Title: MUCOSAL CYTOTOXIC T LYMPHOCYTE RESPONSES

Abstract

The invention provides methods for induction of an antigen-specific, mucosal cytotoxic T lymphocyte response useful in preventing and treating infections with pathogens that gain entry via a mucosal surface.
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MUCOSAL CYTOTOXIC T LYMPHOCYTE RESPONSES

Related Applications

The present application is a continuation-in-part application of, and claims the benefit under Title 35 of U.S. Provisional Application Nos. 60/058,523 filed on September 11, 1997, and 60/074,894 filed on February 17, 1998.

Technical Field

The present invention relates to methods and compositions for stimulating immune responses in mammals. More particularly, the invention relates to methods and compositions for stimulating mucosal immunity.

Background of the Invention


Numerous questions remain, however, concerning which vaccine candidates may afford the most effective protection against mucosal challenge with virus, and what
mechanisms may be involved in mediating protective immunity. While a number of studies have shown a role for CTL in protection against infections such as influenza that have a mucosal component (Taylor and Askonas, *Immunology* 58:417-420, 1986; Epstein et al., *J. Immunol.* 160:322-327, 1998; Kulkarni et al., *J. Virol.* 69:1261-1264, 1995), these reports have not established whether the CTL need to be in a local mucosal site to protect. Conversely, while other studies have shown the induction of CTL in the mucosa, they have not established that these cells have a role in protection (Gallichan and Rosenthal, *J. Exp. Med.* 184:1879-1890, 1996; Bennink et al., *Immunology* 35:503-509, 1978; Lohman et al., *J. Immunol.* 155:5855-5860, 1995); and Klavinskis, et al., *J. Immunol.* 157:2521-2527, 1996 *J. Immunol.* 155:5855-5860, 1995. Yet other studies have shown the induction by vaccines of protective immunity in the mucosa, but in the face of multiple immune responses, have not been able to sort out which responses are involved in protection (Lehner et al., *Nature Medicine* 2:767-775, 1996; Putkonen et al., *J. Virol.* 71:4981-4984, 1997; Miller et al., *J. Virol.* 71:1911-1921, 1997; Quesada-Rolander et al., *AIDS Res Hwn Retroviruses* 12:993-999, 1996; Bender et al., *J. Virol.* 70:6418-6424, 1996; Wang et al., *Vaccine* 15:821-825, 1997).

Thus, although the role of CTL in protection against mucosal infections has been of interest for decades, especially in the case of influenza virus, prior investigations have failed to identify fundamental mechanisms linking immune responses to protection. In this regard, because mucosal infection by virus induces a local IgA response, it has been too readily assumed that this response, and not a concomitant CTL response, was responsible for protection against viral infection through the mucosal route. However, the role of secretory IgA in neutralizing and protecting against mucosal HIV challenge is also not clear.

CTL are crucial mediators of immunity to intracellular microorganisms such as viruses as well as certain bacteria and protozoan parasites. CTL specifically recognize "non-self" antigenic peptides bound to major
histocompatibility complex (MHC) class I molecules on the surface of "target cells" and then kill the target cells expressing the non-self antigenic peptides. Non-self polypeptides from which the non-self peptides are derived can be a) proteins encoded by intracellular microbes, b) host-encoded proteins whose expression is induced by a microbe, or c) mutant host encoded proteins expressed by, for example, tumor cells.

Thus, generation of CTL responses in the inductive and the effector mucosal immune system may be important to establishing effective protective immunity to intracellular microbial pathogens that establish infection via the mucosal barriers. In some cases, administration of antigens via parenteral routes (subcutaneous, intramuscular, intravenous or intraperitoneal, for example) either fails to induce mucosal immunity or does so extremely inefficiently.

As noted above, previous reports of mucosal immune responses elicited by mucosal challenge with viruses have disclosed that the latter induces antiviral antibody responses, and in some cases CTL responses, in the intraepithelial lymphoid populations. Chen et al., J. Virol. 71:3431-3436, 1997; Sydora, et al., Cell Immunol. 167:161-169, 1996. However, it is not clear if either of such responses is relevant to protection against viral infection in general, or HIV infection in particular. Additional studies have suggested a role for CTL in protection against infections that involve the mucosa, such as influenza or respiratory syncytial virus, Taylor et al., Immunology 58:417-420, 1986; Epstein et al., J. Immunol. 160:322-327, 1998; Kulkarni et al., J. Virol. 69:1261-1264, 1995. However, these studies have not addressed the question of whether CTL must be present at the mucosal site of infection, or if their principal activity occurs systemically.

Accordingly, a need exists in the art to better define the roles and mechanisms of CTL in mediating immunity and to develop new tools for mediating immune protection against HIV and other pathogens, particularly by conferring
immune protection at mucosal sites where such pathogens initially proliferate.

Summary of the Invention

The present invention is directed to methods and compositions for inducing a protective mucosal CTL response in a subject. The methods of the invention involve administering either a soluble antigen itself, or a polynucleotide encoding the soluble antigen, to a mucosal surface. The soluble antigens can be full length, naturally occurring polypeptides or fragments (i.e., peptides) derived from them. Peptides to be administered can be any length less than that of the naturally occurring polypeptide. They can be, for example, five to one hundred amino acid residues long, preferably twenty to seventy five amino acid residues long, more preferably twenty five to sixty amino acid residues long and most preferably thirty to fifty amino acid residues long.

The soluble antigen is administered with an adjuvant at the mucosal site or without an adjuvant. Adjuvants can be, for example, cholera toxin (CT), mutant CT (MCT), E. coli heat labile enterotoxin (LT) or mutant LT (MLT). IL-12 and/or IFNγ can be administered with the soluble antigen either in the presence or absence of an adjuvant. Alternatively, the two cytokines (IL-12 and/or IFNγ) can be administered systemically and separately from the soluble antigen which is administered mucosally, optionally with adjuvant. Mucosal routes of administration include IR, intranasal (IN), intragastric (IG), intravaginal (IVG) or intratracheal (IT).

Soluble antigens can be derived from pathogenic viruses (e.g., HIV-1, influenza virus or hepatitis A virus), bacteria (e.g., Listeria monocytogenes), protozoans (e.g., Giardia lamblia). Alternatively, the soluble antigen can be a tumor-associated antigen, e.g., prostate specific antigen produced by prostate tumor cells or tyrosinase produced by melanoma cells. Peptide antigens can be cluster peptide vaccine constructs (CLUVAC). For example, an HIV-1 CLUVAC can include one or more of the following sequences:

EQMHEDISSLWDQSLKPCVKRIQRGPGRAFVTIGK (SEQ ID NO:1),
KQIINMWQEYKAMYAPPISEQGIRRIQRGPGRAVFVTIGK (SEQ ID NO:2),
RDNWRSELYKVKVIEPLGVAPTRIQRGPGRAVFVTIGK (SEQ ID NO:3),
AVAEGTDVFIEVQGAYRAIRHPRRIRQGLERRRIQRGPGRAVFVTIGK (SEQ ID NO:4),
DRVIEVQGAYRAIRHPRRIRQGLERRRIQRGPGRAVFVTIGK (SEQ ID NO:5),
DRVIEVQGAYRAIRRIQRGPGRAVFVTIGK (SEQ ID NO:6),
AQGAYRAIRHPRRIRQGLERRRIQRGPGRAVFVTIGK (SEQ ID NO:7),
EQMHEDIISSLWQLPCVKRIHIHGPGRAFYTTKN (SEQ ID NO:8),
KQIINMWQEYKAMYAPPISEQGIRRIIHGPGRAFYTTKN (SEQ ID NO:9),
RDNWRSELYKVKVIEPLGVAPTRIHIHGPGRAFYTTKN (SEQ ID NO:10),
AVAEGTDVFIEVQGAYRAIRHPRRIRQGLERRRIHIHGPGRAFYTTKN (SEQ ID NO:11),
DRVIEVQGAYRAIRHPRRIRQGLERRRIHIHGPGRAFYTTKN (SEQ ID NO:12),
DRVIEVQGAYRAIRRIHIHGPGRAFYTTKN (SEQ ID NO:13) or
AQGAYRAIRHPRRIRRIHIHGPGRAFYTTKN (SEQ ID NO:14).

Preferably, the CLUVAC includes the amino acid sequence of SEQ ID NO:2, SEQ ID NO:9 or SEQ ID NO:12.
Antigenic peptides can be longer than the length specified by the SEQ ID NO:recipe herein, i.e., the peptide can be extended by adding one or more (e.g., 5, 10, 15, 20) amino acids at the amino and/or carboxy termini of the peptide with any given SEQ ID NO.

The invention also encompasses methods for inducing a protective mucosal CTL response in a subject in which the soluble antigen is delivered IR. Preferably, the level of CTL activity induced by IR immunization is at least 10% greater than that induced by other routes of mucosal administration (e.g., IN). More preferably, mucosal CTL activity induced by IR immunization is at least 2-fold, more preferably, at least 5-fold, and most preferably, at least 10-fold greater than that induced by other routes of mucosal immunization (e.g., IN or IG).

Subjects to which the methods of the invention are applied are mammals, e.g., humans, non-human primates, cats or mice.

Also provided within the invention are immunogenic compositions for inducing a protective mucosal CTL response in a subject which are adapted for intrarectal administration. The compositions comprise a purified soluble antigen formulated for intrarectal delivery to the rectum, colon,
sigmoid colon, or distal colon. They may be formulated as a rectal enema, foam, suppository, or topical gel and generally comprise a base, carrier, or absorption-promoting agent adapted for intrarectal delivery.

In more detailed aspects, the immunogenic compositions of the invention may include a rectal emulsion or gel preparation, preferably wherein the soluble antigen is admixed with a homogenous emulsion or gel carrier, eg., a polyoxyethylene gel. Alternatively, the soluble antigen may be admixed with a rectally-compatible foam.

In other preferred aspects, the immunogenic compositions of the invention are formulated in a suppository. The suppository is comprised of a base or carrier specifically adapted for intrarectal delivery of the antigen. Preferred bases may be selected from a polyethyleneglycol, witepsol H15, witepsol W35, witepsol E85, propyleneglycol dicaprylate (Sefsol 228), Miglyol810, hydroxypropylcellulose-H (HPC), or carbopol-934P (CP). More preferably, the suppository comprises at least two base materials to optimize structural and delivery performance. In other aspects, the suppository includes a stabilizing agent to minimize intrarectal degradation of the soluble antigen.

To optimize intrarectal delivery, the immunogenic compositions of the invention also preferably include an absorption-promoting agent, for example a surfactant, mixed micelle, enamine, nitric oxide donor, sodium salicylate, glycerol ester of acetoacetic acid, cyclodextrin or beta-cyclodextrin derivative, or medium-chain fatty acid.

In yet additional aspects of the invention, immunogenic compositions are provided which include an adjuvant which enhances the CTL response. Suitable adjuvants are detoxified bacterial toxins, for example detoxified cholera toxin (CT), mutant cholera toxin (MCT), mutant- E. coli heat labile enterotoxin, and pertussis toxin.

Preferably, the adjuvant is conjugated to a mucosal tissue or T cell binding agent, such as protein A, an antibody that binds a mucosal tissue- or T-cell-specific protein, or a ligand or peptide that binds a mucosal tissue- or T-cell-
specific protein. In more preferred aspects, the adjuvant is
a recombinant cholera toxin (CT) having a B chain of CT
substituted by protein A conjugated to a CT A chain, which
exhibits reduced toxicity and enhances mucosal tissue binding
mediated by protein A. Alternatively the adjuvant may be
conjugated to a protein or peptide that binds specifically to
T cells, for example by binding CD4 or CD8 (eg., the HIV V3
loop or a T cell-binding peptide fragment of the HIV V3 loop).

Unless otherwise defined, all technical and
scientific terms used herein have the same meaning as commonly
understood by one of ordinary skill in the art to which this
invention belongs. Although methods and materials similar to
those described herein can be used in the practice or testing
of the present invention, suitable methods and materials are
described below. In case of conflict, the present
application, including definitions, will control. In
addition, the materials, methods, and examples described
herein are illustrative only and not intended to be limiting.

Other features and advantages of the invention,
e.g., prevention of viral or other infectious diseases, will
be apparent from the following detailed description, from the
drawings and from the claims.

Brief Description of the Drawings

Fig. 1 is a series of line graphs showing the
results of a CTL assay. IR HIV-1 peptide immunization induced
long lasting mucosal and systemic CTL activity.

Fig. 2 is a series of line graphs showing the
results of a CTL assay. CT enhanced (but was not essential)
for induction of CTL by IR administration of antigenic
peptide.

Fig. 3 is a series of line graphs showing the
results of a CTL assay. CTL induced by IR immunization with
an HIV-1 gp160 peptide specifically lysed target cells
transfected with and expressing an HIV-1 gp160 gene.

Fig. 4 is a series of line graphs showing the
results of a CTL assay which indicated that IR induction of
CTL was IL-12 dependent.
Fig. 5 is a bar graph showing that IR immunization protects against IR challenge with HIV-1 gp160 expressing recombinant vaccinia virus.

Fig. 6A is a line graph showing the results of a CTL assay using SP cells obtained from animals six months after IR immunization with an antigenic peptide.

Fig. 6B is a line graph showing the results of a CTL assay using SP cells obtained from animals six months after intranasal (IN) immunization with antigenic peptide.

Fig. 7A is a bar graph showing the results of a CTL assay using PP as effector cells.

Fig. 7B is a bar graph showing the results of a CTL assay using SP cells as effector cells. PP and SP cells were obtained from animals thirty five days after mucosal (IR, IN, IG) or systemic (subcutaneous (SC)) immunization with an antigenic peptide.

Fig. 8A is a bar graph showing the results of a CTL assay using effector cells from wild-type BALB/c mice.

