Parainfluenza Virus 5 Priming Followed by SIV/HIV Virus-Like-Particle Boosting Induces Potent and Durable Immune Responses in Nonhuman Primates

Peng Xiao, University of Louisiana Lafayette
Krista Dienger-Stambaugh, Cincinnati Children’s Hospital Medical Center
Xuemin Chen, Emory University
Huiling Wei, University of Georgia
Shannon Phan, University of Georgia
Ashley C Beavis, University of Georgia
Karnail Singh, Cincinnati Children’s Hospital Medical Center
Nihar Deb R Adhikary, University of Louisiana Lafayette
Pooja Tiwari, Georgia Institute of Technology
Francois Villinger, Emory University

Only first 10 authors above; see publication for full author list.

Journal Title: FRONTIERS IN IMMUNOLOGY
Volume: Volume 12
Publisher: FRONTIERS MEDIA SA | 2021-02-25, Pages 623996-623996
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.3389/fimmu.2021.623996
Permanent URL: https://pid.emory.edu/ark:/25593/vsxwt

Final published version: http://dx.doi.org/10.3389/fimmu.2021.623996

Copyright information:
© 2021 Xiao, Dienger-Stambaugh, Chen, Wei, Phan, Beavis, Singh, Adhikary, Tiwari, Villinger, Hê and Spearman

This is an Open Access work distributed under the terms of the Creative Commons Attribution 4.0 International License (https://creativecommons.org/licenses/by/4.0/rdf).
Parainfluenza Virus 5 Priming Followed by SIV/HIV Virus-Like-Particle Boosting Induces Potent and Durable Immune Responses in Nonhuman Primates

Peng Xiao¹, Krista Dienger-Stambaugh², Xuemin Chen³, Huiling Wei⁴, Shannon Phan⁴, Ashley C. Beavis⁴, Karnail Singh², Nihar R. Deb Adhikary¹, Pooja Tiwari⁵, Francois Villinger¹*, Biao He⁴* and Paul Spearman²*

¹ New Iberia Research Center, University of Louisiana at Lafayette, New Iberia, LA, United States, ² Division of Infectious Diseases, Department of Pediatrics, Cincinnati Children’s Hospital Medical Center and University of Cincinnati, Cincinnati, OH, United States, ³ Division of Infectious Diseases, Emory University, Atlanta, GA, United States, ⁴ Department of Infectious Diseases, University of Georgia, Athens, GA, United States, ⁵ Wallace H Coulter Department of Bioengineering, Georgia Institute of Technology, Atlanta, GA, United States

The search for a preventive vaccine against HIV infection remains an ongoing challenge, indicating the need for novel approaches. Parainfluenza virus 5 (PIV5) is a paramyxovirus replicating in the upper airways that is not associated with any animal or human pathology. In animal models, PIV5-vectored vaccines have shown protection against influenza, RSV, and other human pathogens. Here, we generated PIV5 vaccines expressing HIV envelope (Env) and SIV Gag and administered them intranasally to macaques, followed by boosting with virus-like particles (VLPs) containing trimeric HIV Env. Moreover, we compared the immune responses generated by PIV5-SHIV prime/VLPs boost regimen in naïve vs a control group in which pre-existing immunity to the PIV5 vector was established. We demonstrate for the first time that intranasal administration of PIV5-based HIV vaccines is safe, well-tolerated and immunogenic, and that boosting with adjuvanted trimeric Env VLPs enhances humoral and cellular immune responses. The PIV5 prime/VLPs boost regimen induced robust and durable systemic and mucosal Env-specific antibody titers with functional activities including ADCC and neutralization. This regimen also induced highly polyfunctional antigen-specific T cell responses. Importantly, we show that diminished responses due to PIV5 pre-existing immunity can be overcome in part with VLP protein boosts. Overall, these results establish that PIV5-based HIV vaccine candidates are promising and warrant further investigation including moving on to primate challenge studies.

Keywords: parainfluenza virus 5 (PIV5), human immunodeficiency virus (HIV) vaccine, virus-like particles (VLP), rhesus macaques, immune responses
INTRODUCTION

Combination antiretroviral therapy (ART) has markedly changed the clinical outlook for human immunodeficiency virus (HIV)-infected patients and represents a major advance in the fight against HIV/AIDS. Viral suppression from ART has been shown to reduce sexual transmission of HIV, leading to the concept of treatment as prevention as a proven intervention to slow the spread of the epidemic (1–3). Despite this advance, there still are an estimated 1.7 million new HIV infections per year worldwide (4). The development of a safe and effective HIV vaccine remains the most promising strategy for preventing infections on a global basis and eventually ending the HIV pandemic (5–7). Phase III clinical trials of HIV vaccines evaluated thus far have not convincingly led to protection from infection. The RV144 trial employing a recombinant canarypox (ALVAC-HIV) prime and envelope (Env) gp120 protein (AIDSVAX B/E) boost vaccine approach showed evidence of a modest protective effect, and gave hope that a protective HIV vaccine is ultimately achievable (8, 9). However, the more recent failure of HVTN 702 in South Africa, which utilized the same poxvirus prime/gp120 boost strategy, underlines the need for novel approaches (10). The immune correlates required for an HIV vaccine to provide protection in human populations remain incompletely defined. In the absence of well-established correlates of protection, desirable characteristics of an HIV vaccine include the induction of robust and long-lasting functional antibodies against Env and antigen-specific T cell polyfunctional responses for controlling viral replication in the event of infection (11, 12). Vaccine-elicited mucosal immune responses against HIV are less well explored but are also desirable characteristics of vaccine candidates (13).

Parainfluenza virus 5 (PIV5) is a single-stranded, negative-sense RNA virus of the genus Rubulavirus in the family Paramyxoviridae (14). It has several distinguishing characteristics that make it an attractive vector for vaccine development. PIV5 can infect a variety of animals but causes no known disease, and is also non-pathogenic in humans (15–18). Its general use as a kennel cough live vaccine in dogs underscores its safety. PIV5 can be produced in high titers in many cell lines, including Vero cells, which have been approved for use for vaccine production by FDA (19). Furthermore, PIV5 has been used to develop vaccine candidates that elicit mucosal and systemic immune responses and are protective in preclinical studies against diverse viral and bacterial pathogens, including influenza virus, rabies virus, respiratory syncytial virus, and Mycobacterium tuberculosis (20–24).

HIV-1 Env proteins rarely result in the elicitation of highly functional antibodies, even when delivered in the presence of adjuvants (25, 26). Neutralizing antibodies against primary (Tier 2) isolates have been particularly hard to generate in human trials, even when Env-specific antibody titers are robust (27, 28). SOSIP Env trimers have recently shown promise in inducing neutralizing antibodies against homologous Tier 2 HIV isolates in nonhuman primates (29–32). SIV virus-like particles (VLPs) have demonstrated the ability to enhance the titer and breadth of antibody responses in nonhuman primates when employed as a boost following live vector priming (33). Here we employed VLPs displaying native trimeric HIV Env on an SIV Gag core (SHIV VLPs) as a booster following PIV5 priming. We sought to determine if boosting with VLPs enhanced the magnitude and quality of antibody and T cell responses primed by a PIV5-based HIV vaccine regimen.

