 prophylactic vaccine comprising at least one of the epitopes as defined in any one of claims 1 to 4 and one or more of pharmaceutically acceptable adjuvants, excipients, binding agents, carriers or preservatives.

22. Anti-HIV-1 prophylactic vaccine comprising the
combination of the epitopes as defined in claim 10 and one or more of pharmaceutically acceptable adjuvants, excipients, binding agents, carriers or preservatives.

23. Anti-HIV-1 prophylactic vaccine comprising the composition defined in claim 16 and one or more of pharmaceutically acceptable adjuvants, excipients, binding agents, carriers or preservatives.

24. Anti-HIV-1 prophylactic vaccine comprising a recombinant DNA construction having sequences of one or more of the epitopes as defined in any one of claims 1 to 4.

25. Anti-HIV-1 prophylactic vaccine comprising a recombinant protein construction having sequences of one or more epitopes as defined in any one of claims 1 to 4.

26. Anti-HIV-1 prophylactic vaccine comprising a viral vector containing sequences of one or more of the epitopes as defined in any one of claims 1 to 4.

27. Anti-HIV-1 prophylactic vaccine comprising one or more of the epitopes as defined in any one of claims 1 to 4 with another CD4+ HIV-I immunogen.

28. Therapeutic vaccine comprising at least one of the epitopes as defined in any one of the claims 1 to 4 and one or more of pharmaceutically acceptable adjuvants,
excipients, binding agents, carriers or preservatives.

29. Therapeutic vaccine comprising the combination of the epitopes defined in claim 10 and one or more of pharmaceutically acceptable adjuvants, excipients, binding agents, carriers or preservatives.

30. Therapeutic vaccine comprising the composition defined in claim 16 and one or more of pharmaceutically acceptable adjuvants, excipients, binding agents, carriers or preservatives.

31. Therapeutic vaccine comprising a recombinant DNA construction having sequences of one or more of the epitopes as defined in any one of claims 1 to 4.

32. Therapeutic vaccine comprising a recombinant protein construction having sequences of one or more of the epitopes as defined in any one of claims 1 to 4.

33. Therapeutic vaccine comprising a viral vector containing sequences of one or more of the epitopes as defined in any one of claims 1 to 4.

34. Therapeutic vaccine comprising one or more of the epitopes as defined in any one of claims 1 to 4 with another CD4+ HIV-I immunogen.

35. Method for the identification of the epitopes as
defined in any one of claims 1 to 4, the method comprising the steps of:

a) selection of the peptides conserved from the consensus sequence of HIV-I of subtype B, having at least 15 amino-acids;

b) selection, among the peptides selected in stage a), of those binding promiscuously to multiple HLA-DR molecules;

c) expansion of the sequences of the said peptides selected at the N-terminal and C-terminal ends;

d) confirmation of recognition of the selected peptides by T-lymphocytes of HIV-1+ patients.

36. Method according to claim 35, wherein the selection in step b) is achieved by means of the TEPITOPE algorithm, wherein said peptides bind at least 18 HLA-DR molecules available on said algorithm, with a threshold of 3%.

37. Method according to claim 35, wherein step d) is achieved by means of ELISPOT assays with the positive definition point at 30 SFC/10^6 PBMC IFN-\( \gamma \).

38. Method for treating or preventing an infection caused by the HIV-I virus, the method comprising administering to the patient a therapeutically effective amount of the composition as defined in claim 16.

39. Method for treating or preventing an infection caused by the HIV-I virus, the method comprising administering to
the patient a therapeutically effective amount of the combination as defined in claim 10.

40. Method for treating or preventing an infection caused by the HIV-I virus, the method comprising administering to the patient a therapeutically effective amount of the vaccine as defined in any one of claims 21 to 34.

41. Method according to any one of claims 38 to 40, wherein the administration is subcutaneous or intramuscular.
SEQUENCE LISTING

<110> Universidade de Sao Paulo
Fundagao Zerbini
Fundacao de Amparo a Pesquisa do Estado de Sao Paulo - FAPESP

<120> EPITOPES, COMBINED EPITOPES, USE OF THE EPITOPES OR THEIR
COMBINATION, COMPOSITION, USE OF THE COMPOSITION, ANTI-HIV-I
PROPHYLACTIC VACCINES, THERAPEUTIC VACCINES, METHOD FOR THE
IDENTIFICATION OF EPITOPES AND METHODS FOR TREATMENT AND
PREVENTION

<130> P1036

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<151> 09.05.2005

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Ser He
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HIV-1-derived peptide

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1 5 10 15

10

Asp Lys Glu Leu Tyr Pro Leu Ala Ser Leu Arg Ser Leu Phe Gly

1 5 10 15

20

Gln Arg Pro Leu Val Thr leu Lys lie Glu Gly Gln Leu Lys Glu

1 5 10 15

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Gly Lys lie lie Leu Val Ala Val His Val Ala Ser Gly Tyr lie

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Ala Leu Phe Tyr Leu Asp Val Val Pro lie Asp

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**PRT**

*Artificial Sequence*

*HIV-1-derived peptide*
HIV-1-derived peptide

Ser Leu Gin Tyr Leu Ala Leu Val Ala Leu Val Ala Pro Lys Lys

Artificial Sequence

HIV-1-derived peptide

VaI Leu Glu Trp Arg Phe Asp Ser Arg Leu Ala Phe His His VaI

Artificial Sequence

HIV-1-derived peptide

VaI Leu Ala lie VaI Ala Leu Val Val Ala Thr lie He Ala He
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
IPC*: C07K 7/08 (2007.01); C07K 14/16 (2007.01); A61K 38/12 (2007.01)
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)
IPC*: C07K, A61K38/00

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)
Registry, CA, Genbank, EBI Patent, WPI, Epdoc, Medline, Embase

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Further documents are listed in the continuation of Box C.

See patent family annex.

Date of the actual completion of the international search 19 December 2006 (19.12.2006)

Date of mailing of the international search report 10 January 2007 (10.01.2007)

Name and mailing address of the ISA/ A T
Austrian Patent Office
Dresdner Straße 87, A-1200 Vienna
Facsimile No. +43 / 1 / 534 24 / 535

Authorized officer
GÖRNER W.

Telephone No. +43 / 1 / 534 24 / 558
Continuation of first sheet

Continuation No. I:

Nucleotide and/or amino acid sequence(s)

(Continuation of item 1.b 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. type of material: a sequence listing
   b. format of material: a sequence listing
   c. time of filing/furnishing: contained in the international application as filed

Continuation No. II;

Observations where certain claims were found unsearchable

(Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

Claims Nos.: 4,24,25,31,32 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

Claim 4 is unclear since neither mimetic nor equivalent sequences are disclosed in the application and therefore had to be excluded from the search as well as the establishment for novelty, inventive step and industrial applicability.
Claims 24,25,31,32 are unclear due to the formulations "comprising a recombinant DNA construction having sequences of one or more epitopes" and "comprising a recombinant protein construction having sequences of one or more epitopes" since neither the term "DNA construction" nor the term "protein construction" can be reliably interpreted nor the respective sequences for any DNA constructs are disclosed in the application. Therefore, the subject matters of claims 24,25,31,32 had to be excluded from the search as well as the establishment for novelty, inventive step and industrial applicability.
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