Fig. 8B is a bar graph showing the results of a CTL assay using effector cells from BALB/c mice which lack the ability to produce functional IFNγ. These experiments show the dependence of IR induction of CTL on IFNγ.

Fig. 9A is a line graph showing the results of a CTL assay using PP as effector cells.

Fig. 9B is a line graph showing the results of a CTL assay using SP cells as effector cells. PP and SP cells were obtained from BALB/c mice thirty five days after IR immunization with either antigenic peptide, CT and IL-12 (composition A) or antigenic peptide and CT (without IL-12; composition B).

Figs. 10A and 10B demonstrate that IR immunization with PCLUS3-18IIIB or PCLUS3-18MN induces both Peyer's patch (panel A) and spleen (panel B) long-lasting immunity. However the levels of the induction of CTL are different. Killing of P18IIIB-I10 (closed squares) or P18MN-T10 peptide-pulsed targets (closed circles) is compared with killing of unpulsed targets (open squares or circles). In both panel A and panel B, SEM of triplicate were all <5% of the mean.
Fig. 11 demonstrates that protection induced by mucosal immunization with HIV-1 peptide vaccine is specific. On day 35, mice were challenged intrarectally with \(2.5 \times 10^7\) plaque-forming units (pfu) of vaccinia virus expressing gp 160IIIB (vPE16) or with \(2.5 \times 10^7\) pfu of vaccinia virus expressing \(\beta\)-galactosidase (vSC8). Bars = SEM of five mice per group. The difference is significant at \(P < 0.01\) by Student’s test.

Fig. 12 demonstrates that protection induced by mucosal immunization with HIV-1 peptide vaccine is long-lasting. On day 35 or 6 months after the start of the immunization, mice were challenged intrarectally with \(2.5 \times 10^7\) pfu of vaccinia virus expressing gp 160IIIB. Bars = SEM of five mice per group. The difference is significant at \(P < 0.01\) by Student’s test.

Fig. 13 demonstrates that protection induced by mucosal immunization with HIV-peptide is dependent on CD8 positive T-cells. BALB/c mice were treated IP with 0.5 mg monoclonal anti-CD8 antibody (clone 2.43, NIH Frederick, MD) one day before and after each immunization and also two days before and three days after the challenge with vPE16. Mice were challenged intrarectally with \(2 \times 10^7\) pfu of vPE16 vaccinia virus expressing gp 160IIIB.

Figs. 14A and 14B demonstrate that mucosal immunization with HIV-1 peptide induces mucosal CTL responses and stimulates protective immunity against intrarectal recombinant HIV-1 vaccinia challenge. Figure 14A depicts induction of the mucosal and systemic CTL responses by different routes of immunization with synthetic peptide HIV vaccine. Killing of peptide-pulsed targets (closed bars) is compared with killing of unpulsed targets (open bars) at an effector-to-target ratio of 50:1. Panel A depicts results or immunizing on day 35 (IR or SC--bar 1, no immunogen; bar 2, SC; bar 3, IR) BALB/c mice challenged intrarectally with \(2.5 \times 10^7\) plaque-forming units (pfu) of vaccinia virus expressing gp 160IIIB.

Figs. 15A and 15B demonstrate enhancement of the mucosal (Fig. A) and systemic (Fig. B) CTL responses to HIV-1
peptide by the mucosal (not systemic) treatment with rmIL-12. BALB/c mice were treated by the IP route (right panels) or IR route (left panels) with 1μg of the rmIL-12 each day of the IR immunization with PCLUS3-18IIIB (50 μg/mice). On day 35 HIV-specific Peyer’s patch CTL (Fig. 15A) and spleen CTL (Fig. 15B) were studied.

Fig. 16 demonstrates that mucosal treatment with rmIL-12 in DOTAP along with HIV peptide vaccine enhances protection against mucosal challenge with vaccinia virus expressing gp160IIIB (vPE16). Five mice per group were immunized IR on days 0, 7, 14 and 21 with no immunogen (bar 1), with 50 μg PCLUS3-18IIIB alone (bar 2), or with peptide plus 1 μg rmIL-12 in DOTAP (bar 3), and challenged on day 35 intrarectally with 5 X 10⁷ pfu of vaccinia virus expressing gp 160IIIB. Viral pfu in the ovaries were determined six days later.

Figs. 17A and 17B demonstrate that IL-12 cannot act directly in the induction of mucosal CD8+ CTL in the absence of IFNγ. IFNγ⁻/⁻ mice (BALB/c background) were immunized IR with the rmIL-12 (1 μg/mouse) + DOTAP together with peptide. On day 35 HIV-specific Peyer’s patch CTL (Fig. 17A) and spleen CTL (Fig. 17B) were studied. Killing of P18IIIB-I10 pulsed targets by effector cells from immunized IFNγ⁻/⁻ mice (closed squares) or conventional BALB/c mice (closed circles) is compared with killing of unpulsed targets (open squares or circles).

Detailed Description

A. INDUCTION OF MUCOSAL IMMUNITY

The present invention is based on the discovery that IR administration of a peptide, e.g., a synthetic multideterminant HIV-1 gp160 envelope glycoprotein peptide, to a mucosal surface can induce an antigen-specific, protective CTL response in the mucosal immune system, even in the absence of a mucosal adjuvant.

The exemplary synthetic multideterminant peptides (CLUVAC) are composed of subregions containing epitopes that evoke some or all of (a) a helper T cell response, (b) a CTL
response and (c) a high titer of neutralizing antibodies in multiple hosts of a given species expressing a broad range of MHC haplotypes. IR immunization with the HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2), and the mucosal adjuvant CT induced peptide-specific CTL in both the inductive (PP) and the effector (LP) sites of the mucosal immune system, as well as in systemic lymphoid tissue, i.e., SP. In contrast, systemic immunization with peptide vaccine produced HIV-1 peptide-specific CTL only in the SP.

IR immunization induced long-lasting protective immune responses. For example, antigen-specific CTL were found in both mucosal and systemic sites 6 months after immunization. IR immunization with the antigenic peptide elicited significantly stronger CTL responses than IN immunization with the same peptide. While IR administration with PCLUS3-18IIIB (SEQ ID NO:2) induced a significant response when administered alone, the response was enhanced by the inclusion of CT. The CTL were CD8+ T lymphocytes restricted by MHC class I molecules, recognizing MHC class I positive target cells either endogenously expressing HIV-1 gp160 or pulsed with an appropriate gp160 peptide. Induction of both mucosal and systemic CTL response by IR immunization was IL-12-dependent, as shown by inhibition of induction of CTL in mice treated i.p. with anti-IL-12 antibody.

Furthermore, inclusion of IL-12 in the composition of antigenic peptide and CT used for IR immunization resulted in enhanced mucosal and systemic CTL responses relative to the responses elicited by antigenic peptide and CT without IL-12. The dependence on IFNγ of mucosal and systemic CTL generation following IR immunization was demonstrated by the absence of such responses in mice which lack the ability to produce functional IFNγ, e.g., as the result of a premature stop codon in the IFNγ-encoding gene. The stop-codon mutation causes the gene to encode a truncated protein lacking the activity of IFNγ.

IR immunization with PCLUS3-18IIIB (SEQ ID NO:2) protected mice against an IR challenge with a recombinant vaccinia virus expressing HIV-1 IIIB gp160. Thus, an HIV-1
peptide induced CTL responses in the mucosal and systemic immune systems after IR immunization and protected against mucosal challenge with virus expressing HIV-1 gp160.

The immunization method of the invention is useful to induce a mucosal CTL response to any soluble antigen. Accordingly, the invention provides a new method for vaccine administration to elicit immunologic protection against viruses that enter through mucosal barriers, including HIV-1. The method can also be applied to achieving protection from infection by certain bacteria (e.g., L. monocytogenes) and protozoa (e.g., G. lamblia) that establish infection via mucosae. In addition, since mucosal immunization results in systemic generation of CTL activity, the protocol can also be useful in prevention of infection by microorganisms that enter through non-mucosal routes. The method is also useful for immunotherapy of infections that are mucosally or non-mucosally established. Finally, the methods can be utilized for immunotherapy of cancer, both in the region of and remote from a mucosal surface.

B. METHODS OF IMMUNIZATION

The invention features methods for protecting subjects from infection by intracellular microorganisms such as viruses as well as intracellular bacteria and protozoans. The methods involve induction of CTL responses specific for antigenic peptides derived from proteins encoded by genes of relevant microbes and expressed in association with MHC class I molecules on the surface of infected cells. This is achieved by delivery of an appropriate soluble antigen to a mucosal surface, e.g., rectal, vaginal, nasopharyngeal, gastric or tracheal mucosae. Relevant microorganisms include but are not restricted to those that enter their hosts via mucosal barriers, e.g. HIV-1, influenza virus, enteric viruses such as rotaviruses, hepatitis A virus, papilloma virus, feline immunodeficiency virus, feline leukemia virus, simian immunodeficiency virus, intracellular bacterial pathogens, e.g., L. monocytogenes and mycobacteria such as M. tuberculosis and M. leprae. Since the responses elicited by
mucosal immunization occur in the systemic as well as the mucosal immune system, the methods can also be applied to protection from infection by intracellular pathogens that enter their hosts via non-mucosal (e.g., HIV-1 in some scenarios such as a "stick" by a contaminated syringe needle, rabies virus and malarial protozoans) as well as mucosal routes. In light of the above considerations, immunization via mucosae can also be used for immunotherapy of intracellular infections.

Finally, the mucosal immunotherapy of the invention can be applied to subjects with cancer, particularly, but not limited to, those with solid tumors in the region of the relevant mucosa, e.g., colonic, rectal, bladder, ovarian, uterine, vaginal, prostatic, nasopharyngeal, lung or certain melanoma tumors. Tumor immunity is substantially mediated by CTL with specificity for tumor associated peptides (e.g., prostate specific antigen peptides in prostatic cancer, carcinoembryonic antigen peptides in colon cancer, human papilloma virus peptides in bladder cancer and MART1, gp100 and tyrosinase peptides in melanoma) (Rosenberg et al. (1994) J.N.C.I. 76:1159; Kawakami et al. (1994) Proc. Natl. Acad. Sci. USA. 91:3515) bound to MHC class I molecules on the surface of tumor cells.

B.1 Antigenic Polypeptides

A soluble antigen to be administered according to the invention can be any soluble carbohydrate or peptide antigen, e.g., one containing all or part of the amino acid sequence of a peptide which is naturally expressed in association with a MHC class I molecule on the surface of a cell infected with relevant microbe or expressed on a tumor cell. In the case of infected cells, the cell surface expressed peptide is derived from a protein either encoded by genes of the infectious agent or whose expression is induced by the infectious agent. Thus, the soluble antigen can be a full length, naturally occurring polypeptide, e.g., full length HIV-1 gp160 or gp120 or an antigenic fragment thereof.
Antigen-specific recognition by CTL involves interactions between components of the antigen-specific T cell receptor on the surface of the CTL and residues on both the antigenic peptide and the MHC class I molecule to which the peptide is bound. Thus the soluble antigen can also be a fragment (i.e., a peptide) of the naturally occurring polypeptide that is either (a) itself capable of binding to MHC class I molecules of multiple haplotypes on the surface of antigen presenting cells (APC) and stimulating CD8+ T cell responses in subjects expressing these MHC class I haplotypes or (b) which can be proteolytically processed by APC into fragments with these properties. Ways of establishing the ability of a candidate peptide to stimulate a CTL response in the context of multiple MHC class I haplotypes are well known to one of ordinary skill in the art and are amply described in co-pending U.S. patent application 08/060,988 incorporated herein by reference in its entirety.

Antigenic peptides can be engineered to bind to MHC class II molecules of multiple haplotypes and will be recognized by CD4+ helper T cell precursor cells of subjects expressing multiple MHC class II haplotypes. Ways of establishing the ability of a candidate peptide to stimulate a helper T cell response in the context of multiple MHC class II haplotypes are well known to one of ordinary skill in the art and are amply described in co-pending U.S. patent application 08/060,988 incorporated herein by reference in its entirety.

In addition, antigenic peptides, can contain epitopes that elicit neutralizing antibodies, i.e., those that bind to the relevant microbe or a cell infected with the microbe and neutralize or kill it. Ways of establishing these properties of a candidate peptide are well known to one of ordinary skill in the art and are amply described in copending U.S. patent application 08/060,988 incorporated herein by reference in its entirety.

Antigenic peptides can also elicit antibodies that induce antibody-dependent cellular cytolysis (ADCC) of cells infected by the appropriate microbe or tumor cells expressing at their surface the protein from which the antigenic peptide
was derived. ADCC is a protective mechanism by which specialized cells of the immune system (K cells) recognize the Fc portion of IgG antibody molecules bound to the surface of a target cell and lyse the relevant target cell. Sera, or other body fluids such as rectal lavages, from test subjects mucosally immunized with a candidate peptide can be tested for their ability to mediate ADCC by methods known to an ordinary artisan. A standard cell mediated lympholysis (CML) assay is used. Briefly, a source of lymphoid ADCC effector cells (e.g., peripheral blood mononuclear cells (PBMC) or SP cells) is incubated in vitro with target cells expressing the above described cell surface protein in the presence of various dilutions of test sera. Lysis of the target cells, which can be measured by the release of a detectable label (⁵¹Cr, for example) from prelabeled target cells, is an indication of the presence of ADCC inducing antibodies in the test serum.

Peptides of the invention can be cluster peptide vaccine constructs (CLUVAC). A CLUVAC is a chimeric peptide containing a) a subregion with multiple overlapping helper T cell activating epitopes that can be presented by multiple MHC class II molecules (a cluster peptide), b) a subregion with a CTL activating epitope and c) a subregion that elicits the production of a neutralizing antibody. The peptide sequences containing these epitopes can be derived from different parts of a microbial or tumor associated polypeptide. Alternatively, the CTL inducing and antibody neutralizing epitopes can be located in one subregion of an antigen and the helper epitope(s) can be in a second subregion. CLUVAC and their design are extensively described in co-pending U.S. patent application 08/060,988 incorporated herein by reference in its entirety.

