RESEARCH ARTICLE

Immunological Characterization of Plant-Based HIV-1 Gag/Dgp41 Virus-Like Particles

Sarah A. Kessans1,2, Mark D. Linhart1,2, Lydia R. Meador1,2, Jacquelyn Kilbourne2, Brenda G. Hogue1,2,3, Petra Fromme3,4, Nobuyuki Matoba1,2, Tsafrir S. Mor1,2*

1 School of Life Sciences, Arizona State University, Tempe, Arizona, United States of America, 2 Center for Infectious Diseases and Vaccinology, The Biodesign Institute, Arizona State University, Tempe, Arizona, United States of America, 3 Center for Applied Structural Discovery, The Biodesign Institute, Arizona State University, Tempe, Arizona, United States of America, 4 Department of Chemistry and Biochemistry, Arizona State University, Tempe, Arizona, United States of America

¤ Current address: Biomolecular Interaction Centre, Department of Chemistry, University of Canterbury, Christchurch, New Zealand
¤ Current address: Owensboro Cancer Research Program, Owensboro, Kentucky, United States of America, and James Graham Brown Cancer Center and Department of Pharmacology & Toxicology, University of Louisville School of Medicine, Louisville, Kentucky, United States of America
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Abstract

It is widely anticipated that a prophylactic vaccine may be needed to control the HIV/AIDS epidemic worldwide. Despite over two decades of research, a vaccine against HIV-1 remains elusive, although a recent clinical trial has shown promising results. Recent studies have focused on highly conserved domains within HIV-1 such as the membrane proximal external region (MPER) of the envelope glycoprotein, gp41. MPER has been shown to play critical roles in mucosal transmission of HIV-1, though this peptide is poorly immunogenic on its own. Here we provide evidence that plant-produced HIV-1 enveloped virus-like particles (VLPs) consisting of Gag and a deconstructed form of gp41 comprising the MPER, transmembrane, and cytoplasmic domains (Dgp41) provides an effective platform to display MPER for use as an HIV vaccine candidate. Prime-boost strategies combining systemic and mucosal priming with systemic boosting using two different vaccine candidates (VLPs and CTB-MPR—a fusion of MPER and the B-subunit of cholera toxin) were investigated in BALB/c mice. Serum antibody responses against both the Gag and gp41 antigens were elicited when systemically primed with VLPs. These responses could be recalled following systemic boosting with VLPs. In addition, mucosal priming with VLPs allowed for a boosting response against Gag and gp41 when boosted with either candidate. Importantly, the VLPs also induced Gag-specific CD4 and CD8 T-cell responses. This report on the immunogenicity of plant-based Gag/Dgp41 VLPs may represent an important milestone on the road towards a broadly efficacious and inexpensive subunit vaccine against HIV-1.
Introduction

The HIV-1 transmembrane subunit of the envelope protein (Env), gp41, contains the highly conserved membrane proximal external region, located just outside the lipid viral envelope (MPER, amino acids 661–683, [1]). The gp41 domain that encompasses the MPER and extends toward the C-terminal heptad repeat (residues 649–684, sometimes denoted as “MPR” but for simplicity we will refer to both as MPER, [2]) functions as a galactosyl-ceramide-binding lectin and is critical for mediating viral transcytosis across mucosal membranes [3] and other mucosal transmission routes [4, 5]. Both mucosal and systemic antibodies (Abs) raised against immunogens containing the MPER can block the transcytosis of HIV across the epithelial barrier [6, 7], similar to naturally occurring polyclonal mucosal IgAs found in the mucosal secretions of some highly exposed persistently seronegative (HEPS) individuals [8–11]. Revealingly, broadly neutralizing human monoclonal Abs (mAbs) such as 2F5, 4E10 and 10E8 also target this region [12–15]. The MPER, therefore, provides an important target for vaccine design, in addition to the widely-explored but highly-mutable surface subunit of Env (gp120, [16–19], reviewed in [20–22]).