The efficacy of live biologic vector-based vaccines, such as recombinant viral or bacterial vectors, can be limited by pre-existing immunity against vectors from natural exposure or vaccine-induced immunity (34–37). Therefore, one critical question for the use of PIV5 as a vector, is whether prior exposure to PIV5 would limit the induction of immune responses by PIV5-based vaccines. A previous study has shown that pre-existing neutralizing antibodies against PIV5 in dogs did not negatively affect the immunogenicity of a PIV5-based influenza vaccine (38). However, this potential issue remains to be addressed in nonhuman primates or in humans.

In this study, we found that an HIV vaccine regimen employing an intranasal (IN) prime with recombinant PIV5 vector followed by boosting with SHIV VLPs elicited robust and durable systemic responses, including polyfunctional T cell responses and antibodies mediating neutralization and ADCC. Pre-existing immunity to PIV5 resulted in lower responses to the prime, but these differences could be partly overcome following the VLPs boost.

MATERIALS AND METHODS

Construction of Recombinant Parainfluenza Virus 5-Based Vaccine

Madin-Darby bovine kidney (MDBK) cells were cultured in Dulbecco’s modified Eagle medium (DMEM) with 5% fetal bovine serum (FBS), 5% l-glutamine, 100 IU/ml penicillin, and 100 μg/ml streptomycin (Sigma-Aldrich). The recombinant PIV5 vectors expressing HIV-1 JRFL Env (gp140), ASH-SIVmac239 Gag (SivGag p55), or Ebola virus glycoprotein (EboG) were generated as described before (22). Briefly, plasmids containing full-length genome of PIV5 with the gp140 gene between SH and HN of PIV5, SivGag lacking SH coding sequence and insertion of EboG between HN and L were constructed following standard molecular cloning techniques. Infectious viruses were rescued using these plasmids along with necessary support plasmids encoding PIV5 replication machinery. The rescued viruses were plaque-purified and confirmed by RT-PCR sequencing. The viruses were propagated in MDBK cells as previously described (22). Recombinant PIV5 viruses were harvested at 7 days post infection, clarified by centrifugation at 3,000 rpm for 10 min, aliquoted and stored at -80°C. PIV5 virus titers were determined by plaque assay as previously described (24). MDBK cells were infected with each recombinant PIV5-based HIV vaccine (at an MOI of 5) or were infected with PIV5-mock. At 24 or 48 h post infection, cell lysates were separated by SDS-PAGE, transferred to PVDF membranes, and blotted with a monoclonal antibody specific to the HIV-1 Env (D7324) or SIV Gag (CA183), respectively.
The study consisted of two experimental groups. Group 1 was administrated PIV5-VLP HIV immunization, and group 2 was pre-immunized with PIV5-EboG vaccine. The immunization regimen and doses are detailed in Figure 1. Six rhesus macaques in group 1 were administrated PIV5-VLPs (Env+SIVGag) intranasally (IN) at weeks 0, 6, and 12 with three different doses: 2 × 10² PFU (n=2), 2 × 10⁴ PFU (n=2), and 2 × 10⁶ PFU (n=2), respectively, and all received the same 4th dose (1 × 10⁶ PFU, IN) at week 29, followed by two intramuscular (IM) HIV VLPs boosts (35 and 39 weeks). VLP boosts consisted of purified VLPs (containing 10 µg gp120 Env) premixed with CpG (500 µg) and R848 (750 µg) adjuvants. The four animals in group 2 were all pre-immunized with PIV5-EboG vaccine (1 × 10⁶ PFU, IN) at weeks 0, 6, and 12 to induce pre-existing immunity to PIV5 prior to initiating the same prime/boost immunization regimen, as outlined in Figure 1.

Sample Collection
Peripheral blood mononuclear cells (PBMCs) obtained throughout the immunization course were Ficoll-separated and used fresh for ELISPOT assay or T-cell intracellular cytokine staining (ICS) assay. Plasma samples were collected, aliquoted, and stored at -80°C until analyzed. Bronchoalveolar lavage (BAL) fluid samples were collected following each immunization and bone marrow (BM) samples were collected at the end of study. Nasal and rectal secretions were collected in Merocel sponges and processed as previously described (41), and stored at -80°C until analyzed.

ELISPOT Assay
Total and antigen-specific plasmablasts in peripheral blood or plasma cell responses in bone marrow were determined by ELISPOT assay as previously described (42). Briefly, 96-well multiscreen HTS filter plates (Millipore) were coated overnight at 4°C with 100 µl/well of 5 µg/ml of goat anti-monkey IgG or IgA antibodies (Rockland) or of 5 µg/ml of HIV gp120 (Immune Tech) or Ebola GP protein (IBT Bioservices), respectively. Plates were washed with PBS-0.05% Tween 20 (PBS-T) and blocked with complete medium at 37°C for 2 h. Freshly isolated cells were plated in duplicates in serial 3-fold dilutions and incubated overnight in a 5% CO₂ incubator at 37°C. Plates were washed with PBS-T and incubated with biotin-conjugated anti-monkey IgG or IgA antibodies (Rockland) diluted 1:1000 for 1 h at 37°C. After washing, plates were incubated with horseradish peroxidase (HRP)-conjugated streptavidin diluted 1:1000 (Vector labs) at room temperature and developed using the Immunospot ELISPOT Analyzer (Cellular Technology Limited). The number of spots specific for each Ig isotype was reported as the number of either total or antigen-specific antibody producing cells per million.
Envelope Protein Enzyme-Linked Immunosorbent Assay

Systemic and mucosal binding antibodies to HIV Env protein were assessed by enzyme-linked immunosorbent assay (ELISA) as described previously (39, 43). The antibody titer was defined as the reciprocal of the average test sample dilution at which the optical density (OD) of the test sample (run in triplicate) was greater than the preimmunized control sample (run in triplicate) plus three standard deviations, at the same dilution.

SIV Gag Protein Quantitative Western Blot Analysis

SIV Gag antibody responses were assessed by quantitative Western blot analysis. 100 ng (Env-equivalent) of SHIV VLPs were loaded into a single lane and separated on a 4%–12% Bis-Tris NuPAGE gel (Invitrogen), then transferred to nitrocellulose (Invitrogen). Blots were rinsed with water and placed inside the multi-channel MPX blotting system (LI-COR Biosciences). Total SIV Gag protein was detected using Revert 700 Total Protein Stain Kit (LI-COR Biosciences). Each lane of the MPX device was stained for 5 min then rinsed twice with kit wash buffer, imaged and quantitated in the 700 channel on an Odyssey CLx Infrared Imaging System (LI-COR Biosciences). Blots were replaced in the same orientation in the MPX device and blocked with Odyssey blocking buffer (LI-COR Biosciences) for 1 h before addition of diluted serum (1 to 50 in Odyssey blocking buffer + 0.1% Tween20) and incubated overnight at 4°C. Channels were washed five times with wash buffer (1X PBS + 0.05% Tween-20) before blots were removed from MPX device and probed with Goat anti-Human-IgG IRDye 800CW. The SIV Gag-reactive IR800 signal was imaged and quantitated (background subtracted) and total SIV Gag signal (background subtracted) was used to normalize load. Data are reported as fold change over normalized pre-immune plasma sample signal.