HIV-1 CLUVAC can include the following sequences:

EQMHEIDISSLWDQSLKPCVKRIRQGPGRAFVTIGK (SEQ ID NO:1)
KQIIINMQEVBGKAMYAPPISQIRRIRGPGRAFVTIGK (SEQ ID NO:2)
RDNWRESLYKVKVKEIPGLVAPTRIRQGPGRAFVTIGK (SEQ ID NO:3)
AVAEGTDRVIEVQGAYRAIRHIPRIRQGLERRIQGPGRAFVTIGK (SEQ ID NO:4)
DRVIEVQGAYRAIRHIPRIRQGLERRIQGPGRAFVTIGK (SEQ ID NO:5)
DRVIEVQGAYRAIRRIQGPGRAFVTIGK (SEQ ID NO:6)
AQGAYRAIRHIPRRIRRIQRGPGRAFVTIGK (SEQ ID NO:7)
EQMHEIDIISLWDQSLKPCVHKHIHIGPGRAFYTTKN (SEQ ID NO:8)
KQINMWOEVGKAMYAPPISQIRRIHIGPGRAFYTTKN (SEQ ID NO:9)
RDNWRELYKVKVKEPLGVAPTRIHIHGPGRAFYTTKN (SEQ ID NO:10)
AVAEGETDRIVEVQGAYRAIRHIPRRIRIQGGLERRIHIGPGRAFYTTKN (SEQ ID NO:11)
DRIVEVQGAYRAIRHIPRRIRIQGHLERHIHIGPGRAFYTTKN (SEQ ID NO:12)
DRIVEVQGAYRAIRHIPRRIRIQGHLERHIHIGPGRAFYTTKN (SEQ ID NO:13)
AQGAYRAIRHIPRRIRRIHIGPGRAFYTTKN (SEQ ID NO:14)

It is possible, for example, to link a particular cluster peptide to any peptide containing a CTL and/or neutralizing antibody epitope. Examples of cluster peptides include: cluster peptide 1 whose amino acid sequence (EQMHEIDIISLWDQSLKPCVK) (SEQ ID NO:17) is the first 20 amino acids of the HIV-1 CLUVAC with SEQ ID NO:1; cluster peptide 3 whose amino acid sequence (KQINMWOEVGKAMYAPPISQIR) (SEQ ID NO:18) is the first 24 amino acids of the HIV-1 CLUVAC with SEQ ID NO:2; cluster peptide 4 whose amino acid sequence (RDNWRELYKVKVKEPLGVAPT) (SEQ ID NO:19) is the first 24 amino acids of the HIV-1 CLUVAC with SEQ ID NO:3; and cluster peptide 6 whose amino acid sequence (AVAEGETDRIVEVQGAYRAIRHIPRRIRIQGLER) (SEQ ID NO:20) is the first 33 amino acids of the HIV-1 CLUVAC with SEQ ID NO:4.

Of particular interest are peptides containing the amino acid sequences of SEQ ID NO:2, SEQ ID NO:9 and SEQ ID NO:12. Mucosal responses to the first (PCLUS3-18IIIB) are extensively characterized in the Examples presented infra. PCLUS3-18MN (SEQ ID NO:9) and PCLUS6.1-18MN (SEQ ID NO:12) have been tested in human clinical trials. The CTL epitope within PCLUS3-18IIIB (SEQ ID NO:2) is the amino acid sequence RIQRPGGRAFVTIGK (SEQ ID NO:15).

In some instances, mucosal adjuvants are co-administered at the mucosal tissue site with the soluble antigens. Such adjuvants include, but are not limited to, detoxified bacterial toxins, for example detoxified cholera toxin (CT), E. coli heat labile toxin (LT), mutant CT (MCT) (Yamamoto et al. J. Exp. Med. 185:1203 (1997); Yamamoto et al. Proc. Natl. Acad. Sci. USA 58:5267 (1997); Douce et al.
Infect. Immunity 65:2821 (1997)), mutant E. coli heat labile toxin (MLT) (Di Tommaso et al. Infect. Immunity 64:974 (1996); Partidos et al. Immunology 89:83 (1996), and pertussis toxin. MCT and MLT contain point mutations that substantially ablate toxicity without substantially compromising adjuvant activity relative to that of the parent molecules. In other preferred embodiments, the adjuvant is modified for increased binding to mucosal tissues and/or T-cells. Thus, in one aspect of the invention, CT or other bacterial toxins are conjugated to a mucosal tissue binding agent, such as protein A, an antibody that binds a mucosal tissue- or T-cell-specific protein (eg., a receptor), or a ligand or peptide that binds a mucosal tissue- or T-cell-specific protein (eg., CD4 or CD8). For example, the B chain of CT may be substituted by protein A, conjugated to the CT A chain, to eliminate toxicity and enhance mucosal tissue binding mediated by protein A. Alternatively, the HIV V3 loop which binds CD4 on T cells may be conjugated to the adjuvant to enhance delivery to T cells.

The CTL augmenting cytokines, eg., IL-12 and IFNγ, are, in some instances, either administered systemically or, preferably, coadministered at the mucosal tissue site with the soluble antigen.

Variants of disclosed antigenic peptides can contain different amino acids (preferably conservative changes) from the parental molecules but retain the biological activity of the parental molecules, e.g., the ability to induce specific CTL responses subsequent to mucosal immunization. Such variants can be synthesized by standard means, and are readily tested in assays known to those in the art. Antigenic polypeptides are typically longer than the length of the nominal SEQ ID NOS. recited herein, i.e., the polypeptide can be extended by adding amino acids to the amino or carboxy termini of the peptide defined by any given SEQ ID NO.

Peptides and polypeptides of the invention will also include those described above but modified for in vivo use by:
(a) chemical or recombinant DNA methods to include mammalian signal peptides (Lin et al., *J. Biol. Chem.* 270:14255 (1995)) or the bacterial peptide, penetratin (Joliot et al., *Proc. Natl. Acad. Sci. USA* 88:1864 (1991)), that will serve to direct the peptide across cell and cytoplasmic membranes and/or traffic it to the endoplasmic reticulum (ER) of antigen presenting cells (APC), e.g., dendritic cells which are potent CTL inducers;

(b) addition of a biotin residue which serves to direct the polypeptides or peptide across cell membranes by virtue of its ability to bind specifically to a translocator present on the surface of cells (Chen et al., *Analytical Biochem.* 227:168 (1995));

(c) addition at either or both the amino- and carboxy-terminal ends, of a blocking agent in order to facilitate survival of the relevant polypeptide or peptide in vivo. This can be useful in those situations in which the termini tend to be degraded ("nibbled") by proteases prior to cellular or ER uptake. Such blocking agents can include, without limitation, additional related or unrelated peptide sequences that can be attached to the amino and/or carboxy terminal residues of the polypeptide or peptide to be administered. This can be done either chemically during the synthesis of the peptide or by recombinant DNA technology (see Section 3.3 infra). Alternatively, blocking agents such as pyroglutamic acid or other molecules known to those of average skill in the art can be attached to the amino and/or carboxy terminal residues, or the amino group at the amino terminus or carboxyl group at the carboxy terminus replaced with a different moiety. Likewise, the polypeptides or peptides can be covalently or noncovalently coupled to pharmaceutically acceptable "carrier" proteins prior to administration.

Also of interest are peptidomimetic compounds based upon the amino acid sequence of the peptides of the invention. Peptidomimetic compounds are synthetic compounds having a three-dimensional structure (i.e. a "peptide motif") based upon the three-dimensional structure of a selected peptide. The peptide motif provides the peptidomimetic compound with
the activity of binding to MHC molecules of multiple haplotypes and activating CD8+ and CD4+ T cells from subjects expressing such MHC molecules that is the same or greater than the activity of the peptide from which the peptidomimetic was derived. Peptidomimetic compounds can have additional characteristics that enhance their therapeutic application such as increased cell permeability, greater affinity and/or avidity and prolonged biological half-life. The peptidomimetics of the invention typically have a backbone that is partially or completely non-peptide, but with side groups identical to the side groups of the amino acid residues that occur in the peptide on which the peptidomimetic is based. Several types of chemical bonds, e.g. ester, thioester, thioamide, retroamide, reduced carbonyl, dimethylene and ketomethylene bonds, are known in the art to be generally useful substitutes for peptide bonds in the construction of protease-resistant peptidomimetics.

B.2  In vivo methods to deliver soluble antigens to mucosae of a subject

In methods of the invention that induce mucosal CTL responses, a soluble antigen is delivered to antigen presenting cells of the inductive mucosal immune system (e.g., PP of the intestine). Delivery involves administering to a subject either the soluble antigen itself, e.g., an antigenic peptide, an expression vector encoding the soluble antigen, or cells transfected or transduced with the vector.

B.2.1 Administration of soluble antigen

Soluble antigens can be delivered to the mucosal immune system of a mammal using techniques substantially the same as those described infra for delivery to human subjects. Examples of appropriate mammals include but are not restricted to humans, non-human primates, horses, cattle, sheep, dogs, cats, mice, rats, guinea pigs, hamsters, rabbits and goats.

A soluble antigen of the invention can be delivered to the mucosal immune system of a human in its unmodified state, dissolved in an appropriate physiological solution,
e.g., physiological saline. Alternatively, it can be modified as detailed in Section B.1 in order to facilitate transport across cell and/or intracellular membranes and to prevent extracellular or intracellular degradation. Its transport across biological membranes can also be enhanced by delivering it encapsulated in liposomes using known methods (Gabizon et al., Cancer Res. 50:6371 (1990); Ranade, J. Clin. Pharmacol. 29:685 (1989)) or an appropriate biodegradable polymeric microparticle (also referred to as a "microsphere", "nanosphere", "nanoparticle", or "microcapsule"). Naturally, it is desirable that the soluble antigens be selectively targeted to the mucosae. This can be achieved by contacting the soluble antigens directly with the relevant mucosal surface, e.g., by IR, IVG, IN, intrapharyngeal (IPG) or IT infusion or implantation. CTL activity (systemic or mucosal) induced by IR immunization can be two-fold, preferably five-fold, more preferably twenty-fold, even more preferably fifty-fold and most preferably two hundred-fold greater than that induced by IN immunization.

Soluble antigens of the invention can be delivered in liposomes into which have been incorporated ligands for receptors on relevant cells (e.g., dendritic cell or macrophage APC) or antibodies to cell-surface markers expressed by these cells. Thus, for example, an antibody specific for a dendritic cell surface marker can direct liposomes containing both the anti-dendritic cell antibody and the relevant soluble antigen to dendritic cells.

The soluble antigens can be administered mucosally either alone or together with a mucosal adjuvant. Suitable adjuvants include, but are not restricted to, CT, MCT and MLT. IL-12 and/or IFNγ can also be administered to the subject. Thus, a soluble antigen can be administered with an adjuvant, with IL-12 (in the absence of an adjuvant), with IFNγ (in the absence of an adjuvant), with both IL-12 and IFNγ (in the absence of an adjuvant), with an adjuvant and IL-12, with an adjuvant and IFNγ, or with an adjuvant and both IL-12 and IFNγ. The IL-12 and IFNγ can be administered systemically or co-administered mucosally with the peptide. When the soluble
antigen is administered encapsulated in liposomes or microparticles, the IL-12 and/or IFNγ can be co-incorporated into the same liposomes/microparticles or incorporated into separate liposomes/microparticles. Alternatively, the cytokines can be administered in a free, soluble form. Examples of other cytokines that can be used, singly or in combination, are granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), or tumor necrosis factor α (TNFα). These and other immunopotentiating cytokines are administered as discussed above for IL-12 and IFNγ. Administration can be single or multiple (two, three, four, five, six, eight or twelve administrations, for example). Where multiple, they can be spaced from one day to one year apart. When using peptides without a helper T-cell epitope, it can be necessary to carry out such multiple administrations.

In preferred aspects of the invention formulations of soluble antigen are prepared for intrarectal administration (i.e., delivery to the rectum, colon, sigmoid colon, or distal colon), eg., by formulating the soluble antigen in a rectal enema, foam, suppository, or topical gel. These rectal delivery formulations are adapted for improved delivery, stability, and/or absorption in the rectum, eg., by combining the soluble antigen with one or more known intrarectal delivery base or carrier materials, intrarectal absorption-promoting materials, and/or stabilizers.

Preferred base formulations for rectal delivery of soluble antigen within the methods of the invention include hydrophilic and hydrophobic vehicles or carriers such as those commonly used in formulating suppositories and rectal emulsion or gel preparations. Thus, emulsion vehicles are used which incorporate soluble antigen in an oil/water emulsion suitable for intrarectal administration. Alternatively, gel formulations are provided which incorporate the soluble antigen in a homogenous gel carrier, for example a polyoxyethylene gel such as polyoxyethylene(20)cetyl ether (BC-20TX). When the antigen is formulated in a rectally compatible foam, a non-CFC propellant foam is preferred.
Preferred carriers or delivery vehicles for use within the invention are conventional suppository base materials that are adapted for intrarectal use, such as polyethyleneglycols. Exemplary polyethyleneglycols known and available in art include PEG 400, PEG 1500, PEG 2000, PEG 4000 or PEG 6000. Preferred bases in this context include witepsol H15, witepsol W35, witepsol E85. These are selected for their desired lipophilic and/or hydrophilic properties, and are often selected to form a multiple layer suppository with both lipophilic and hydrophilic base layers. Preferred lipophilic base materials are macrogols of low molecular weight. Particular examples of suitable lipophilic bases include propyleneglycol dicaprylate (Sefsol 228) and Miglyol810. A preferred hydrophilic base is PEG, eg., PEG 400. In one formulation useful within the invention, hydroxypropylcellulose-H (HPC) and/or carbopol-934P (CP) are used as bases for an inner suppository layer, and a witepsol base is used for the outer layer.