The proximity of the MPER to the viral envelope is increasingly recognized as a major factor in the antigenicity and immunogenicity of the domain [23–25], suggesting that the presentation of the MPER in the context of a membrane, e.g. in virus-like particles (VLPs) may be of value. This notion and the recent success of prophylactic VLP-based vaccines such as those aimed at human papillomaviruses [26] provide the motivation for VLP-based vaccines against HIV-1. Gag, a polyprotein that gives rise to the main structural proteins of HIV-1, is both necessary and sufficient for the formation of enveloped VLPs [27, 28]. Gag contains the highest density of cytotoxic T-lymphocyte (CTL) epitopes of any HIV protein [29] and Gag-based VLPs are capable of inducing strong CTL responses without adjuvant [30]. CD8 T cell responses to Gag have been correlated with control of viral replication in infected individuals [31]. In addition, Gag VLPs can display HIV Env proteins on their surface in their native conformation [32], and these VLPs have been shown to induce both Env- and Gag-specific Abs and CTLs [33], making Gag VLPs attractive candidates as an HIV vaccine platform [34].

Plant-based production systems for biologics and vaccines lately reached several critical milestones gaining FDA approval for large-scale clinical trials and commercialization [35–39]. We previously reported that Gag VLPs displaying a deconstructed form of gp41 (Dgp41, comprising MPER, transmembrane, and cytoplasmic domains) could be produced in Nicotiana benthamiana plants (Fig 1) [40]. Here we report on immunization studies employing plant-based HIV-1 Gag/Dgp41 VLPs and demonstrate their immunogenicity.

Materials and Methods

Immunogens’ preparation

Highly-enriched plant-based VLPs were prepared by transiently expressing Dgp41 in Gag-expressing transgenic N. benthamiana plants and transiently expressing Dgp41 as previously described (see Fig 1 for expression and purification strategy) [40]. Quantitative immunoblots were use to quantify Gag and Dgp41 as previously described [40].

CTB-MPR is a fusion protein consisting of the HIV-1’s MPER fused to the carboxy-terminus of cholera toxin B-subunit (CTB, [41]). Expression of CTB-MPR in Escherichia coli and its purification previously described [41]. CTB-MPR preparation quality and yield were determined by Coomassie stained gels, quantitative immunoblots (using a pure standard), and the absorbance at 280 nm using $\varepsilon = 2.1 \text{ mM}^{-1} \text{ cm}^{-1}$ [41]. Total protein was determined as previously described [42].
Animal Care and Use

This study was reviewed and approved by the Arizona State University Institutional Animal Care and Use Committee (IACUC) under protocol number 11-1174R.

**Housing and husbandry.** All animals were housed in accordance with the American Association for Laboratory Animal Care (AALAC) standards. The animals are housed in Thoren ventilated racks that are HEPA filtered on both the supply and exhaust air on Irradiated Sani-Chip 7990. Environmental enrichment includes social housing and nestlets to encourage nesting activities. Animals are provided unlimited access to food and water. They are handled in accordance with the Animal Welfare Act and Institutional Animal Care and Use Committee.
(IACUC) regulations. Experiments involving animals were conducted in a facility fully accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International (Unit #000765) and an assurance is on file with the Office for Laboratory Animal Welfare (#A3217-01). Experiments were planned and conducted utilizing the three R’s (reduce, replace and refine), which included environmental enrichment, veterinary oversight, numbers reflecting statistical significance and the use of appropriate analgesics and anesthesia when appropriate.

**Animal Monitoring.** All animals are observed daily by trained DACT personnel for signs of illness or abnormal behavior by 10 AM. Training is provided by the Department of Animal Care and Technologies (DACT) veterinarians, experienced DACT personnel (i.e., Lead Technologists, Supervisors). Personnel performing the daily observations report sick or injured animals to the DACT vet team. During regular work hours, personnel can contact the veterinary team in person, by phone, or via email using a veterinary team distribution list. Outside of working hours, a veterinary team member carries a dedicated on-call cell phone. A call list with contact numbers for supervisory and veterinary personnel is posted in each animal facility.

The mice are monitored for activity level that may be indicative of illness such as hypoactivity (abnormally low), hyperactivity (abnormally high), lethargy, restlessness. They are monitored for behavioral signs such as vocalization, self-trauma, aggressiveness, isolation from cage mates, or ataxia. They are monitored for changes to their appearance such as unkempt or greasy fur, porphyrin staining around eyes and nostrils, hunched posture, pale mucous membranes, pale paws, soiled anogenital area, labored breathing, weight loss, dehydration or diarrhea. Animals that show obvious signs of illness are either removed from the study and treated if it is relevant, or euthanized immediately, based upon the recommendation of the veterinary team.