TZM-bl Neutralization Assay

The HIV-1 primary cell-free viruses were obtained from the NIH AIDS Reagent Program and used in TZM-bl neutralization assay adapted from Dr. Montefiori’s laboratory at Duke University. All viruses were propagated in human peripheral blood mononuclear cells activated with Concanavalin A and recombinant interleukin-2 (IL-2). Virus stocks were titrated in TZM-bl cells to derive the 50% tissue culture infective dose (TCID50). For neutralization assay, viruses were titrated at a concentration that resulted in 10^5 relative light units (RLU). Briefly, 50 µl of 2-fold serial dilutions of heat-inactivated plasma and 50 µl of titrated virus were mixed and incubated for 1 h at 37°C in a 96-well flat-bottom plate. Next, 100 µl of TZM-bl cells (10,000/well) in complete DMEM medium containing 15 µg/ml diethylaminoethyl (DEAE) dextran (Sigma-Aldrich) were added to each well, and the 96-well plates were incubated for 48 h. Assay controls included TZM-bl cells alone (cell control, no virus) and TZM-bl cells with virus only (virus control, no test sample). At 48 h, the cells were lysed and luciferase activity was measured immediately on a BioTek luminometer. Neutralization activities were determined by GraphPad Prism (v7.01) software, where values from the experimental wells were compared with the value from control wells. The 50% inhibitory concentration (IC50) was calculated based on the plasma dilution that caused a 50% reduction of RLU compared to the control wells.
**Antibody Dependent Cellular Cytotoxicity Assay**

ADCC was measured using HIV-infected targets and was performed as previously described (44). Briefly, target cells of CEM.NKR.CCR5.CD4+,Luc were infected with 50 ng of HIV-1 JRFL virus and cultured for 4 days. Two-fold serial dilutions of each heat-inactivated plasma sample were added to the infected targets for 20 min at room temperature. CD16-KHYG-1 effector cells were added at a 10:1 effector-to-target ratio and incubated for additional 8 h. The cells were lysed and luciferase activity was measured immediately on a BioTek luminometer. Fifty percent (50%) ADCC killing indicates the plasma titers required for half-maximal cell lysis (similar to 50% inhibitory concentration [IC50] for neutralization).

**Neutralizing Antibody Titers to Parainfluenza Virus 5**

PIV5 neutralizing antibody titers were measured by ELISA as described previously (38). Briefly, serial dilutions of monkey plasma were incubated with 200 TCID50 of PIV5 virus and incubated for 2 h before being added to 96-well plates containing 90% confluent MDBK cells. After 3 days of incubation, the wells were scored after staining with mouse anti PIV5 followed by FITC labeled goat anti-mouse Ig. The neutralizing antibody titer was the highest plasma dilution completely neutralizing 200 TCID50 of PIV5.

**T Cell Intracellular Cytokine Staining Assay**

To detect cytokine production by T cells, freshly isolated PBMC (2x10^6) at two weeks post each immunization were stimulated with peptide pools of HIV Env or SIV Gag at a 1-μg/ml final concentration in the presence of anti-CD28 ECD (1 μg/ml, clone CD28.2; Beckman Coulter), anti-CD107a FITC (1 μg/ml, clone eBioH4A3; eBioscience) and anti-CD95 PE-Cy5 (clone DX2, BD Biosciences). Cells were cultured for 2 h at 37°C before adding Brefeldin A (10 μg/ml, BD Biosciences) for an additional 4 h. For each assay, an unstimulated control (dimethyl sulfoxide, DMSO only) and a positive control (PMA/ionomycin) were included. After 6 h, cells were washed with PBS and an Aqua viability dye (Invitrogen) was used to stain for Live/Dead cells at room temperature (RT) for 15 min. Cells were washed with PBS containing 2% fetal bovine serum (FBS), and surface stained for viability dye (Invitrogen) was used to stain for Live/Dead cells at room temperature (RT) for 15 min. Cells were washed with PBS containing 2% fetal bovine serum (FBS), and surface stained for formaldehyde solution for acquisition on a BD FACSARia Fusion cell sorter. CD3+ T cells were gated for CD4+ and CD8+ T cells, and each population was further phenotyped into memory/naïve subsets by gating CD28 and CD95. The net percent of cytokine-secreting cells was determined by subtraction of the values obtained with DMSO only stimulated samples. Boolean combinations and cytokine positive cells frequency were determined with Flowjo software (version 9.2; ThreeStar Inc). Polyfunctional responses of CD4+ and CD8+ T cells were assessed using the Pestle 2.0 and SPICE 6.0 software (https://niaid.github.io/spice/; Vaccine Research Center, NIAID, NIH) (45).

**Statistical Analyses**

Differences between the two groups were evaluated by the exact Wilcoxon rank sum test. The one-way ANOVA test was used for comparisons across three or more groups. The Spearman rank correlation test was used to assess the relationships between two-group parameters. The distribution of the polyfunctional responses was analyzed by using Student’s t-test and permutation comparison test for pie charts between groups. All statistical analyses were considered significant if they produced P values of < 0.05.

**RESULTS**

**Generation and Characterization of Parainfluenza Virus 5 and Virus-Like Particles Vaccines**

Previously, we reported that the SH-HN junction of PIV5 was the optimal site for inserting the hemagglutinin of influenza virus to give the best protection against infection (24). We also found that deletion of the SH gene resulted in a more efficacious vector (46). To generate PIV5-based HIV vaccine candidates, we inserted HIV-1 JRFL gp140 at the SH-HN junction to generate PIV5-HIVgp140 and inserted SIV Gag between HN and L in a PIV5 vector that included a deletion of the SH gene to produce PIV5ΔSH-SIVGag (Figure 2A). To test for expression and packaging of recombinant PIV5-based HIV vaccines, MDBK cells were infected with recombinant PIV5 viruses or were mock infected. Supernatants were collected and analyzed by SDS-PAGE and Western blot. Protein bands at sizes appropriate for HIV gp140 protein (Figure 2B) and SIV Pr55Gag (Figure 2C) from PIV5 expressing vectors were readily visible in test samples. Similarly, we successfully generated a PIV5 expressing a heterologous Ebola glycoprotein from the Ebolavirus Zaire (EBOV) strain, termed PIV5-EboG for the study (data not shown).

Next, we generated SHIV VLPs as booster immunogen using an inducible 293F cell expressing SIVmac239 Gag and HIV-1 JRFL Env. An uncleaved Gag core was included in the VLP design, as we have previously shown that this creates an immunogen with stable Env incorporation under a variety of conditions (47). Electron microscopic analysis of SIV Gag-Env
particles budding from the plasma membrane was previously shown (33). To further confirm that SHIV VLPs were generated, cellular supernatants were harvested and centrifuged on a 20 to 60% sucrose gradient, followed by Western blotting. As expected, fractions 14 and 15 corresponding to lanes 4 and 5, respectively (density of 1.16 g/ml) were highly enriched in VLPs, as measured by Env and Gag in these fractions (Figure 2D). Purified VLPs used for immunization were quantified by gp120 ELISA.