Crystalline cellulose or other common stabilizing agents can be added in selected amounts (eg., 30-60% by weight) to the base to promote sustained release.

Selection of base materials, stabilizers, and absorption-promoting agents for formulating intrarectal delivery compositions, particularly suppositories, is determined according to conventional methods, eg., based on melting and drop rates, breaking hardness, disintegration and special breaking times, spreading properties, and diffusion rates. Preferred values of these various properties are known and can be readily determined to adjust antigen formulations to be suitable for intrarectal use. In particular, appropriate values for formulating a time-release suppository having appropriate structural and chemical properties for rectal delivery are known or readily ascertained. In this context, conventional additives, eg., softeners such as neutral oils, Estasan, etc. may be included to optimize consistency, rate of delivery and other characteristics of the formulation.
In formulating mucosal delivery compositions, it is also desired to include absorption-promoting agents to enhance delivery of the soluble antigen and, optionally, the CTL-stimulatory cytokine, to the mucosal surface. A variety of absorption promoting agents (i.e., agents which enhance release or diffusion of the antigen and/or CTL-stimulatory cytokine from the delivery vehicle or base, or enhance delivery of the antigen and/or CTL-stimulatory cytokine to the mucosal tissue or T-cells, for example by enhancing membrane penetration) are known in the art and are useful in mucosal delivery formulations of the invention, eg., for inclusion in intrarectal delivery formulations, particularly suppositories. These include, but are not limited to, surfactants (eg., tween 80), mixed micelles, enamines, nitric oxide donors (eg., S-nitroso-N-acetyl-DL-penicillamine, NOR1, NOR4—which are preferably co-administered with an NO scavenger such as carboxy-PITO or doclofenac sodium), sodium salicylate, glycerol esters of acetoacetic acid (eg., glyceryl-1,3-diaceotoacetor or 1,2-isopropylideneglycerine-3-acetoacetate) and other release-diffusion/absorption-promoting agents adapted for mucosal delivery.

Absorption-promoting agents useful within the invention include a variety of compounds specifically adapted for intrarectal use. In this context, the rate and extent of rectal drug absorption are often lower than with oral absorption, possibly an inherent factor owing to the relatively small surface area available for drug uptake. In addition, the composition of rectal formulations (solid vs liquid, nature of the suppository base) is an important factor in the absorption process. This relationship between formulation and drug uptake has been clearly demonstrated. Thus, coadministration of absorption-promoting agents (eg., surfactants, sodium salicylate, enamines) represents a key approach towards optimizing rectal drug absorption.

Rectal drug delivery in a site- and rate-controlled manner using suppositories, enemas, osmotic pumps, or hydrogel formulations provides a range of options for manipulating mucosal delivery, eg., controlling concentrations, time-
release, and immunogen-drug effects. Absorption from aqueous and alcoholic solutions may occur very rapidly, but absorption from suppositories is generally slower and very much dependent on the nature of the suppository base, the use of surfactants or other additives, particle size of the active ingredients, etc.

Accordingly, preferred formulations for administering soluble antigens and CTL-stimulatory cytokines within the methods of the invention are designed to optimize mucosal delivery. These agents may thus include clycodextrins and beta-cyclodextrin derivatives (eg., 2-hydroxypropyl-beta-cyclodextrin and heptakis(2,6-di-O-methyl-beta-cyclodextrin). These compounds, preferably conjugated with one or more of the active ingredients and formulated in an oleaginous base, are well documented to enhance bioavailability in intrarectal formulations. Other absorption-enhancing agents adapted for intrarectal delivery include medium-chain fatty acids, including mono- and diglycerides (eg., sodium caprate--extracts of coconut oil, Capmul), and triglycerides (eg., amyloexetin, Estaram 299, Miglyol 810).

It is well known in the medical arts that dosages for any one human subject depend on many factors, as well as the particular compound to be administered, the time and route of administration and other drugs being administered concurrently. Dosages for the soluble antigens of the invention will vary, but can be approximately 0.01 mg to 100 mg per administration. Dosages for the mucosal adjuvants will be approximately 0.001 mg to 100 mg per administration.

Dosages for IL-12 will be approximately 25 μg/kg to 500 μg/kg and for IFNγ will be 300 KU to 30,000 KU per administration--comparable dosages will be used for other cytokines. For example, 3,000 KU of IFNγ can be administered to a human patient once per week. Methods of determining optimal doses are well known to pharmacologists and physicians of ordinary skill. Routes will be, as recited supra, mucosal, e.g., IR, IG, IVG, IN, IPG or IT.
B.2.2 Administration of soluble antigens utilizing expression vectors encoding the soluble antigens.

An expression vector is composed of or contains a nucleic acid in which a polynucleotide sequence encoding a peptide or polypeptide of the invention is operatively linked to a promoter or enhancer-promoter combination. A promoter is a transcriptional regulatory element composed of a region of a DNA molecule typically within 100 nucleotide pairs upstream of the point at which transcription starts. Another transcriptional regulatory element is an enhancer. An enhancer provides specificity in terms of time, location and expression level. Unlike a promoter, an enhancer can function when located at variable distances from the transcription site, provided a promoter is present. An enhancer can also be located downstream of the transcription initiation site. A coding sequence of an expression vector is operatively linked to a transcription terminating region. To bring a coding sequence under control of a promoter, it is necessary to position the translation initiation site of the translational reading frame of the peptide or polypeptide between one and about fifty nucleotides downstream (3') of the promoter. Examples of particular promoters are provided infra. Expression vectors and methods for their construction are known to those familiar with the art.

Suitable vectors include plasmids, and viral vectors such as herpes viruses, retroviruses, vaccinia viruses, attenuated vaccinia viruses, canary pox viruses, adenoviruses and adeno-associated viruses, among others.

The application of antigen-encoding genes to the mucosal induction of CTL in humans can utilize either in vivo or ex vivo based approaches.

The ex vivo method includes the steps of harvesting cells (e.g., dendritic cells) from a subject, culturing the cells, transducing them with an expression vector, and maintaining the cells under conditions suitable for expression of the soluble antigen. These methods are well known in the art of molecular biology. The transduction step is accomplished by any standard means used for ex vivo gene
therapy, including calcium phosphate, lipofection, electroporation, viral infection, and biolistic gene transfer. Cells that have been successfully transduced are then selected, for example, for expression of a drug resistance gene. The cells can then be lethally irradiated (if desired) and injected or implanted into the subject. IL-12 and IFNγ can be administered either systemically or to the relevant mucosal surface as discussed above (Section B.2.1).

The in vivo approach requires delivery of a genetic construct directly into the mucosa of a subject, preferably targeting it to the cells or tissue of interest (e.g., dendritic cells in PP). This can be achieved by administering it directly to the relevant mucosa (e.g., IR in the case of PP). Tissue specific targeting can also be achieved by the use of a molecular conjugate composed of a plasmid or other vector attached to poly-L-lysine by electrostatic or covalent forces. Poly-L-lysine binds to a ligand that can bind to a receptor on target cells (Cristiano et al. J. Mol. Med 73:479 (1995)). Similarly, cell specific antibodies of the type described supra in Section B.2.1 can be bound to vectors and thereby target them to the relevant cells of the mucosal immune system. A promoter inducing relatively tissue or cell-specific expression can be used to achieve a further level of targeting. Appropriate tissue-specific promoters include, for example, the inducible IL-2 (Thompson et al., Mol. Cell. Biol. 12: 1043 (1992)), IL-4 (Todd et al., J. Exp. Med. 177:1663 (1993)) and IFNγ (Penix et al., J. Exp. Med. 178:483 (1993)) T-cell targeting promoters. These promoters would allow production of the soluble antigens in lymphoid tissue, including mucosal lymphoid tissue, e.g., PP. Naturally, an ideal promoter would be a dendritic cell specific promoter.

Vectors can also be delivered by incorporation into liposomes or other delivery vehicles either alone or co-incorporated with cell specific antibodies, as described supra in Section B.2.1.

DNA or transfected cells can be administered in a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers are biologically compatible vehicles which
are suitable for administration to a human, e.g., physiological saline. A therapeutically effective amount is an amount of the DNA of the invention which is capable of producing a medically desirable result in a treated animal. As is well known in the medical arts, the dosage for any one patient depends upon many factors, including the patient’s size, body surface area, age, the particular compound to be administered, sex, time and route of administration, general health, and other drugs being administered concurrently. Dosages will vary, but a preferred dosage for administration of DNA is from approximately $10^6$ to $10^{12}$ copies of the DNA molecule. This dose can be repeatedly administered, as needed. Routes of administration will be the mucosal routes recited for soluble antigens supra in Section B.2.1. The mucosal adjuvants, IL-12 and IFNγ can also be administered in the same combinations, by the same routes and at the same dosages recited in Section B.2.1.

B.3 Sources of peptides and polypeptides

Peptides and polypeptides used in the methods of the invention can be obtained by a variety of means. Smaller peptides (less than 100 amino acids long) can be conveniently synthesized by standard chemical methods familiar to those skilled in the art (e.g., see Creighton, Proteins: Structures and Molecular Principles, W.H. Freeman and Co., N.Y. (1983)). Larger peptides (longer than 100 amino acids) can be produced by a number of methods including recombinant DNA technology (see infra). Some polypeptides (e.g., HIV-1, gp160, CT, LT, IL-12, or IFNγ) can be purchased from commercial sources.

Polypeptides such as HIV-1, gp160, CT, IL-12, or IFNγ can be purified from biological sources by methods well-known to those skilled in the art (Protein Purification, Principles and Practice, second edition (1987) Scopes, Springer Verlag, N.Y.). They can also be produced in their naturally occurring, truncated, chimeric (as in the CLUVC, for example), or fusion protein forms by recombinant DNA technology using techniques well known in the art. These methods include, for example, in vitro recombinant DNA

A variety of host-expression vector systems can be utilized to express the nucleotide sequences. Where the peptide or polypeptide is soluble, it can be recovered from: (a) the culture, i.e., from the host cell in cases where the peptide or polypeptide is not secreted; or (b) from the culture medium in cases where the peptide or polypeptide is secreted by the cells. The expression systems also encompass engineered host cells that express the polypeptide in situ, i.e., anchored in the cell membrane. Purification or enrichment of the polypeptide from such an expression system can be accomplished using appropriate detergents and lipid micelles and methods well known to those skilled in the art. Alternatively, such engineered host cells themselves can be used in situations where it is important not only to retain the structural and functional characteristics of the protein, but also to assess biological activity.

The expression systems that can be used for purposes of the invention include but are not limited to microorganisms such as bacteria (for example, E. coli and B. subtilis) transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing the nucleotide sequences; yeast transformed with recombinant yeast expression vectors; insect cells infected with recombinant viral expression vectors (baculovirus); plant cell systems infected with recombinant viral expression vectors (e.g., cauliflower mosaic virus, CAMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors; or
mammalian cells (e.g., COS, CHO, BHK, 293, 3T3) harboring recombinant expression constructs containing promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses.

In bacterial systems, a number of expression vectors can be advantageously selected depending upon the use intended for the gene product being expressed. For example, when a large quantity of such a protein is to be produced, e.g., for in vivo immunization, vectors which direct the expression of high levels of fusion protein products that are readily purified can be desirable. Such vectors include, but are not limited to, the E. coli expression vector pUR278, (Ruther et al., EMBO J. 2:1791 (1983)), in which the coding sequence can be ligated individually into the vector in frame with the lacZ coding region so that a fusion protein is produced; pIN vectors (Inouye & Inouye, Nucleic Acids Res. 13:3101 (1985); Van Heeke & Schuster, J. Biol. Chem. 264:5503 (1989)); and the like. pGEX vectors can also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution in the presence of free glutathione. The PGEX vectors are designed to include thrombin or factor Xa protease cleavage sites so that the cloned target gene product can be released from the GST moiety.

In mammalian host cells, a number of viral-based expression systems can be utilized. In cases where an adenovirus is used as an expression vector, the nucleotide sequence of interest can be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene can then be inserted in the adenovirus genome by in vitro or in vivo recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable of expressing the gene product in infected hosts (e.g., See Logan and Shenk, Proc. Natl. Acad. Sci. USA 81:3655 (1984)). Specific
initiation signals can also be required for efficient
translation of inserted nucleotide sequences. These signals
include the ATG initiation codon and adjacent sequences. In
cases where an entire gene or cDNA, including its own
initiation codon and adjacent sequences, is inserted into the
appropriate expression vector, no additional translational
control signals may be needed. However, in cases where only a
portion of the coding sequence is inserted, exogenous
translational control signals, including, perhaps, the ATG
initiation codon, must be provided. Furthermore, the
initiation codon must be in phase with the reading frame of
the desired coding sequence to ensure translation of the
entire insert. These exogenous translational control signals
and initiation codons can be of a variety of origins, both
natural and synthetic. The efficiency of expression can be
enhanced by the inclusion of appropriate transcription
enhancer elements, transcription terminators, etc. (Bittner et

In addition, a host cell strain can be chosen which
modulates the expression of the inserted sequences, or
modifies and processes the gene product in the specific
fashion desired. Such modifications (e.g., glycosylation) and
processing (e.g., cleavage) of protein products may be
important for the function of the protein. Appropriate cell
lines or host systems can be chosen to ensure the correct
modification and processing of the foreign protein expressed.
Mammalian host cells include but are not limited to CHO, VERO,
BHK, HeLa, COS, MDCK, 293, 3T3, and WI38.