**Euthanasia.** Mice used in this study were euthanized by CO2 asphyxiation, which is consistent with the most recent recommendations of the American Veterinary Medical Association [43]. Cervical dislocation or secondary thoracotomy was used as a subsequent secondary measure. No animals died during the experiments. One mouse belonging to the control group (see below) was found to be in lateral recumbency and experiencing labored breathing. Since this was the day before the final bleeding, the mouse was euthanized immediately. Gross necropsy was performed and revealed slightly enlarged kidneys, gas in the gastrointestinal tract (indicative of not eating), and dehydration. This was not deemed as a result of the study as the mouse was in our control group. This was one day before scheduled termination so there was no significant impact on our study.

**Systemic immunization**

The immunization samples was prepared by diluting the concentrated protein preparations into PBS supplemented with Ribi Adjuvant (Sigma-Aldrich, final concentration of 2% oil as per manufacturer’s instructions). Female BALB/c mice (6-wk old, Charles River) were immunized intraperitoneally (i.p.) with 200 μL of the immunization sample containing either CTB-MPR (3.5 μg ≡ 1.2 μg MPER ≡ 0.2 nmol MPER), VLP preparation (111 μg total protein containing 4.8 μg dgp41 and 4.8 μg Gag, equivalent to 0.2 nmol and 0.1 nmol, respectively), or a negative control sample (100 μg total protein, mock-purified in the same manner as the VLP proteins). Four experimental groups (n = 8) were given either VLP or CTB-MPR during each of three priming immunizations, and were then given either VLP or CTB-MPR during each of two boosting immunizations (Fig 2A). A fifth group was immunized with the negative control sample under the same regimen. Retro-orbital vein blood samples were taken as indicated. Serum was prepared following clotting and centrifugation and kept at −80°C until further use.
For intranasal (i.n.) administration, immunization cocktails were made by mixing the indicated proteins with PBS. Each dose (20 μl total, 10 μl per nostril) contained either CTB-MPR (35 μg/17 μg MPER) with cholera toxin (CT, 1 μg, List Biological Laboratories),
VLP preparation (1.1 mg total protein containing 48 μg dgp41 and 48 μg Gag, equivalent to 2 nmol and 1 nmol, respectively) with either CT (1 μg), or murabutide (200 μg, InvivoGen). Female BALB/c mice (6-wk old, Charles River, n = 8 per group) were given four i.n. priming immunizations (Fig 2B). Group 1 received VLPs+murabutide, Groups 2 and 3 received VLPs +CT, Groups 4 and 5 received CTB-MPR+CT. Group 6, serving as a control, was immunized i.p. with VLPs, as described above. Mice in all groups were given two i.p. boosting immunizations with either VLPs (4.8 μg dgp41 and 4.8 μg Gag, Groups 1, 2, 4 and 6) or CTB-MPR (3.5 μg), Groups 3 and 5, Fig 2B), administered with Ribi Adjuvant as per the systemic trial above. Serum, vaginal secretions, and fecal pellets were collected from all mice as indicated. Abs were extracted from fecal pellets by soaking five pellets (~50 mg) in PBS containing 0.02% Na-azide (500 μL) for 30 min at 4°C with occasional vortex and clarification by centrifugation (14,000 ×g, 10 min). Vaginal secretions were collected by lavage using PBS (100 μL) with a blunt-tipped syringe needle. Serum, fecal Abs, and vaginal lavages were kept at -80°C.

Antibody Titer Assays
ELISA plates were coated with 20 μg of streptavidin (Sigma-Aldrich) and 2 μg of biotinylated MPR peptide (for detection of anti-MPER Abs) or 1 μg of p24-CTA2 (for detection of anti-Gag Abs). Binding of the MPR peptide through its N-terminal biotin should maximize accessibility the accessibility of the peptide to interactions with cognate antibodies [41]. The wells were overlaid with a threefold serial dilution of serum, vaginal secretions or fecal samples (starting with 1:50, 1:5 or 1:2, respectively) in PBS containing 0.5% Tween 20 and 5% dry milk and the procedure was continued as previously described [41]. Endpoint titers were determined as the reciprocal of the dilution factor of sample giving background levels of OD490. Statistical analysis of data was by the Kruskal-Wallis test followed by Dunn’s Multiple Comparison test.