Evaluation of Antigen-Specific Plasmablast Responses in Blood and Long-Lived Plasma Cell Responses in Bone Marrow

Plasmablasts and plasma cells are short- and long-lived antibody secreting cells (ASC), respectively. To assess the ASC response to the vaccine regimen we initially examined the dynamics of antigen-specific plasmablasts in the peripheral blood over the course of immunization. We found that Env-specific IgG plasmablasts were induced steadily following each of the 4 PIV5-(Env+SivGag) prime immunizations in group 1, but were boosted significantly upon the 1st SHIV VLPs boost ($P<0.01$) (Figure 3A). Of note, there was no consistent dose response noted among the animals receiving different doses of PIV5 SHIV vaccine for the first three doses. PIV5 pre-immunized Group 2 animals exhibited a significantly lower level of Env-specific IgG plasmablast responses after immunization with the four PIV5-SHIV vaccines and the 1st VLPs booster compared to animals in group 1 ($P<0.05$). However, the HIV Env plasmablast levels obtained after the 2nd SHIV VLPs boost were equivalent to those seen in Group 1 animals (Figure 3A). A similar pattern was also observed for Env-specific IgA producing plasmablasts in the two groups (data not shown).

We also evaluated the frequencies of long-lived plasma cells (LLPCs) in the bone marrow (BM) that are responsible for the steady state antibody production. Env-specific IgG and IgA plasma cells in BM were detected at the end of study for group 1 (week 81, 42 weeks post-2nd SHIV VLPs boost) and group 2 (week 71, 16 weeks post-2nd SHIV VLPs boost). Animals in group 1 had relatively high levels of Env-specific IgG and IgA LLPCs at the end of study, which were significantly higher than those measured in group 2 animals ($P<0.01$) (Figure 3B), suggesting attenuation of memory humoral responses by PIV5 pre-existing immunity.

Evaluation of Systemic and Mucosal Antibody Responses

To evaluate systemic humoral immune responses, plasma samples were collected at multiple time points after each of the PIV5 prime and the VLPs boost immunizations. In group 1 animals, binding IgG antibody responses to homologous HIV
Env gp120 were detectable as early as week 8 (2 weeks post 2\textsuperscript{nd} PIV5 immunization). These titers were markedly boosted at week 31 (2 weeks post 4\textsuperscript{th} PIV5 prime), without any significant difference between the animals primed initially with 2×10\textsuperscript{5}, 2×10\textsuperscript{6} or 2×10\textsuperscript{7} PFU PIV5 doses (Fig 4A). Binding antibody responses were significantly enhanced after the 2\textsuperscript{nd} SHIV VLPs boost (week 41, P<0.01), with median peak titers of 3.1×10\textsuperscript{4} (Figure 4A). These responses showed excellent longevity in memory with a median titer of 1.2×10\textsuperscript{8} by 10 weeks which were maintained at these levels for the next 20 weeks. In contrast, Group 2 animals showed no detectable seroconversion to HIV Env after four PIV5 SHIV vaccinations, indicating that pre-existing immunity to the vector limited their responses (Figure 4B). Only after VLPs boosting were HIV Env titers detectable, with a median peak titer of 1.2×10\textsuperscript{9}, significantly lower compared to the group 1 (P<0.01). The differences between non-immune and vector-immune animals was even more apparent when plotted as the geometric mean of Env specific IgG (Figure 4B). This figure emphasizes that Group 2 animals failed to mount an Env-specific response until after the first VLPs administration. Gag-specific immune responses were measured by quantitative Western blotting in Group 1 animals, shown in Figure 4C. This method was chosen because of a favorable signal-to-noise ratio as compared to ELISA in our hands. Specific responses were detectable following the 4\textsuperscript{th} immunization with PIV5 vectors, but became much more prominent following the SHIV VLPs boost (Figure 4C). Gag responses were near background levels in Group 2 but were significantly enhanced following VLPs boosting (Figure 4D). There were no significant correlations between any plasmablast levels measured after the various booster immunizations (PIV5 and VLPs) and corresponding plasma titers for HIV Env antibodies (P>0.05).

Stimulation of antiviral immunity in mucosal immune compartments may be desirable in an HIV vaccine, because HIV transmission occurs primarily at mucosal sites. We therefore investigated whether this intranasal prime/parenteral boost immunization regimen elicited virus-specific mucosal antibody responses. Env-specific IgA and IgG antibodies were detected at low levels in nasal and rectal secretions at week 8 in a few animals, and IgG titers significantly increased at week 31 after the 4\textsuperscript{th} PIV5 prime (P<0.05) correlating with the plasma IgG titers. Both nasal IgA and IgG titers were significantly increased at week 37 after the 1\textsuperscript{st} SHIV VLPs boost (P<0.05) (Figures 5A, B). These responses declined and were maintained at low levels for the next 20 weeks. Env-specific IgA levels in rectal secretions were low and only observed in a few animals even after the 1\textsuperscript{st} SHIV VLPs boost (Figure 5C). Of note, Env-specific rectal IgG levels were significantly increased after the 4\textsuperscript{th} PIV5 prime (P<0.05) in 4 out of the 6 monkeys and after the 1\textsuperscript{st} and 2\textsuperscript{nd} SHIV VLPs boosts (P<0.05) (Figure 5D). In contrast to Group 1, mucosal antibodies were completely undetectable in Group 2 after the 4 PIV5 SHIV immunizations, and barely detectable after two SHIV VLPs boosts (data not shown). Overall, despite employing an intranasal prime, the vaccine did not elicit robust mucosal antibody responses. However, there was some evidence of parenteral boosting of PIV5-primed mucosal IgA and IgG responses.

**Antibody Dependent Cellular Cytotoxicity Activity Against Human Immunodeficiency Virus -1**

To determine whether the vaccine-induced binding antibodies were functional, we initially tested them for non-neutralizing ADCC activity against homologous (HIV-1 JRFL), a tier 2 virus. ADCC activity mediated by Env-specific IgG binding antibody was detectable in three animals of Group 1 at week 31 (2 weeks after the 4\textsuperscript{th} PIV5 vaccine), and all demonstrated ADCC at week 41 (2 weeks after the 2\textsuperscript{nd} SHIV VLPs) (Figure 6A, upper panel). Although the ADCC titer weakened somewhat by week 49, ADCC activity was still detected at week 79 (Figure 6A, upper panel). In contrast, the ADCC titers in Group 2 were essentially negative except for two animals at week 67 (Figure 6A, lower panel). A comparison of both groups for corresponding time points relative to PIV5 SHIV prime/VLPs boost immunization for the 50% ADCC titers (half-maximal cell lysis) showed significantly higher ADCC titers in Group 1 compared to the Group 2 (P<0.01) (Figure 6B), except for the last time point analyzed. There also was a significant correlation between Env-
specific IgG binding titer and 50% ADCC titers for Group 1 animals at week 41 ($r=0.9411$, $P=0.0167$) (Figure 6C). However, there was no such correlation for Group 2 (data not shown).