For long-term, high-yield production of recombinant
proteins, stable expression is preferred. For example, cell
lines which stably express the sequences described above can
be engineered. Rather than using expression vectors which
contain viral origins of replication, host cells can be
transformed with DNA controlled by appropriate expression
control elements (e.g., promoter, enhancer sequences,
transcription terminators, polyadenylation sites, etc.), and a
selectable marker. Following the introduction of the foreign
DNA, engineered cells can be allowed to grow for 1-2 days in
an enriched medium, and then are switched to a selective medium. The selectable marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines. This method can advantageously be used to engineer cell lines which express the gene product. Such engineered cell lines can be particularly useful in screening and evaluation of compounds that affect the endogenous activity of the gene product.

A fusion protein can be readily purified by utilizing an antibody or a ligand that specifically binds to the fusion protein being expressed. For example, a system described by Janknecht et al., Proc. Natl. Acad. Sci. USA 88:8972 (1991) allows for the ready purification of non-denatured fusion proteins expressed in human cell lines. In this system, the gene of interest is subcloned into a vaccinia recombination plasmid such that the gene’s open reading frame is translationally fused to an amino-terminal tag consisting of six histidine residues. Extracts from cells infected with recombinant vaccinia virus are loaded onto Ni$_2^+$ nitriloacetic acid-agarose columns and histidine-tagged proteins are selectively eluted with imidazole-containing buffers. If desired, the histidine tag can be selectively cleaved with an appropriate enzyme.

The following examples are meant to illustrate the invention and not to limit it.

EXAMPLES

Materials and Methods

Human IR immunization Soluble antigens of the invention are dissolved in a pharmaceutically acceptable carrier, e.g., physiological saline and administered with and without adjuvant and with or without cytokines, e.g., IL-12 and/or IFNγ. The latter can be administered systemically or together with the antigen. Dosages of soluble antigen are approximately 0.001 mg to 100 mg per administration. Administration can be single, or multiple. In the case of
multiple immunizations, there can, for example, be four weekly administrations, followed (if desired) by a booster administration several months (e.g., two, three, four, six, eight or twelve) or several years (e.g., two, three, four, five, ten, twenty or thirty) thereafter. Administration of the antigen (with and without adjuvant and with and without cytokine) can be by IR insertion of a suppository into which the immunizing composition has been incorporated, by enema or by flexible sigmoidoscope.

**CTL activation cultures** At various time points after immunization (different for each experiment), CTL activity was assessed. BALB/c mice were sacrificed and the intestines and SP were surgically removed. Single cell suspensions were prepared from the organs by methods familiar to those of average skill in the art. In the case of PP and LP, the PP were first dissected from the outer surface of the intestine. The rest of the intestine was cut into small fragments 1 to 3 mm square which were suspended in phosphate buffered saline (PBS). The tissue fragment suspension was stirred for 20 minutes at room temperature and shaken under the same conditions to liberate the intraepithelial lymphocytes (IEL) from the tissue. The suspended IEL were discarded and tissue fragments were treated with collagenase type VIII (Sigma) (300U/ml) dissolved in PBS for 1 hour at room temperature to release the LP cells. PP cells were extracted from the dissected PP by the same collagenase treatment. Cells (PP or LP) were then placed on a discontinuous gradient containing 75% and 40% of Percoll, followed by centrifugation at 2,000 r.p.m. for 20 minutes. Lymphocytes were harvested from the 75%/40% interface and washed two times. Immune cells from SP, PP, LP were cultured with 1μM P18IIIIB-I10 peptide at 5 x 10^6 per/milliliter in 24-well culture plates in complete T cell medium (CTM): RPMI 1640 containing 10% fetal bovine serum, 2mM L-glutamine, penicillin (100U/ml), streptomycin (100μg/ml), and 5 x 10^-5 M 2-mercaptoethanol. P18IIIIB-I10 is a peptide containing the minimal essential CTL epitope of P18IIIIB (SEQ ID NO:15) and has the sequence: RGPGRAFVTI (SEQ ID NO:16).
Three days later, 10% supernatant from concanavalin A activated spleen cells was added as a source of interleukin-2 (IL-2). After 7 days of culture, LP lymphocytes (LPL) were stimulated in a second 7 day culture with 1μM P18IIIB-110 peptide (SEQ ID NO:16) together with 4 X 10^6 of 3300-rad irradiated syngeneic SP cells. Immune SP and PP cells were similarly stimulated in vitro for two 7-day culture periods before assay. Cytolytic activity of CTL was measured by a 4-hour assay with ^51Cr labeled P815 targets using a method familiar to those of ordinary skill in the art. For testing the peptide specificity of CTL, ^51Cr-labeled P815 targets were pulsed for 2 hours with peptide at the beginning of the assay. The percent specific ^51Cr release was calculated as 100X (experimental release-spontaneous release)/(maximum release-spontaneous release). Maximum release was determined from supernatants of cells that were lysed by addition of 5% Triton-X 100. As a control, spontaneous release was determined from target cells incubated without added effector cells. Standard errors of the mean of triplicate cultures were all <5% of the mean.

**Viral plaque assay** Six days after IR challenge with recombinant vaccinia virus expressing HIV-1 IIIB gp160 (vPE16), mice were sacrificed. Their ovaries were removed, dissociated into single cell suspensions and assayed for vPE16 titer by plating serial 10-fold dilutions on a plate of BSC-2 indicator cells, staining with crystal violet and counting plaques at each dilution. The minimal detectable level of virus was 70 plaque forming units (pfu).

**Example 1. Comparison of mucosal and systemic CTL responses after mucosal and systemic immunization**

Mice were immunized IR with 4 doses (on days 0, 7, 14 and 21) of the HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50μg/mouse). 5 weeks to 6 months after the first dose, antigen-specific T cells were isolated from PP, LP and SP and tested for the presence of HIV-1 P18IIIB peptide specific CTL (Fig. 1), as described above. Closed squares show killing of
P18IIIIB-I10 (SEQ ID NO:16) -pulsed targets and open diamonds show killing on unpulsed targets. IR immunization induced long-lasting protective immune responses: antigen-specific CTL were detected in mucosal inductive (PP) and effector (LP) sites and in a systemic site (SP) at least 6 months after immunization. In contrast, systemic immunization (s.c. in incomplete Freund’s adjuvant) induced CTL in the spleen but not in the mucosal immune system (i.e., PP and LP).

Example 2. Comparison of CTL responses with and without a mucosal adjuvant

BALB/c mice were immunized IR with 4 doses of the synthetic HIV-1 CLUVAC PCLUS3-18IIIIB (SEQ ID NO:2) (50μg/mouse per immunization) alone, i.e., in the absence of adjuvant or cytokine on days 0, 7, 14 and 21. In parallel, another group of mice was immunized IR with PCLUS3-P18IIIIB (SEQ ID NO:2) HIV-1 peptide in combination with CT (1μg/mouse). On day 35 antigen-specific T cells were isolated from PP, LP and SP. Immune cells from SP, PP, or LP were cultured and tested for antigen specific CTL (Fig. 2), as described above. Closed squares show killing of P18IIIIB-I10 (SEQ ID NO:16) pulsed targets, and open diamonds show killing of unpulsed targets. IR administration of peptide alone induced a significant response. The response was enhanced by the co-administration of CT.

Example 3. CTL induced by mucosal immunization lyse targets expressing HIV-1 gp160 envelope protein

Mice were immunized IR with 4 doses (on days 0, 7, 14 and 21) of the HIV-1 CLUVAC PCLUS3-18IIIIB (SEQ ID NO:2) (5μg/mouse per immunization) in the presence of CT (1μg/mouse). On day 35, antigen-specific T cells were isolated from PP, LP and SP. Immune cells from SP, PP, or LP were cultured as described above. Cytolytic activity of CTL was measured using a standard ⁵¹Cr release assay (Fig. 3).

Three different cell lines were used as target cells: 15-12 cells, 3T3 18 Neo BALB/c cells and P815 cells.
15-12 cells are BALB/c 3T3 fibroblasts transfected with a vector encoding HIV-1 gp160 (Takahashi et al. Proc. Natl. Acad. Sci. USA 85:3105 (1988)); 3T3 18 Neo BALB/c cells are BALB/c 3T3 fibroblasts transfected with an expression vector containing a Neor gene but no gp160 gene; and P815 cells are untransfected cells that present antigenic peptides added to the culture. CTL lysis of gp160 expressing 15-12 cells (closed squares in right panels) was compared to that of control gp160 non-expressing 3T3 18 Neo BALB/c cells (open diamonds right panels). P815 target cells (left panel) were tested in the presence (closed squares) or absence (open diamonds) of P18IIIB-I10 peptide (SEQ ID NO:16) (1µM). The percent specific 51Cr release was calculated as described above. CTL induced by IR immunization killed target cells either endogenously expressing HIV-gp160 or pulsed with P18IIIB peptide (SEQ ID NO:15).

Example 4. CTL induced by IR immunization are IL-12 dependent.

BALB/c mice were immunized IR with 4 doses (on days 0, 7, 14 and 21) of CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50µg/mouse per immunization) in combination with CT (1µg/mouse). One day before and one day after immunization with peptide mice were treated intraperitoneally (i.p.) with anti-IL-12 antibody (0.5 mg/per injection; 4mg/mouse total dose (Fig. 4, right panels) or were untreated (Fig. 4, left panels). On day 35, antigen-specific T cells were isolated from PP, LP and SP. Immune cells from SP, PP, or LP were cultured as described above. Cytolytic activity of CTL was measured (Fig. 4) as described above. For testing the peptide specificity of CTL, 51Cr labeled P815 targets were pulsed with peptide P18IIIB-I10 (SEQ ID NO:16) at the beginning of the assay (closed squares), or (as a control) left unpulsed i.e., without peptide (open diamonds). Induction of both mucosal and systemic CTL responses by IR immunization was IL-12-dependent, as shown by inhibition of induction of CTL in mice treated i.p. with anti-IL-12 antibody.
BALB/c mice were immunized IR on days 0, 7, 14 and 21 with either composition A containing the synthetic HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50μg/mouse per immunization), CT (10μg/mouse per immunization), and recombinant IL-12 (1μg/mouse per immunization), or composition B containing the synthetic HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50μg/mouse per immunization), and CT (10μg/mouse per immunization), on day 35, antigen-specific T cells were isolated from PP and SP. Immune cells from PP (Fig. 9A) and SP (Fig. 9B) were separately cultured and tested for antigen specific CTL, as described above. Fig. 9A shows killing of peptide P18IIIB-I10 (SEQ ID NO:16) pulsed target cells by effector cells from mice immunized with composition A (i.e., antigen with IL-12) (open squares), and Fig. 9B shows killing of the P18IIIB-I10 (SEQ ID NO:16) pulsed target cells by effector cells from mice immunized with composition B (i.e., antigen without IL-12) (open diamonds). In both Figs. 9A and 9B, control data was obtained using the two effector populations tested on unpulsed targets. The enhanced CTL responses of both PP (Fig. 9A) and SP (Fig. 9B) effector cells from mice immunized IR with composition A compared to the responses elicited by IR immunization with composition B indicate that co-administration of IL-12 augments mucosal and systemic CTL responses. These data both provide independent evidence for the role of IL-12 in eliciting mucosal (and systemic) immunity by mucosal administration of antigens and confirm the findings of the antibody inhibition data described above.

Example 5. Intrarectal peptide immunization protects against mucosal challenge with HIV-gp160 expressing recombinant vaccinia virus.

Groups of 5 mice were immunized IR with 4 doses of the HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50μg/mouse per immunization) on days 0, 7, 14 and 21 in combination with CT. On day 35, mice were challenged IR with 2.5 x 10⁷ pfU of vaccinia virus expressing gp160IIIB (vPE16). After 6 days, the mice were killed and their ovaries assayed for the
presence of virus (Fig. 5) as described above. The left bar of Fig. 5 shows virus titer in the ovaries of unimmunized mice and the right bar shows virus titer in the ovaries of immunized mice. IR immunization with PLCUS3-18IIIIB (SEQ ID NO:2) protected mice against IR challenge with vPE16 as shown by a 4.5-log reduction in viral pfu in ovaries compared to unimmunized animals (p<0.005).

Example 6. Comparison of systemic CTL responses six months after IR and IN immunization

Mice were immunized IR or IN with 4 doses (on days 0, 7, 14 and 21) of the HIV-1 CLUVAC PCLUS3-18IIIIB (SEQ ID NO:2) (50μg/mouse). Six months after the first dose, antigen-specific T cells were isolated from SP and tested for the presence of HIV-1 P18IIIIB-I10 (SEQ ID NO:16) peptide specific CTL (Fig. 6A and 6B) as described above. Closed squares show killing of P18IIIIB-I10 (SEQ ID NO:16)-pulsed targets and open diamonds show killing of unpulsed targets. The level of CTL activity induced by IR immunization (Fig. 6A) was higher than that induced by IN immunization (Fig. 6B). These data indicate that IR immunization induced potent, long-lasting, antigen-specific splenic immune responses.

To evaluate different routes of immunization, mice were immunized mucosally (IR, IN, IG) or systemically (SC) with 4 doses (on days 0, 7, 14 and 21) of the HIV-1 CLUVAC PCLUS3-18IIIIB (SEQ ID NO:2) (50μg/mouse). Thirty-five days after the first dose, antigen-specific T cells were isolated from PP and SP and tested for the presence of HIV-1 P18IIIIB--I10 (SEQ ID NO:16) peptide-specific CTL as described above (Fig. 7). Closed bars show killing of P18IIIIR-I10 (SEQ ID NO:16) pulsed targets and open bars show killing on unpulsed targets. The level of CTL activity induced by IR immunization was significantly higher in inductive mucosal tissue (PP) and much higher in systemic immunological tissue (SP) than that induced by the other mucosal routes (IN and IG). CTL activity was detected in both PP and SP after immunization via all three mucosal routes. Systemic immunization (SC) only resulted in significant CTL activity in SP.
Example 7. CTL induced by IR immunization are IFNγ dependent.