IFN-γ ELISPOT Assay
Splenocytes were prepared from pooled harvested spleens on Week 12. Interferon-gamma (IFN-γ) Enzyme-Linked Immunosorbent Spot Assay (ELISPOT) responses were measured using a mouse IFN-γ set (BD Biosciences). Threefold serially diluted triplicates of splenocytes (starting at 1 x 10⁶ splenocytes/well) were applied to the plates in a final volume of 200 μl RPMI 1640 culture medium (with 10% heat-inactivated fetal bovine serum, 100 U/mL penicillin, 100 μg streptomycin). The peptides AAMQMLKDTINEEAA (corresponding to the GagCD8 epitope, from HIV-1 Consensus C Gag (15-mer) Peptides, Cat#8118, NIH AIDS Reagent Program) and SNPPVPVDYKWRW/VPVDYKWRWILGL (corresponding to the GagCD4 epitope, from HIV-1 Consensus C Gag (15-mer) Peptides, Cat#8118, NIH AIDS Reagent Program) were used as stimuli in the assay at 5 μg/mL. Reactions without peptide served as background controls. Reactions were allowed to proceed for 30 h at 37°C in a humidified 5% CO₂ atmosphere. Spots were detected with the detection antibody, developed with 3-amino-9-ethyl-carbazole (BD Biosciences), and analyzed using a CTL ImmunoSpot plate reader and counting software (Cellular Technology Ltd). For each group of mice, the number of background spots in the absence of peptide was subtracted from the average of the triplicate values in order to determine the number of peptide-relevant spots.

Results
Serum anti-Gag responses to systemic immunizations
Anti-p24 Abs were quantified after the third priming immunization (Week 6), before the first boost (Week 8), after the first boost (Week 10), and after the second boost (Week 12, Fig 3). In
Fig 3. Serum anti-p24 (anti-Gag) responses. (A) Mice were immunized as per Fig 2A. Serum samples were diluted 1:50 and IgGs were detected by direct ELISA on indicated weeks. Shown are average net Immunogenicity of Plant-Based HIV-1 VLPs
all mice (16/16) primed with VLPs, significant Ab titers (p < 0.01, in comparison to the negative control group) were elicited after priming and remained steady until the first boost (Fig 3B–3E). The VLP-primed animals were then split into two groups and boosted with either CTB-MPR or VLPs. CTB-MPR-boosted mice retained significant titers of anti-p24 Abs through the end of the trial, despite the fact that the mice did not receive any further Gag protein. Boosting with VLPs resulted in increased antibody titers, with final antibody titers reaching extremely significant (p < 0.001) values over the negative control mice. The majority of mice (5/8) primed with CTB-MPR and boosted with VLPs developed anti-p24 Ab responses at similar titers to VLP-primed animals with two doses. All of these data demonstrate that plant-produced VLPs can elicit robust, long-lived Ab responses against the Gag protein, which are detectable at high titers at least eight weeks after the final VLP immunization (our final time point) for VLP/CTB-MPR-immunized animals.

**Serum anti-MPER responses to systemic immunizations**

Most (11/16) mice primed with CTB-MPR responded to the MPER moiety before the second boost (Fig 4). One of these mice (#3 from the CTB-MPR-primed and boosted group) responded remarkably better than the rest of the mice (shown on its own, Fig 4). VLP priming induced detectable anti-MPER Abs in 7/16 mice. All animals boosted with CTB-MPR, regardless of priming group, displayed significant anti-MPER antibody titers (p < 0.001 and p < 0.01, respectively) after the second boost. In 7/8 mice primed with VLPs and boosted with VLPs, significant (p < 0.05) titers were elicited after the first boost, which steadily increased following the second boost. All four experimental groups elicited statistically significant Ab levels following the second boost. However, CTB-MPR appears to be a slightly more potent inducer of anti-MPER Abs, though this difference was not statistically significant.

**Serum anti-Gag antibody response to mucosal immunization**

Because HIV-1’s main mode of transmission mode is the crossing of mucosal barriers in the female genital tract and the lower gastro-intestinal tract, we further tested the ability of plant-derived VLPs to stimulate mucosal responses. To this end, immunogens were used to test the effectiveness of intranasal priming with systemic boosting of plant-produced VLPs, CTB-MPR, or a combination of both.