Neutralizing Activity to Human Immunodeficiency Virus-1

Next, we determined whether antibody-neutralizing activity was induced following PIV5 immunization and SHIV VLPs boosts. Virus neutralization was tested against both homologous (HIV-1 JRFL) and heterologous (clade A and C, tier 2) HIV-1 isolates using the TZM-bl assay. Similar to ADCC, Env-specific neutralizing antibodies were detected in Group 1 monkeys at week 31 (2 weeks after the 4th PIV5 vaccine), and the peak neutralizing titers were reached at week 41 (2 weeks after the 2nd SHIV VLPs boost) (Figure 7A, upper panel). Unlike ADCC, the neutralizing antibody titers gradually declined over time (Figure 7A, upper panel). We further tested Group 1 week 41 plasma samples for neutralization against heterologous HIV-1 viruses (clade A or C). However, among the five heterologous isolates tested, neutralization was observed only against HIV-1 98/BR/004 virus, for two animals (Figure 7B). In contrast to Group 1, neutralization titers were only detected in two animals in Group 2 by week 57 (2 weeks post the 2nd SHIV VLPs) and 67 (Figure 7A, lower panel). A comparison of the 50% neutralization titers (IC50) between the two groups showed statistical significant differences ($P<0.01$), except for the last time point tested (Figure 7C). Moreover, we observed a striking correlation between neutralization and ADCC activity against HIV-1 JRFL in
Group 1 monkeys at week 41 ($r=0.8478$, $P=0.0331$) (Figure 7D). No such correlation was observed for Group 2 (data not shown).

Evaluation of Antigen-Specific T Cell Responses

We investigated the induction of Env- and Gag-specific memory CD4+ and CD8+ T cell responses in PBMC at 2 weeks post each prime and boost immunization via intracellular cytokine staining for T cells secreting of IFN-γ, TNF-α, IL-17A, MIP-1β, and CD107a following HIV Env or SIV Gag peptide pool stimulation (Figure S1). We found that both CD4+ and CD8+ T cells specific for the HIV Env and SIV Gag were expanded and detectable in Group 1 monkeys including those administered the lowest (2 × 10^2 PFU) intranasal dose of PIV5 (Figure S2). The sum of the responses by group are illustrated in Figures 8A, B for CD4 and CD8 T cell responses respectively with a predominance of responses to HIV Env rather than SIV Gag. These responses were low but detectable after even the first PIV5 immunization in Group 1, but markedly boosted after VLPs boosts (Figures 8A, B). In contrast, T cell responses in Group 2 monkeys were significantly lower for both CD4 and CD8 T cells ($P<0.01$). We next evaluated the polyfunctional responses of antigen-specific CD4+ and CD8+ T cells, defined by those cells that simultaneously produce multiple cytokines. As shown in Figure 8C, Env-specific CD4+ and CD8+ responses in Group 1 were highly polyfunctional ($P<0.01$) compared to those in Group 2, with more than 50% cytokine secreting cells exhibiting three (yellow) or four (pink) functions, though no combination of five (blue) were detected. In addition, the IFN-γ+ TNF-α+ CD107a+ secreting cells were the most representative T cell population induced in Group 1 monkeys. No statistical difference was observed for Gag-specific polyfunctional CD4+ T cell responses between the two groups. However, highly polyfunctional Gag-specific CD8+ T cells were induced in Group 1 with about 20% of three or four functions compared to that of 2% in Group 2 ($P<0.05$) (Figure 8C). Indeed, the Group 2 had a significantly higher percentage of monofunctional cytokine profile compared to the Group 1.

Parainfluenza Virus 5 Neutralizing Antibody Responses to Successive Immunizations

PIV5 neutralizing antibodies became detectable as early as 2 weeks after the initial immunizations reaching a titer plateau comprised between 10^2 and 10^3 by week 8 (Figure S3) in animals from both groups, irrespective of the initial doses of PIV5 vaccine in Group 1 (data not shown). These vector-specific neutralizing responses clearly diminished the initial response to immunizations using the same vector in Group 2 (Figures 3–8), although there was still antigen priming as judged by the binding antibody response to a single VLP boost (Figure 4B). Of note also, the PIV5 neutralizing antibody titer plateau suggests that these responses reached a maximum titer following two intranasal doses that was not increased by additional immunizations.

DISCUSSION

The development of a safe and effective vaccine regimen to prevent HIV infection or disease remains an elusive goal.
Evaluating new vectors and prime-boost combinations is warranted as part of the quest for a protective vaccine. PIV5 is a well-characterized live, nonpathogenic virus strain that has been shown to elicit protective responses against multiple viral and bacterial pathogens in animal models, but had not previously been evaluated as an HIV vaccine candidate vector. In this study, we evaluated the immunogenicity of IN administration of recombinant PIV5 vectors expressing HIVgp140 and SIV Gag, followed by boosting with SHIV VLPs displaying trimeric Env on their surface as a mucosal and systemic-targeted vaccine candidate regimen. We included SIV Gag and HIV Env in the vector design, a combination chosen with future SHIV challenge experiments in mind to establish protective efficacy. SHIV VLPs were created that matched the antigens included in the live vector, and were administered parenterally with the expectation that magnitude and durability of B and T cell responses would be enhanced.

The study provided three important observations. First, IN immunizations with PIV5-based HIV vaccine elicited potent systemic and more limited mucosal antibody titers, functional antiviral antibodies mediating ADCC and neutralization, and SHIV-specific T cell responses. Second, boosting with TLR7/8/9-adjuvanted VLPs markedly enhanced antiviral responses at the systemic level in both magnitude and functionality, as demonstrated by higher levels of ADCC, neutralizing antibodies, and the development of polyfunctional antigen-specific CD4+ and CD8+ T cell responses. Notably, antibody responses were maintained for a substantial duration following this prime-boost regimen. Third, pre-existing immunity against PIV5 (induced by PIV5 EboG) markedly reduced HIV and SIV-specific immune responses.

Inhibition of mucosal HIV acquisition is thought to rely primarily on neutralizing antibodies based on passive protection experiments in macaque models (48–51). However, there is also accumulating evidence for a role for Fcγ receptor-mediated effector antibody functions (such as ADCC) to contribute to protection (8, 9, 52–54). We therefore evaluated the magnitude and functionality of humoral responses elicited by the PIV5 prime/VLPs boost regimen. Binding antibody levels were generated with intranasal PIV5 priming, and were significantly boosted by a late dose of PIV5 (at week 29) and by VLPs administration (Figure 6A). ADCC responses were also generated by PIV5 administration, and were substantially boosted by VLPs administration (Figure 6). Remarkably, Env-specific antibodies persisted at high levels 24 weeks following the last VLPs boost, and ADCC responses were also long-lived (Figure 6A). We observed a correlation between neutralization
and ADCC activity against HIV-1 JRFL at the peak time point (Figure 7D), indicating that these two functions may overlap in vivo following this vaccine regimen. Overall, the magnitude, functionality, and durability of the humoral responses were encouraging, supporting plans to address protective efficacy in future studies. We note, however, that neutralization was limited to the homologous JRFL virus, indicating that further refinements will be needed to achieve the difficult goal of substantial neutralization breadth.