Wild-type BALB/c mice and BALB/c mice lacking the ability to produce functional IFNγ (IFNγ⁻/⁻ mice) (Dalton et al., Science 259:1739 (1993); Tishon et al., Virology 212:244 (1995)) were immunized IR with 4 doses (on days 0, 7, 14 and 21) of CLUVAC PCLUS3-18IIIB (SEQ ID NO:2) (50μg/mouse per immunization) in combination with CT (1μg/mouse). On day 35, antigen-specific T cells were isolated from PP, LP and SP. Immune cells from SP, PP, or LP of wild type BALB/c and IFNγ⁻/⁻ mice were cultured as described above. Cytolytic activity of CTL was measured (Fig. 8A and Fig. 8B) as described above. The peptide specificity of CTL was tested as follows: ⁵¹Cr labeled P815 targets were pulsed with peptide P18IIIB-I10 (SEQ ID NO:16) at the beginning of the assay (solid bars), or (as a control) left unpulsed i.e., without peptide (open bars). Induction of both mucosal and systemic CTL responses by IR immunization was IFNγ dependent, as shown by the lack of detectable CTL activity in SP, PP, and LP cells from IFNγ⁻/⁻ mice (Fig. 8B) and potent CTL activity in SP, PP, and LP cells from wild-type BALB/c mice (Fig. 8A).

Summarizing the foregoing results, mucosal, administration of an antigenic peptide to mice results in the induction of a protective CTL response detectable in both the inductive (Peyer’s patch (PP)) and effector (lamina propria (LP)) sites of the intestinal mucosal immune system, as well as in systemic lymphoid tissue, i.e., spleen (SP). The mucosal CTL response is enhanced by co-administering the mucosal adjuvant, cholera toxin (CT) with the antigenic peptide, is inhibited by antibody that specifically binds to (thereby neutralizing the activity of) interleukin-12 (IL-12), and is not detectable in mice lacking the ability to produce functional interferon-γ (IFNγ). Furthermore, including IL-12 in the composition of antigenic peptide and CT used for IR immunization resulted in enhanced PP and SP CTL responses relative to those obtained by IR immunization with antigenic peptide, CT and no IL-12. IR immunization with the viral
peptide resulted in protection from viral infection upon subsequent IR challenge with the appropriate virus.

Example 8. Comparison of different HIV vaccine peptides for use in eliciting viral mucosal protection

Because of the variability of the V3 loop of HIV, further studies were conducted comparing two cluster peptide constructs using the V3 loop and incorporating the CTL epitope P18 from strain IIIB (Ratner et al., Nature 313:277-284, (1985)) or MN (Gurgo et al., Virology 164:531-536, (1998)) of HIV-1. These studies were conducted as an exemplary analysis to demonstrate that HIV vaccine cluster peptide constructs can be prepared from different HIV strains and screened in side-by-side assays to optimize induction of mucosal CTL immunity.

Animals For each of the following Examples, female BALB/c mice were purchased from Frederick Cancer Research Center (Frederick, MD). IFN-γ−/− mice were purchased from Jackson Laboratories (Bar Harbor, ME). Mice used in this study were 6-12 weeks old. The IFN-γ−/− mice were maintained in a specific pathogen-free microisolator environment.

Immunization Mice were immunized with 4 doses of the synthetic HIV peptide vaccine construct PCLUS3-18IIIB (Ahlers et al., J. Immunol. 150:5647-5665, (1993)) (50 μg/mouse for each immunization) on days 0, 7, 14 and 21 in combination with cholera toxin (CT) (10 μg/mouse) (List Biological Laboratories, Campbell, CA) by intrarectal administration. For subcutaneous immunization, incomplete Freund’s adjuvant was used. rm IL,-12 (a generous gift of Genetics Institute, Inc., Cambridge, MA) was delivered either intraperitoneally (IP) (1μg) or intrarectally (1μg) mixed with DOTAP (Boehringer Mannheim), a cationic lipofection agent, along with the peptide vaccine.

Cell purification Five weeks to 6 months after the first dose, antigen-specific T cells were isolated from Peyer’s patches (PP), lamina propria (LP) and the spleen (SP). The Peyer’s patches were carefully excised from the intestinal wall and dissociated into single cells by use of collagenase type VIII, 300U/ml (Sigma) as described, Mega et al., Int.
Our data showed that most PP CD3+ T cells isolated from normal mice were CD4+, while CD3+CD8+ T cells were less frequent. Further, collagenase did not alter expression of CD3, CD4, or CD8 on splenic T cells treated with this enzyme. Lamina propria lymphocyte (LPL) isolation was performed as described, Mega et al., *Int. Immunol.* 3:793-805, (1991). The small intestines were dissected from individual mice and the mesenteric and connective tissues carefully removed. Fecal material was flushed from the lumen with unsupplemented medium (RPMI 1640). After the PP were identified and removed from the intestinal wall, the intestines were opened longitudinally, cut into short segments, and washed extensively in RPMI containing 2% fetal bovine serum (FBS). To remove the epithelial cell layer, tissues were placed into 100 ml of 1 mM EDTA and incubated twice (first for 40 min and then for 20 min) at 37°C with stirring. After the EDTA treatment, tissues were washed in complete RPMI medium for 10 min at room temperature and then placed into 50 ml of RPMI containing 10% FCS and incubated for 15 min at 37°C with stirring. The tissues and medium were transferred to a 50 ml tube and shaken vigorously for 15 seconds, and then the medium containing epithelial cells was removed. This mechanical removal of cells was repeated twice more, using fresh medium each time, in order to completely remove the epithelial cell layer. Histologic examination revealed that the structure of the villi and lamina propria were preserved. To isolate LPL, tissues were cut into small pieces and incubated in RPMI 1640 containing collagenase type VIII 300 U/ml (Sigma) for 50 min at 37°C with stirring. Supernatants containing cells were collected, washed and then re-suspended in complete RPMI 1640. This collagenase dissociation procedure was repeated two times and the isolated cells pooled and washed again. Cells were passed through a cotton-glass wool column to remove dead cells and tissue debris and then layered onto a discontinuous gradient containing 75% and 40% Percoll (Pharmacia Fine Chemicals, Pharmacia Inc., Sweden). After centrifugation (4°C, 600g, 20min), the interface layer between the 75% and 40% Percoll was carefully removed and washed with incomplete
medium. This procedure provided >90% viable lymphocytes with a cell yield of 1.5-2 X 10^6 lymphocytes/mouse. The SP were aseptically removed and single cell suspensions prepared by gently teasing them through sterile screens. The erythrocytes were lysed in Tris-buffered ammonium chloride and the remaining cells washed extensively in RPMI 1640 containing 20/o FBS.

**Cytotoxic T lymphocyte assay** Immune cells from SP, PP, LP were cultured at 5X10^6 per/milliliter in 24-well culture plates in complete T cell medium (CTM): RPMI 1640 containing 10% FBS, 2 mM L-glutamine, penicillin (100 U/ml), streptomycin (100 µg/ml), and 5 x 10^{-5} M 2-mercaptoethanol. Three days later we added 10% concanavalin A supernatant-containing medium as a source of IL-2. LPL were studied after 7 days stimulation with 1µM P18IIIB-I10 peptide together with 4X10^6 of 3300-rad irradiated syngeneic spleen cells. SP and PP cells were stimulated *in vitro* similarly for one or two 7-day culture periods before assay. Cytolytic activity of CTL lines was measured by a 4-hour assay with ^51^Cr labeled targets. Two different cell lines were used as a target cells: 1) 15-12 cells, Takahashi et al., Proc. Natl. Acad. Sci. USA 85:3105-3109 (1988) (BALB/c 3T3 fibroblasts transfected with HIV-IIIB gp160 and endogenously expressing HIV gp160), compared with 18 Neo BALB/c 3T3 fibroblasts transfected with Neo^k^ alone as a control, and 2) P815 targets tested in the presence or absence of I10 peptide (IµM). For testing the peptide specificity of CTL, ^51^Cr labeled P815 targets were pulsed for 2 hours with peptide at the beginning of the assay. The percent specific ^51^Cr release was calculated as 100 X (experimental release-spontaneous release)/(Maximum release - spontaneous release). Maximum release was determined from supernatants of cells that were lysed by addition of 5% Triton-X 100. Spontaneous release was determined from target cells incubated without added effector cells.

**Vaccinia virus** Recombinant vaccinia virus vPE16 expresses the HIV-1 gp160 gene from isolate IIIB (BH8) (Earl et al., J. Virol. 64:2448-2451 (1990)). Expression is directed by the compound early/late P7.5 promoter. Two copies
of the sequence T5NT, which serves as a transcription termination signal for early vaccinia virus genes, are present in the IIIB gp 160 gene. Both of these have been altered in vPE16, so as to retain the original coding sequence and allow early transcription of the gene. The virus, vSC8, is used as a negative control without gp160 (Chakrabarti et al., Mol. Cell Biol. 5:3403-3409 (1985)). Both vPE16 and VSC8 express beta-galactosidase.

**Determination of virus titer in the ovary** On day 35 or 6 months after cluster peptide HIV vaccine immunization, mice were challenged intrarectally with 2.5 X 10^7 or 5 X 10^7 pfu of vaccinia virus expressing gp160IIIB (vPE16). Six days after the challenge with recombinant vaccinia virus expressing HIV-gp160, the mice were killed and ovaries were removed, homogenized, sonicated, and assayed for vPE16 titer by plating serial 10-fold dilutions on a plate of BSC-2 indicator cells staining with crystal violet and counting plaques at each dilution. The minimal detectable level of virus was 100 pfu.

To compare different HIV vaccine cluster peptide constructs, BALB/c mice were immunized intrarectally with peptide (PCLUS3-18IIIB or PCLUS3-18MN) in the presence of CT as a mucosal adjuvant weekly for four weeks (on days 0, 7, 14, and 21). Mice were studied either two weeks later (day 35) or at six months for memory CTL responses in the Peyer’s Patches (PP) or spleen (SP). IR immunization with both peptides PCLUS3-18IIIB or PCLUS3-18MN in combination with CT induces a P18-specific CTL response in the intestinal PP (Fig. 10, panel A) and in the spleen (Fig. 10, panel B).

However, the level of CTL response after IR immunization with PCLUS3-18MN was significantly lower than after IR immunization with PCLUS3-18IIIB. The difference may reflect a higher affinity of the minimal 10-mer P18IIIB-I10, compared to P18MN-T10, for H-2D^d (Takahashi et al., Science 246:118-121 (1989); Takeshita et al., J. Immunol. 154:1973-1986 (1995)). Also, much higher production of IFN-γ by mucosal P18IIIB-specific CD8^+ CTL was observed compared to P18-MN-specific CTL after stimulation with specific peptide in vitro for 48 hours. When tested on 15-12 gp160IIIB-
transfected fibroblast targets endogenously expressing gp 160IIIB, the CTL elicited by immunization with PCLUS3-18IIIB also killed these targets. On the basis of these observations, the PCLUS3-18IIIB construct was used in the protection experiments described below.

Example 9. Mucosal immunization of mice with cluster peptide construct provides longlasting protection from infection with recombinant vaccinia virus expressing HIVg160

In the foregoing Examples, the ability of the mucosal immune responses induced by the HIV cluster peptide vaccine to protect against virus challenge via a mucosal route is demonstrated. To determine the specificity of this protection for recombinant protein HIV-1 IIIB gp160, IR immunized mice were challenged on day 35 after the start of immunization by IR infusion with vaccinia virus expressing HIV-1 IIIB gp 160 (vPE16), or with control vaccinia virus expressing β-galactosidase (vSC8). Unimmunized animals challenged with vPE16 or vSC8 served as controls. Six days after the challenge, mice were sacrificed and the ovaries were removed and assayed for vaccinia titer (6 days after infection with vaccinia, the ovaries contain the highest titer of virus).

IR immunization with the synthetic HIV peptide vaccine protected mice against an IR challenge with vaccinia virus expressing HIV-1 IIIB gp160 compared to unimmunized controls, but did not protect against IR challenge with vaccinia virus expressing only an unrelated protein, β-galactosidase (Fig. 11). Thus, the protection was specific for virus expressing HIV-1 gp160, and any nonspecific inflammatory response induced by the peptide infusion intrarectally was not sufficient to protect against viral challenge two weeks after the last dose of the immunization.

Although the presence of mucosal memory CTL precursors was observed, requiring restimulation in vitro for activity 6 months after IR immunization (Belyakov et al., Proc. Nati. Acad. Sci. 95:1709-1714 (1998)), the strength and duration of protection remained unclear. To resolve this
question, the IR immunized mice were challenged 6 months after
the start of immunization with PCLUS3-18IIIB by IR
administration with vaccinia virus expressing HIV-1 IIIB
(vPE16). This study showed that, even 6 months after HIV
cluster peptide immunization, BALB/c mice exhibit protection
against recombinant HIV-vaccinia challenge (Fig. 12).

Example 10. Protection of mice against mucosal viral
challenge is mediated by CD8 CTL in the mucosal site

To determine the immune mechanism responsible for
protection against mucosal challenge with virus expressing HIV
gp160, mice were treated IP with 0.5 mg monoclonal anti-CD8
antibody (clone 2.43, NIH, Frederick, MD) one day before and
after each of the four immunizations and also two days before
and three days after the challenge with vPE16. This treatment
led to a significant inhibition of the protection against
mucosal challenge with vPE16 (Fig. 13). Thus, protection in
the mucosal site against the virus expressing HIV-1 gp160 is
mediated by CD8 lymphocytes.

Because the HIV peptide constructs disclosed herein
elicit both strong mucosal and systemic MHC class I restricted
95:1709-1714 (1998)), the role of these responses in mediating
protection were further investigated. Because SC immunization
with peptide vaccine elicits splenic but not mucosal CTL,
whereas IR immunization elicits both (Fig. 14), SC and IR
immunizations can be compared to determine whether systemic
CTL are sufficient to protect against mucosal challenge, or
whether local mucosal CTL are necessary. Accordingly, mice
were immunized with PCLUS3-18IIIB plus IFA by the SC route or
with PCLUS3-18IIIB and CT by the IR route on days 0, 7, 14 and
21, and compared these. On day 35 after the start of
immunization, these groups of mice as well as unimmunized
control mice were challenged by IR administration of vaccinia
virus expressing HIV-1 gp160 (vPE16). Finally, six days after
the challenge, mice were sacrificed and their ovaries assayed
for viral titer.
SC immunization with PCLUS3-18IIIB did not protect mice against mucosal challenge with vPE16, whereas IR immunization with the same peptide did protect (Fig. 14B). Thus, protection against mucosal challenge with virus expressing HIV-1 gp160 can be induced only by mucosal immunization of mice, and correlates with local mucosal CTL activity, not with splenic CTL activity. On this basis one can conclude that the CD8⁺ CTL-mediated protection from mucosal challenge with recombinant vaccinia expressing HIV-1 gp160 requires local mucosal CD8⁺ CTL, whereas a systemic CTL response is not sufficient.

Example 11. Cytokine dependence and enhancement of protection by local administration of IL-12 with the vaccine

Induction of mucosal CTL by peptide vaccine is dependent on endogenous IL-12 in that it can be blocked by in vivo treatment of mice with anti-IL-12 (Belyakov et al., Proc. Nati. Acad. Sci. 95:1709-1714, (1998)). To further define the role of IL-12 in the CTL response and protection, BALB/c (H-2D⁺) mice were treated by the IP route with 1 µg of the rmIL-12 each day of the IR immunization with PCLUS3-18IIIB (50 µg/mice). This treatment did not lead to significant changes in the HIV-specific CTL activity in either mucosal or systemic sites (Fig. 15). However, when the mice were treated with the rmIL-12 (1 µg) + DOTAP intrarectally together with peptide, we found a significant increase in the CTL level in both mucosal and systemic sites 35 days after the start of immunization (Fig. 15).

In view of the above results, the possibility that rmIL-12 administered at the local site and time of mucosal immunization might increase protection against mucosal challenge with vaccinia virus expressing HIV-1 gp160 was investigated. To address this question, BALB/c mice were immunized with the rmIL-12 + DOTAP intrarectally together with peptide. The mice were then challenged mice on day 35 after the start of immunization by IR administration of vaccinia virus expressing HIV-1IIIB gp160. In this study, twice the dose of challenge virus was used, whereby the unimmunized mice
had a titer of several times $10^{10}$ rather than several times $10^8$
seen in the previous Examples using a lower challenge dose.
Nevertheless, the immunized mice showed a reduction of greater
than 4 logs in virus titer, as had been seen in the earlier
experiments (Fig. 16, bar 2).

Importantly, IR immunization with the synthetic HIV
peptide vaccine plus rmIL-12 protected mice against an IR
challenge with this gp-160-recombinant vaccinia virus even
more effectively than after the IR immunization with peptide
alone (6-log reduction in viral pfu versus 4-log reduction,
$p<0.05$) (Fig. 16, bar 3 versus bar 2).

As the induction of mucosal CD8+ CTL is strongly
dependent on IL-12 and IFN-γ (Belyakov et al., Proc. Natl.
Acad. Sci. 95:1709-1714 (1998)), further studies were
undertaken herein to determine which cytokine acts directly
for the generation of mucosal CTL, and which acts through a
secondary mechanism. To address this question, IFNγ−/− mice
(BALB/c background) and conventional BALB/c mice were treated
with the rmIL-12 (1 μg/mouse) + DOTAP IR together with
peptide. Mucosal treatment of IR-immunized IFNγ−/− mice with
rmIL-12 did not lead to the induction of mucosal or systemic
CTL (Fig. 17). It thus appears that IL-12 cannot act directly
in the induction of mucosal CD8+ CTL in the absence of IFNγ.

In summary, the foregoing Examples incorporate a
novel viral challenge system in which recombinant vaccinia
virus expressing HIV-1 gp160 is used as a surrogate for HIV-1,
since we cannot infect the mice with HIV-1. Importantly, in
this system, neutralizing antibodies to gp160 cannot protect
against recombinant vaccinia expressing gp160, because the
virus does not incorporate gp160 in the virus particle but
expresses it only in the infected cell. Thus, the protective
immune response must be directed at the infected cell.

The results herein demonstrate that the protective
response is completely dependent on CD8+ cells., by the
abrogation of protection after in vivo depletion of CD8+ cells. Thus, the results show unequivocally that it is CD8+ CTL (whether via lytic activity or via secretion of cytokines or other soluble factors) that protect. Since it has been
shown that protection against vaccinia infection can be mediated by interferon-γ which is secreted by CD8+ CTL in response to antigen stimulation (Harris et al., J. Virol. 69:910-915 (1995)), it is possible that the mechanism involves local secretion of this cytokine by the CTL rather than lysis of infected cell. By either mechanism, the CTL are responsible for mediating protection.

However, since the mucosal immunization induces CTL in both the local mucosal site and the spleen, this result does not distinguish which CTL are responsible for protection. To address this important question, the present Examples take advantage of the fact that subcutaneous immunization with the peptide vaccine induces systemic CTL in the spleen at a level at least as high as that induced by mucosal immunization, but does not induce mucosal CTL. Splenic CTL resulting from both immunization routes kill target cells endogenously expressing HIV-1 gp160. Thus, if systemic CTL against this epitope protected against mucosal challenge, then the subcutaneously immunized mice would have been expected to be protected.

However, the subcutaneously immunized mice showed no evidence of protection against mucosal challenge. Thus, the protection correlated with CTL activity in the local mucosal sites, Peyer’s patches and lamina propria, not with CTL activity in the spleen. The protection was not only mediated by CTL, but also required CTL in the local mucosal site of challenge. Systemic CTL were not sufficient.

Protection mediated by local mucosal CTL appears independently sufficient to mediate a protective immune response. This conclusion is supported by the observation that a two-log reduction in viral pfu occurs in the ovary even at day 2 after mucosal viral challenge (from 2.37 x 10^6 pfu in unimmunized mice to 3.34 x 10^4 pfu in IR immunized mice), before much replication could have occurred in the ovary. This finding indicates that the reduction in titer in the ovary reflects a reduction in the amount of virus that can escape the initial mucosal site of infection. In addition, the enhancement of CTL activity and protection by rmIL-12
depended on local mucosal administration of the cytokine, not systemic administration.

One possible difference between CTL induced by mucosal versus systemic immunization is that the CTL resulting from the SC immunization do not have homing receptors for the GI mucosa, as evidenced by the fact that they are not detected in the lamina propria or Peyer's patches.

The present disclosure represents the first demonstration of protection against GI mucosal challenge requiring local mucosal CTL at the site of challenge. These results have important implications for the development of protective vaccines against mucosal exposure to viruses.

In addition to these results, the Examples provided herein also demonstrate a surprising persistence, not only of memory CTL in the mucosa, but also of protective immunity against mucosal viral challenge. Factors controlling CTL memory, and the role of persistent antigen in maintaining memory CTL, represent another issue that has been of interest for some time (Ahmed and Gray, Science 272:54-60 (1996); Kundig et al., Proc. Natl. Acad. Sci. U.S.A. 93:9716-9723 (1996); Ehl et al., Eur. J. Immunol. 27:3404-3413 (1997)) but has been little studied in the context of mucosal immune responses and protection. In studies of systemic immunity, it was shown (Slifka et al., Blood 90:2103-2108, (1997)) that after infection with lymphocytic choriomeningitis virus (LCMV), CTL memory responses were present in the bone marrow for at least 325 days, indicating long-term persistence of antiviral T cells at this site. While the antigen-specific CD8+ T cell number dropped precipitously following viral clearance, substantial numbers persisted for the life of the mouse (Muraii-Krishna et al., Immunity 8:177-187, (1998)). Upon rechallenge with LCMV, there was rapid expansion of memory CD8+ T cells.

These results indicate that systemic infection with virus can lead to long-term memory and protection. In the case of mucosal CTL memory, it was shown that memory CTL remained at the mucosal site longer if the immunization was via the mucosal route (Gallichan and Rosenthal, J. Exp. Med.
184:1879-1890 (1996)), but the duration of protection by such mucosal CTL was not studied. The ability of mucosal memory CTL to protect will depend in part on the rapidity by which they can expand and be activated after virus exposure, which may relate to the level of virus replication in the mucosal site. For these reasons, it is important to determine the duration of mucosal protection dependent on local mucosal CD8+ CTL. The results herein show persistence of CTL even six months after mucosal immunization with the peptide vaccine construct, without additional reimmunization beyond the initial three-week course. This response is accompanied by protection against mucosal challenge with vaccinia virus expressing gp160.

This latter result is particularly striking because the exemplary immunogen is a peptide administered without any depot form of adjuvant that would maintain the presence of antigen for extended periods. It would be expected that free peptide delivered to the lumen of the gut, or even after transport by mucosal cells, would have a very short half-life. Therefore, either memory CTL can persist in the mucosa at levels sufficient to mediate protection in the absence of persistent antigen, or antigen must persist locally in some cell-bound form, perhaps on N4HC molecules of dendritic cells. Yet another possibility is that the peptide crossreacts with one of the antigens in the mucosal flora and that these crossreactive antigens maintain the memory CTL.

Protection against viral infection by CTL involves more than just the number of CTL induced. Quality of CTLs is as important for in vivo protection as quantity. Previous reports have shown that high-avidity P18IIIB-specific CTLs adoptively transferred into severe combined immunodeficient (SCID) mice were 100- to 1000-fold more effective at viral clearance than the low-avidity CTLs specific for the same peptide-MHC complex, despite the fact that all CTL lines lysed virus infected targets in vitro (Alexander-Miller et al., Proc. Natl. Acad. Sci. U.S.A. 93:4102-4107 (1996)). Thus, the CTL induced by mucosal immunization with the synthetic peptide
vaccine must be present not only in sufficient quantity to protect, but also must be of high enough avidity to protect.

Other aspects of the present disclosure enable optimization of induction of the CTL response and protective immunity. Delivery of certain cytokines such as IL-12 at the site of antigen immunization systemically have been shown to enhance systemic CTL responses (Ahlers et al., J. Immunol. 159:3947-3958 (1997); Iwasaki et al., J. Immunol. 159:4691-4601 (1997); Xiang and Ertl, Immunity 2:129-135 (1995); Irvine et al., J. Immunol. 156:238-245 (1996)). However, no comparable studies have been conducted for mucosal CTL responses.

In the model system disclosed herein, induction of mucosal CTL is dependent on endogenous production of IL-12 by the mouse, because it could be inhibited by anti-IL-12 antibody given in vivo before and after each immunization. In the present Examples, IL-12 co-administered with the antigen intrarectally significantly enhanced CTL induction, and also increased protection against intrarectal vaccinia viral challenge. However, it was striking that IL-12 delivered systemically (i.p.) did not enhance CTL induction either in the systemic sites (eg. spleen) or in the mucosa. This difference may be due to the short half life of IL-12 delivered systemically, which prevented it from surviving long enough to get to the sites of CTL induction. Therefore, for mucosal CTL induction as well as for systemic CTL induction, it is important to deliver the cytokine directly to the site of antigen administration and CTL induction.

Enhancement of CTL induction in the mucosa with recombinant IL-12 is a useful strategy for mucosal vaccine development. In this context, small doses given locally in the mucosal sites are not likely to have the global toxicity that has been associated with systemic administration of this cytokine.

The results herein further show that enhancement of the CTL response in vivo by rmIL-12 is dependent on interferon-γ, as no enhancement was observed in IFNγ−/− mice. At least two mechanisms can explain this result. First, IL-12
may be acting through its well-defined ability to induce production of IFN-γ (Trinchieri, G., Blood 94:4008-4027 (1994)), which then acts directly on CTL precursors. Alternatively, since IFN-γ is important for expression of the IL-12 receptor (Szabo, et al., J. Exp. Med. 185:817-824 (1997)) IL-12 may act directly on CTL, but may not be able to act in IFNγ−/− mice because of the lack of IL-12R expression.

al., Science 234:1563-1566 (1986); Tsubota et al., J. Exp.
Med. 169:1421-1434 (1989); Walker and Levy, Immunology 66:628-
630 (1989); Mackewicz et al., J. Clin. Invest. 87:1462-1466,
Such soluble factors include the chemokines RANTES, MIP-1α,
and MIP-1β (Cocchi et al., Science 270:1811-1815 (1995)).

The present disclosure provides the first
demonstration of protection against mucosal viral challenge
mediated by CTL, which must be present in the local mucosa.
Also provided are methods and compositions for enhancing the
induction of such local mucosal CTL by a peptide vaccine given
together with recombinant IL-12 at the mucosal site. These
methods and compositions for stimulating CTL immunity at local
sites of mucosal exposure represent important tools for
vaccine development to prevent HIV infection or disease in
humans. Because the gastrointestinal tract appears to be a
major site of early SIV and HIV replication, among other
pathogens, and because this and other mucosal sites are
frequent sites of entry for these pathogens, it is
particularly critical to achieve protection at mucosal sites.
The present disclosure satisfies these objects and provides
other advantages as set forth above.