T-cell polyfunctionality has been considered an important parameter reflecting the quality of a potent T-cell response (55). Such responses have been demonstrated in HIV-1-infected individuals who are able to control virus without ART and inversely correlate with viremia and disease progression (56, 57), suggesting that CD8+ T-cell polyfunctionality would be a correlate of virus control (58, 59). Conversely, lack of T-cell polyfunctionality would lead to cell exhaustion and functional impairment (60, 61). Thus, the capacity of vaccine induced T cells to produce IFN-γ, TNF-α, IL-17A, MIP-1β, and CD107a was evaluated in this study. We found that PIV5/VLPs vaccine generated Env-specific CD4+ and CD8+ T-cells that were polyfunctional, with >50% of cytokine secreting cells exhibiting three or four functions. Polynfunctional responses were less profound in Group 2 animals with pre-existing immunity. A significant proportion of Gag specific CD8+ T-cells was also induced, which could be advantageous for control of acquired virus. In contrast, monofunctional cells dominated the Gag-specific CD8+ T cell responses induced in Group 2 animals.

Group 1 animals received different amounts of the priming vectors for the initial three doses in an attempt to derive the optimal dose of PIV5 in this model. Statistically-significant differences were not found between the animals administered doses ranging from $2 \times 10^2$ to $2 \times 10^6$ PFU of SHIV PIV5 in Group 1, although there was a trend towards more reliable response with higher doses. This was especially apparent for ADCC titers, suggesting a potential association between the PIV5 dose and antibody maturation. The

![Figure 7](image-url)
antibody maturation process is dependent on both time and amount of antigen exposure, and our data suggest that the lowest (2x10^2 PFU) dose was suboptimal (Figure 6). However, only minimal differences were observed for neutralization antibody titers relative to the different PIV5 doses (Figure 7).

Mucosal IgA antibodies have been associated with protection or delayed acquisition of SHIV (62–64). Intranasal immunization induces antigen-specific immune responses in the upper and lower respiratory tract and genital tract (65, 66). It is hypothesized that intranasal PIV5 vaccine immunization leads to the attraction and maturation of ASCs localized to the mucosa. While we detected Env-specific mucosal IgA in both nasal and rectal secretions following immunization, these levels were very low prior to VLPs boosting. Although IgA is the most abundant antibody isotype in mucosal secretions, we noted that vaccine-induced Env-specific IgG antibodies were higher than those of IgA in both nasal and rectal secretions, indicating that mucosal IgG seen here may represent transudate from the systemic compartment.

Group 2 animals demonstrated a marked reduction in cellular and humoral immune responses that was elicited by induction of vector-specific immunity. This effect was surprising to us, and seemed contradicted by the increasing immune response induced by successive PIV5 immunization in Group 1. Indeed, humoral responses were markedly boosted by PIV5-Gag and PIV5-Env immunization at week 29 (Figures 4–7), as were cell-mediated immune responses (Figure 8). The reasons for the discrepancy between the effectiveness of the day 29 dose in Group 1 and the suppression of PIV5-elicited responses by pre-immunizing with PIV5 in Group 2 are not certain, but we can consider several possibilities. First, the timing of the initial SIV Gag/Env PIV5 immunization in Group 2 was 4 weeks after the last EboG PIV5 administration, whereas the fourth SHIV PIV5 immunization in Group 1 animals was given 17 weeks following the prior dose of PIV5. Thus, it is possible that anti-PIV5 responses had diminished by week 29. A second possibility is that HIV- and SIV-specific anamnestic responses were less inhibited by pre-existing antibodies than was the induction of initial responses (priming). This idea is supported by studies showing that boosting a response requires less antigenic stimulation than initiating a de novo immune response (67–70). A third possibility stems from our observation that humoral responses elicited by EboG PIV5 surpassed those elicited for SHIV by PIV5 (data not shown) suggesting that the EboG be a more potent immunogen, potentially acting as an adjuvant for PIV5 specific responses. The suppression of responses to the vector insert in Group 2 is different from with the one we had previously shown in dogs pre-immunized with PIV5, where subsequent administration of PIV5 expressing influenza hemagglutinin successfully generated influenza-specific immune responses (38). It is possible that there is a species difference in their responses to PIV5. This finding, coupled with the fact that levels of detectable anti-PIV5 titers in canine owning humans are lower than in dogs, suggest that giving multiple doses of PIV5-vectored vaccines may remain an effective mucosal vaccine strategy for human immunization.

In conclusion, we have shown that a PIV5-based SHIV vaccine has the ability to induce potent systemic humoral and
cell-mediated responses, and more limited mucosal antiviral responses. Boosting with adjuvanted trimeric Env VLPs promoted robust and durable systemic antibody responses and polyfunctional T cell responses, with a limited boosting of mucosal responses. While a stringent induction of pre-existing anti PIV5 immunity clearly affected the development of such potent SIV and HIV-specific responses, the significance of this in human populations remains to be determined. Our results support the further preclinical development of PIV5-based vectors as HIV vaccine candidates, including determination of protective efficacy in nonhuman primate models.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The animal study was reviewed and approved by IACUC, University of Louisiana at Lafayette.

AUTHOR CONTRIBUTIONS

BH, PS, and FV designed the study. PX, KD-S, XC, HW, SP, KS, NA, and PT performed the analyses. PX, KS, FV, BH, and PS redacted the manuscript. All authors contributed to the article and approved the submitted version.

REFERENCES

1. Cohen MS, Chen YQ, McCauley M, Gamble T, Hosseinipour MC, Kamarasamy N, et al. Antiretroviral Therapy for the Prevention of HIV-1 Transmission. N Engl J Med (2016) 375:830–9. doi: 10.1056/NEJMoal1600693
2. Cohen MS, Chen YQ, McCauley M, Gamble T, Hosseinipour MC, Kamarasamy N, et al. Prevention of HIV–1 infection with early antiretroviral therapy. N Engl J Med (2011) 365:495–505. doi: 10.1056/NEJMc1110588
3. McNairy ML, Cohen M, El-Sadr WM. Antiretroviral therapy for prevention is a combination strategy. Curr HIV/AIDS Rep (2013) 10:152–8. doi: 10.1007/s11904-013-0152-1
4. UNAIDS. Global HIV & AIDS statistics fact sheet. (2018).
5. Beyrer C, Baral SD, van Griensven F, Goodreau SM, Charliyartaks S, Wirtz AL, et al. Global epidemiology of HIV infection in men who have sex with men. Lancet (2012) 380:367–77. doi: 10.1016/S0140-6736(12)60821-6
6. Frieden TR, Fofoti KE, Mermin J. Applying Public Health Principles to the HIV Epidemic—How Are We Doing? N Engl J Med (2015) 373:2281–7. doi: 10.1056/NEJMsa1513641
7. Fauci AS, Folkers GK, Marston HD. Ending the global HIV/AIDS pandemic: the critical role of an HIV vaccine. Clin Infect Dis (2014) 59(Suppl 2):S80–4. doi: 10.1093/cid/ciu420
8. Reker-Ngarm S, Pitsutithum P, Nitayaphan S, Kaewkungwal J, Chiu J, Paris R, et al. Vaccination with ALVAC and AIDSVAX to prevent HIV-1 infection in Thailand. N Engl J Med (2009) 361:2209–20. doi: 10.1056/NEJMoa0904892
9. Haynes BF, Gilbert PB, McElrath MJ, Tomaras GD, Alam SM, et al. Immune correlates analysis of an HIV–1 vaccine efficacy trial. N Engl J Med (2012) 366:1275–86. doi: 10.1056/NEJMoai1113425