Although the invention has been described with
reference to the presently preferred embodiment, it should be
understood that various modifications can be made without
departing from the spirit of the invention. Accordingly, the
invention is limited only by the following claims.
WHAT IS CLAIMED IS:

1. A method for inducing a protective mucosal cytotoxic T lymphocyte (CTL) response in a mammalian subject comprising contacting a mucosal tissue of the subject with a composition comprising a purified soluble antigen.

2. The method of claim 1, wherein the soluble antigen is an antigenic peptide.

3. The method of claim 1, wherein said composition further comprises an adjuvant.

4. The method of claim 3, wherein the adjuvant is selected from cholera toxin (CT), mutant cholera toxin (MCT), or mutant- E. coli heat labile enterotoxin (MLT).

5. The method of claim 1, further comprising administering a purified cytokine to the subject.

6. The method of claim 1, wherein the cytokine is contacted with a mucosal surface of the subject.

7. The method of claim 5, wherein the purified cytokine is selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor a (TNFa).

8. The method of claim 1, further comprising administering purified interferon-γ to the subject.

9. The method of claim 8, wherein the purified interferon-γ is contacted with a mucosal surface of the subject.

10. The method of claim 5, further comprising administering purified interferon-γ to the subject.
11. The method of claim 10, wherein the purified interferon-\(\gamma\) is contacted with a mucosal surface of the subject.

12. The method of claim 1, wherein said composition further comprises a purified cytokine selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor.

13. The method of claim 1, wherein said composition further comprises purified interferon-\(\gamma\).

14. The method of claim 12, wherein said composition further comprises purified interferon-\(\gamma\).

15. The method of claim 1, wherein the antigen is a peptide derived from a pathogenic virus.

16. The method of claim 15, wherein the pathogenic virus is HIV-1.

17. The method of claim 15, wherein the pathogenic virus is influenza virus.

18. The method of claim 15, wherein the pathogenic virus is rotavirus.

19. The method of claim 1, wherein the antigen is a peptide derived from a pathogenic bacterium or protozoan.

20. The method of claim 1, wherein the antigen is a tumor-associated peptide.

21. The method of claim 1, wherein the antigen is a peptide comprising an HIV-1 cluster peptide vaccine construct (CLUVAC) selected from the group consisting of:

\[
\text{EQMHEDIISLWDQSLPCVKRIQRPGAFVTIK} \quad (\text{SEQ ID NO:1}),
\]
KQIINMWQEVGKAMYAPPSGQIRRIQRGPGAFVTIGK (SEQ ID NO:2),
RDWNRSELYKVKVVKIEPLGVAPTRIQRGPGAFVTIGK (SEQ ID NO:3),
AVAEGRDRIEVEVQGAYRAIRHIPRRIRIQRRIQPGLRRIQGPGAFVTIGK (SEQ ID NO:4),
DRVIEVVQGAYRAIRHIPRRRIRIQRGRELRIQGPGAFVTIGK (SEQ ID NO:5),
DRVIEVVQGAYRAIRRIQRGPGAFVTIGK (SEQ ID NO:6),
AQGAYRAIRHIPRRRIRIQRGPGAFVTIGK (SEQ ID NO:7),
EQMHEERLWQSLKPCVRHIHIGPGAFYTTKN (SEQ ID NO:8),
KQIINMWQEVGKAMYAPPSGQIRRIHIGPGAFYTTKN (SEQ ID NO:9),
RDWNRSELYKVKVVKIEPLGVAPTRIHIHIGPGAFYTTKN (SEQ ID NO:10),
AVAEGRDRIEVEVQGAYRAIRHIPRRRIRIQRGRELRIHIGPGAFYTTKN (SEQ ID NO:11),
DRVIEVVQGAYRAIRHIPRRRIRIQRGRELRIHIGPGAFYTTKN (SEQ ID NO:12),
DRVIEVVQGAYRAIRRIHIGPGAFYTTKN (SEQ ID NO:13) and
AQGAYRAIRHIPRRRIRIHRHIGPGAFYTTKN (SEQ ID NO:14).

22. The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2).

23. The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18MN (SEQ ID NO:9).

24. The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS 6.1-18MN (SEQ ID NO:12).

25. A method for inducing a protective mucosal CTL response in a subject, comprising contacting a mucosal tissue of the subject with a composition comprising a soluble antigen, wherein said composition does not comprise an adjuvant.

26. The method of claim 25, further comprising administering a purified cytokine to the subject.

27. The method of claim 25, wherein the cytokine is contacted with a mucosal surface of the subject.

28. The method of claim 27, wherein the purified cytokine is selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2),
interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor α (TNFα).

29. The method of claim 25, further comprising administering purified interferon-γ to the subject.

30. The method of claim 29, wherein the purified interferon-γ is contacted with a mucosal surface of the subject.

31. The method of claim 26, further comprising administering purified interferon-γ to the subject.

32. The method of claim 31, wherein the purified interferon-γ is contacted with a mucosal surface of the subject.

33. The method of claim 25, wherein said composition further comprises a purified cytokine selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor.

34. The method of claim 25, wherein said composition further comprises purified interferon-γ.

35. The method of claim 33, wherein said composition further comprises purified interferon-γ.

36. The method of claim 25, wherein the antigen is a peptide derived from a pathogenic virus.

37. The method of claim 36, wherein the pathogenic virus is HIV-1.

38. The method of claim 36, wherein the pathogenic virus is influenza virus.
39. The method of claim 36, wherein the pathogenic virus is rotavirus.

40. The method of claim 25, wherein the antigen is a peptide derived from a pathogenic bacterium or protozoan.

41. The method of claim 25, wherein the antigen is a tumor-associated peptide.

42. The method of claim 25, wherein the antigen is a peptide comprising an HIV-1 cluster peptide vaccine construct (CLUVAC) selected from the group consisting of:
   EQMHEIIISLWSDSLPCKVRIRQPGPRAFTIGK (SEQ ID NO:1),
   KQIINMWQEVGKAMYAPPISGQIRRIQRPGRPAFTIGK (SEQ ID NO:2),
   RDNWRESLYKVKVKVKEIPPLGVAPRIRQPGPRAFTIGK (SEQ ID NO:3),
   AVAEQGTRVQVQAYRAIRHPRIRQGRLEIRIRQGPGRAFTIGK (SEQ ID NO:4),
   DRVIEVQVQAYRAIRHPRIRQGRLEIRIRQGPGRAFTIGK (SEQ ID NO:5),
   DRVIEVQVQAYRAIRHPRIRQGRLEIRIRQGPGRAFTIGK (SEQ ID NO:6),
   AQGAYRAIRHPRIRQGRLEIRIRQGPGRAFTIGK (SEQ ID NO:7),
   EQMHEIIISLWSDSLPCKVRIRQPGPRAFTTKN (SEQ ID NO:8),
   KQIINMWQEVGKAMYAPPISGQIRRIHIGPGRAPHTKN (SEQ ID NO:9),
   RDNWRESLYKVKVKVKEIPPLGVAPRIRHIGPGRAPHTKN (SEQ ID NO:10),
   AVAEQGTRVQVQAYRAIRHPRIRQGRLEIRHIGPGRAPHTKN (SEQ ID NO:11),
   DRVIEVQVQAYRAIRHPRIRQGRLEIRHIGPGRAPHTKN (SEQ ID NO:12),
   DRVIEVQVQAYRAIRHPRIRHIGPGRAPHTKN (SEQ ID NO:13) and
   AQGAYRAIRHPRIRHIGPGRAPHTKN (SEQ ID NO:14).

43. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2).

44. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18MN (SEQ ID NO:9).

45. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS 6.1-18MN (SEQ ID NO:12).

46. An immunogenic composition for inducing a protective mucosal CTL response in a subject and adapted for
intrarectal administration comprising a purified soluble antigen formulated for intrarectal delivery to the rectum, colon, sigmoid colon, or distal colon.

47. The immunogenic composition of claim 46, which comprises a rectal enema, foam, suppository, or topical gel.

48. The immunogenic composition of claim 46, further comprising a base, carrier, or absorption-promoting agent adapted for intrarectal delivery.

49. The immunogenic composition of claim 48, which includes a rectal emulsion or gel preparation.

50. The immunogenic composition of claim 48, wherein the soluble antigen is admixed with a homogenous gel carrier.

51. The immunogenic composition of claim 48, wherein the homogenous gel carrier is a polyoxyethylene gel.

52. The immunogenic composition of claim 48, wherein the soluble antigen is admixed with a rectally-compatible foam.

53. The immunogenic composition of claim 48, wherein the soluble antigen is formulated in a suppository.

54. The immunogenic composition of claim 53, wherein the suppository is comprised of a base selected from a polyethyleneglycol, witepsol H15, witepsol W35, witepsol E85, propyleneglycol dicaprylate (Sefsol 228), Miglyol810, hydroxypropylcellulose-H (HPC), or carbopol-934P (CP).

55. The immunogenic composition of claim 53, comprising at least two base materials.
56. The immunogenic composition of claim 46, including a stabilizing agent to minimize intrarectal degradation of the soluble antigen.

57. The immunogenic composition of claim 46, including an absorption-promoting agent.

58. The immunogenic composition of claim 57, wherein the absorption-promoting agent is selected from a surfactant, mixed micelle, enanimes, nitric oxide donor, sodium salicylate, glycerol ester of acetoacetic acid, cyclodextrin or beta-cyclodextrin derivative, or medium-chain fatty acid.

59. The immunogenic composition of claim 46, further comprising an adjuvant.

60. The immunogenic composition of claim 59, wherein the adjuvant is selected from cholera toxin (CT), mutant cholera toxin (MCT), mutant- E. coli heat labile enterotoxin, or pertussis toxin.

61. The immunogenic composition of claim 59, wherein the adjuvant is conjugated to a mucosal tissue or T cell binding agent.

62. The immunogenic composition of claim 61, wherein the mucosal tissue or T cell binding agent is selected from protein A, an antibody that binds a mucosal tissue- or T-cell-specific protein, or a ligand or peptide that binds a mucosal tissue- or T-cell-specific protein.

63. The immunogenic composition of claim 59, wherein the adjuvant comprises a recombinant cholera toxin (CT) having a B chain of CT substituted by protein A conjugated to a CT A chain to eliminate toxicity and enhance mucosal tissue binding mediated by protein A.
64. The immunogenic composition of claim 59, wherein the adjuvant is conjugated to a protein or peptide that binds specifically to T cells.

65. The immunogenic composition of claim 64, wherein the protein or peptide binds to CD4 or CD8.

66. The immunogenic composition of claim 66, wherein the protein or peptide is an HIV V3 loop or T cell-binding peptide fragment thereof.

67. The immunogenic composition of claim 59, further comprising purified IL-12.

68. The immunogenic composition of claim 59, further comprising purified interferon-γ.

69. The immunogenic composition of claim 68, further comprising purified IL-12.
FIG. 1

<table>
<thead>
<tr>
<th>SYSTEMIC</th>
<th>INTRARECTAL</th>
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<td>5 WEEKS</td>
<td>5 WEEKS</td>
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</tbody>
</table>

- **Spleen**
- **Lamina Propria**
- **Peyer's Patch**

% Specific Lysis vs. E:T ratios (12.5:1, 25:1, 50:1) for different tissues over different time periods.
**FIG. 4**

CONTROL

ANTI-IL12

Spleen

% Specific Lysis

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<th>12.5:1</th>
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<tr>
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**FIG. 5**

Log_{10} vPE16 Titer

<table>
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1.08 x 10^8

3.21 x 10^3

(P = 0.004)
**FIG. 11**

VACCINIA VIRUS TITER (log10)

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<tr>
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<td>vPE16</td>
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<td>IR</td>
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</tr>
<tr>
<td>None</td>
<td>vSC8</td>
</tr>
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<td>IR</td>
<td>vSC8</td>
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**FIG. 12**

VACCINIA VIRUS TITER (log10)

<table>
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<th>Time After Immunization</th>
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<td>Unimmun</td>
<td>1.0E+10</td>
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<td>2 months</td>
<td>1.0E+06</td>
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<td>6 months</td>
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FIG. 15A

IL-12 BY IR ROUTE

% SPECIFIC LYSIS

60
40
20
0

6:1 12:1 25:1 50:1
E: T

IL-12 BY IP ROUTE

IMMUNIZATION:
- PEPTIDE + CT + rIL12
- PEPTIDE + CT + rIL12
- PEPTIDE + CT + rIL12
- PEPTIDE + CT + NO rIL12
- PEPTIDE + CT + NO rIL12

TARGET:
P18I1B-I10
NO P18I1B-I10
P18I1B-I10
NO P18I1B-I10
FIG. 15B

 IL-12 BY IR ROUTE

 IL-12 BY IP ROUTE

 IMMUNIZATION:
 - PEPTIDE+CT+IL12
 - PEPTIDE+CT+IL12
 - PEPTIDE+CT+NO rIL12
 - PEPTIDE+CT+NO rIL12

 TARGET:
 - P18IIIIB-110
 - NO P18IIIIB-110
 - P18IIIIB-110
 - NO P18IIIIB-110

% SPECIFIC LYSIS

12:1 25:1 50:1 100:1 E: T

12:1 25:1 50:1 100:1 E: T
FIG. 17B

TARGET:
P18iIB-110
NO P18iIB-110

IMMUNIZATION:
IFN-γ-/-
IFN-γ-/-
BALB/c
BALB/c

% SPECIFIC LYSS
60
40
20
0
6:1
12:1
25:1
50:1
E:1

SUBSTITUTE SHEET (RULE 26)
SEQUENCE LISTING

<110> Berzofsky, Jay A.
       Belyakov, Igor M.
       Derby, Michael A.
       Kelsall, Brian L.
       Strober, Warren
       THE GOVERNMENT OF THE UNITED STATES OF AMERICA, as

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Human immunodeficiency virus type 1

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Human immunodeficiency virus type 1

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Arg