ACKNOWLEDGMENTS

The authors are indebted to the veterinary, research, and animal care staff of the New Iberia Research center for their outstanding support of the animal studies for this report. The studies were supported by a grant from NIH-NIAID R01AI111863 to PS and BH.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.623996/full#supplementary-material

Supplementary Figure 1 | (A) Flow cytometry gating strategy used to identify total memory CD4+ and CD8+ T cell responses in PBMC. (B) Representative ICS staining of CD8+ cells in Group 1 after the 2nd SHIV VLPs boost. Flow dot plots illustrate individual cytokine with no stimulation, HIV Env (N and C) peptides or with SIV Gag peptides pool stimulation. A positive control stimulation with PMA/ionomycin was included in each assay.

Supplementary Figure 2 | Induction of antigen-specific cytokine producing CD4+ and CD8+ cells by intranasal administration of different PIV5 doses. (A, B) Group 1 of CD4+ or CD8+ T cells secreting cytokines specific for the sum of Env and Gag after administration with three different PIV5 doses of 2×106 PFU (green), 2×105 PFU (blue) and 2×106 PFU (red), respectively. (C, D) Group 2 of CD4+ or CD8+ T cells secreting cytokines specific for the sum of Env and Gag after administration with a single PIV5 dose of 2×105 PFU (grey or earth yellow).

Supplementary Figure 3 | (A) Comparison of titers of anti-PIV5 neutralizing antibodies between Group 1 and Group 2 at the indicated time points receiving the same immunization, respectively. (B) Correlation of anti-PIV5 titers relative to PIV5 doses at weeks 2, 8 and 14. The correlation coefficients (r) and P values were derived from Spearman rank analysis. Significant P values are in red font.
Virus Vaccine Candidates after In Vitro and In Vivo Passage. J Virol (2017) 91(19):e00559–17. doi: 10.1128/JVI.00559–17

20. Chen Z, Zhou M, Gao X, Zhang G, Ren G, Granadurui CW, et al. A novel rabies vaccine based on a recombinant parainfluenza virus 5 expressing rabies virus glycoprotein. J Virol (2013) 87:2986–93. doi: 10.1128/JVI.02886-12

21. Li Z, Gabbard JD, Mooney A, Gao X, Chen Z, Place RJ, et al. Single-dose vaccination of a recombinant parainfluenza virus 5 expressing NP from H5N1 virus provides broad immunity against influenza A viruses. J Virol (2013) 87:5985–93. doi: 10.1128/JVI.00120-13

22. Phan SL, Zengel JR, Wei H, Li Z, Wang D, He B. Parainfluenza Virus 5 Expressing Wild-Type or Prefusion Respiratory Syncytial Virus (RSV) Fusion Protein Protects Mice and Cotton Rats from RSV Challenge. J Virol (2017) 91(19):e00560–17. doi: 10.1128/JVI.00560–17

23. Chen Z, Gupta T, Xu P, Phan S, Pickar A, Yau W, et al. Efficacy of parainfluenza virus 5 (PIV5)-based tuberculosis vaccines in mice. Vaccine (2015) 33:7217–24. doi: 10.1016/j.vaccine.2015.10.124

24. Mooney AJ, Li Z, Gabbard JD, He B, Tompkins SM. Recombinant parainfluenza virus 5 vaccine encoding the influenza virus hemagglutinin protects against H5N1 highly pathogenic avian influenza virus infection following intranasal or intramuscular vaccination of BALB/c mice. J Virol (2013) 87:363–71. doi: 10.1128/JVI.02330-12

25. Klasse PJ, Sanders RW, Cerutti A, Moore JP. How can HIV-type-1-Env immunogenicity be improved to facilitate antibody-based vaccine development? AIDS Res Hum Retroviruses (2012) 28:1–15. doi: 10.1089/aid.2011.0053

26. Wright PF, Mestecky J, McElrath MJ, Keefer MC, Gorse GJ, Goepfert PA, et al. VACCINES. HIV-1 neutralizing antibodies but limited breadth of neutralization in human- or chimpanzee-derived adenovirus vectors. J Virol (2007) 81:6594–604. doi: 10.1128/JVI.02497–06

27. Leong ML, Hampi J, Liu W, Mathur S, Bahjat KS, Luckett W, et al. Impact of preexisting vector-specific immunity on vaccine potency: characterization of listeria monocytogenes-specific humoral and cellular immunity in humans and modeling studies using recombinant vaccines in mice. Infect Immun. (2009) 77:3958–68. doi: 10.1128/IAI.01274-08

28. Chen Z, Xu P, Salyards GW, Harvey SB, Rada B, Fu ZP, et al. Evaluating a parainfluenza virus 5-based vaccine in a host with pre-existing immunity against parainfluenza virus 5. PLoS One (2012) 7:e50144. doi: 10.1371/journal.pone.0050144

29. Singh K, Marasini R, Chen X, Ding L, Wang JJ, Xiao P, et al. A Bivalent, Spherical Virus-Like Particle Vaccine Enhances Breadth of Immune Responses against Pathogenic Ebola Viruses in Rhesus Macaques. J Virol (2020) 94(9):e01884–19. doi: 10.1128/JVI.01884–19

30. Weeratna RD, Makinen SR, McCluskie MJ, Davis HL. TLR agonists as vaccine adjuvants: comparison of CpG ODN and Resiquimod (R-848). Vaccine (2005) 23:2563–70. doi: 10.1016/j.vaccine.2005.06.024

31. Kozlowski PA, Lynch RM, Patterson RR, Cu-Uvin S, Fligan TN, Neutra MR. Modified wicked method using Weck-Cel sponges for collection of human rectal secretion and analysis of mucosal HIV antibody. J Acquir Immune Defic Syndr (2004) 36:297–309. doi: 10.1097/00126334-2000080100-00001

32. Wrammert J, Smith K, Miller J, Langley WA, Kokko K, Larsen C, et al. Rapid cloning of high-affinity human monoclonal antibodies against influenza virus. Nature (2008) 453:667–71. doi: 10.1038/nature06890

33. Kasturi SP, Kozlowski PA, Nakaya HI, Burger MC, Russo P, Pham M, et al. Adjuvanted a Simian Immunodeficiency Virus Vaccine with Toll-Like Receptor Ligands Encapsulated in Nanoparticles Induces Persistent Antibody Responses and Enhanced Protection in TRIM5alpha Restrictive Macaques. J Virol (2017) 91(4):e01844–16. doi: 10.1128/JVI.01844–16

34. Alpert MD, Harvey JD, Lauwer WA, Reeves RK, Piatak M Jr., Carville A, et al. ADCC, develops over time during persistent infection with live-attenuated SIV and is associated with complete protection against SIV(mac)251 challenge. PLoS Pathog (2012) 8:e1002890. doi: 10.1371/journal.ppat.1002890

35. Roederer M, Nozzi JL, Nason MC. SPICE: exploration and analysis of post-crosslink multivariate datasets. Cytometry A (2011) 79:167–74. doi: 10.1002/cyto.a.21015

36. Li Z, Gabbard JD, Mooney A, Chen Z, Tompkins SM, He B. Efficacy of parainfluenza virus 5 mutants expressing hemagglutinin from H5N1 influenza A virus in mice. J Virol (2013) 87:9604–9. doi: 10.1128/JVI.01289–13

37. Hammond J, Chen X, Ding L, Foutz T, De Vivo A, zur Megede J, et al. Gp120 stability on HIV-1 viruses and Gag Env pseudovirions is enhanced by an unsealed Gag core. Virology (2003) 314:636–49. doi: 10.1016/S0042-6822(03)00467-7

38. Baba TW, Liska V, Hofmann-Lehmann R, Vlask J, Xu W, Aeyubhunie S, et al. Human neutralizing monoclonal antibodies of the IgG1 subtype protect against mucosal simian-human immunodeficiency virus infection. Nat Med (2000) 6:200–6. doi: 10.1038/72309

39. Mascola JR, Steigler G, VanCott TC, Katinger H, Carpenter CB, Hanson CE, et al. Protection of macaques against vaginal transmission of a pathogenic HIV-1/SIV chimeric virus by passive infusion of neutralizing antibodies. Nat Med (2000) 6:207–10. doi: 10.1016/S1078-2764(00)00029-7

40. Shibata R, Igarashi T, Haigwood N, Buckler-White A, Ogert R, Ross W, et al. Neutralizing antibody directed against the HIV-1 envelope glycoprotein can completely block HIV-1/SIV chimeric virus infections of macaque monkeys. Nat Med (1999) 5:204–10. doi: 10.1038/5568

41. Pegu A, Hessel AJ, Mascola JR, Haigwood NL. Use of broadly neutralizing antibodies for HIV-1 prevention. Immunol Rev (2017) 275:296–312. doi: 10.1111/imr.12511

42. Hessel AJ, Hangartner L, Hunter M, Havenith CE, Beurskens FJ, Bakker JM, et al. Fc receptor but not complement binding is important in antibody

43. Bournazos S, Klein F, Pietzsch J, Seaman MS, Nussenzeig MC, Ravetch JV. Broadly neutralizing anti-HIV-1 antibodies require Fc effector functions for in vivo activity. Cell (2014) 158:1243–53. doi: 10.1016/j.cell.2014.08.023
54. Xiao P, Zhao J, Patterson LJ, Brocca-Cofano E, Venzon D, Kozlowski PA, et al. Multiple vaccine-elicited nonneutralizing antienvvelope antibody activities contribute to protective efficacy by reducing both acute and chronic viremia following simian/human immunodeficiency virus SHIV89.6P challenge in rhesus macaques. *J Virol* (2010) 84:7161–73. doi: 10.1128/JVI.00410-10
55. Seder RA, Darrah PA, Roederer M. T-cell quality in memory and protection: implications for vaccine design. *Nat Rev Immunol* (2008) 8:247–58. doi: 10.1038/nri2274
56. Peretz Y, Ndongala ML, Boullet S, Boullass MR, Rouleau D, Cote P, et al. Functional T cell subsets contribute differentially to HIV peptide-specific responses within infected individuals: correlation of these functional T cell subsets with markers of disease progression. *Clin Immunol* (2007) 124:57–68. doi: 10.1016/j.clim.2007.04.004
57. Pereyra F, Addo MM, Korber B, Goonetilleke N, Liu MK, Teruya-Feldstein J, et al. Genetic and immunologic heterogeneity among persons who control HIV infection in the absence of therapy. *J Infect Dis* (2008) 197:563–71. doi: 10.1086/526786
58. Riou C, Burgers WA, Mlisana K, Koup RA, Roederer M, Abdool Karim SS, et al. Differential impact of magnitude, polyfunctional capacity, and specificity of HIV-specific CD8+ T cell responses on HIV set point. *J Virol* (2014) 88:1819–24. doi: 10.1128/JVI.02968-13
59. Betts MR, Nason MC, West SM, De Rosa SC, Migueles SA, Abraham J, et al. HIV nonprogressors preferentially maintain highly functional HIV-specific CD8+ T cells. *Blood* (2006) 107:4781–9. doi: 10.1182/blood-2005-12-4818
60. Streeck H, Brumme ZL, Kardani K, Bolhassani A, Shahbazi S. Prime-boost vaccine strategy against HIV/SIV infection: strategies and challenges for vaccine design. *Int Rev Immunol* (2012) 31:455–65. doi: 10.1080/08830180802684331
61. Siegrist CA. Vaccine immunology (General Aspects of Vaccination). In: Plotkin SA, Ollier WE (eds). *Vaccines*. 7th ed. New York: Saunders Elsevier; (2018) p 16–34.e7. doi: 10.1016/C2013-0-18914-3
62. Shiver JW, Fu TM, Chen L, Casimiro DR, Davies ME, Evans RK, et al. Replication-competent adenoviral vaccine vector elicits effective anti-immunodeficiency-virus immunity. *Nature* (2002) 415:331–5. doi: 10.1038/415331a
63. Amara RR, Villinger F, Altman JD, Lydy SL, O’Neil SP, Staprans SI, et al. Control of a mucosal challenge and prevention of AIDS by a multiprotein DNA/MVA vaccine. *Science* (2001) 292:69–74. doi: 10.1126/science.1058915
64. Bomsel M, Tudor D, Drillet AS, Alfsen A, Ganor Y, Roger MG, et al. Immunization with HIV-1 gp41 subunit virosomes induces mucosal antibodies protecting nonhuman primates against vaginal SHIV challenges. *Immunology* (2011) 134:269–80. doi: 10.1016/j.immuni.2011.01.015
65. Demberg T, Robert-Guroff M. Mucosal immunity and protection against HIV/SIV infection: strategies and challenges for vaccine design. *Int Rev Immunol* (2012) 29:20–48. doi: 10.1080/08830180802684331
66. Lycke N. Recent progress in mucosal vaccine development: potential and limitations. *Nat Rev Immunol* (2012) 12:492–505. doi: 10.1038/nri3251

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Xiao, Dienger-Stambaugh, Chen, Wei, Phan, Beavis, Singh, Adhikary, Tiwari, Villinger, He and Spearman. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.