Anti-p24 Abs were not detected in any of the mucosally-primed mice prior to boosting (Fig 5), although systemically-primed mice displayed anti-p24 serum IgG titers consistent with the first immunization trial (see above) with 6/8 mice responding to Gag following the first priming immunization. The majority of mice (30/32) in all groups that were i.p.-boosted with VLPs responded to the Gag protein by eliciting very high Ab titers (Fig 5). These results confirm the effectiveness of using VLPs in a systemic prime/boost regimen to elicit a response against Gag, and also suggest the effectiveness of mucosal priming with systemic boosting to elicit a similar response.

**Serum anti-MPER antibody response to mucosal immunization**

As was the case of the anti-p24 response, anti-MPR serum Abs were below the detection limit in all mucosally-primed mice prior to boosting immunizations (Fig 5). In accordance with the
Fig 4. Serum anti-MPER IgG levels. (A) Mice were immunized as per Fig 2A. Serum samples were diluted 1:50 and IgGs were detected by direct ELISA on indicated weeks. Shown are average net OD490 values.
(mean +/- SEM). Mouse #3 (from CTBMPR-CTBMPR group) responded significantly higher than other mice from this group and was considered an outlier, and antibody response from this mouse is shown by itself. Also shown are Ab endpoint titers at response peaks after priming (B, Week 6); before first boost (C, Week 8), after first boost (D, Week 10), and after second boost (E, Week 12). Symbols indicate statistical significance as compared to week zero within the group evaluated by Kruskal-Wallis test and Dunn’s Multiple Comparison test: * p < 0.05, ** p < 0.01, *** p < 0.001.

Fig 5. Serum IgG and mucosal IgA responses across all weeks of mucosal immunization. Anti-Gag (A, C, E) and Anti MPER (B, D, F) serum IgG (A and B), fecal mucosal IgA (C and D), and vaginal mucosal IgA (E and F) were assayed in samples obtained on weeks 0, 1, 3, 5, 7, 8, 10, and 12. Time points are averages of all mice in each group. Symbols indicate statistical significance as compared to week zero within the group evaluated by Kruskal-Wallis test and Dunn’s Multiple Comparison test: * p < 0.05, ** p < 0.01, *** p < 0.001.
systemic immunization experiment, no significant Ab titers against MPER were raised in the systemically-primed mice prior to boosting, either. Systemic boosting with CTB-MPR after mucosal priming with the same immunogen (Group 5) elicited significant ($p < 0.001$, Fig 6) levels of Abs as compared to naïve mice, consistent with previous studies [6]. Interestingly, mucosal priming with VLPs and boosting with either VLPs or CTB-MPR (Groups 1, 2, and 3) elicited only marginal levels of Abs, and the same was observed for mucosal priming with CTB-MPR and boosting with VLPs. Although all the mice responded in Group 6 (systemically primed and boosted with VLPs), the levels of Abs were lower in this group than in the CTB-MPR primed and boosted group (Group 5). However, we note that the differences between the two groups were not statistically significant.

**Fecal anti-Gag and anti-MPER responses to mucosal immunizations**

Mucosal response against Gag was limited in all groups either primed or boosted with VLPs, although 6/8 mice were positive for IgAs in fecal samples in Group 6 (systemically primed and boosted with VLPs). Responses from Groups 1, 2, and 3 (mucosally primed with VLPs) were all statistically similar. Additionally, mucosal VLP priming induced slightly higher Ab titers with a greater number of responders as compared to those primed with CTB-MPR (Groups 4 and 5, Fig 6). While the difference was not statistically significant, the trend suggests that mucosal priming with VLPs might assist in eliciting a mucosal response against Gag, though this response is not as strong as that elicited with systemic priming with VLPs.

Similar to the anti-p24 mucosal response, we could not detect any fecal anti-MPER IgAs in any of the treatment groups until after the final immunization. Two weeks after the final boost, low anti-MPER IgA responses could be detected in 12/48 mice. The two responders in Group 5 had two of the highest responses, but no group reached statistically significant responses in comparison to naïve mice (Figs 4 and 5). Mice in Groups 2, 3, and 4 had similar responses. Group 1 had a single responder while all mice in Group 6 had fecal anti-MPER responses below the limit of detection. The results of MPER response in fecal samples are inconclusive, but suggest that mice mucosally primed and systemically boosted with CTB-MPR can elicit a moderate IgA response in fecal samples, while systemic priming and boosting with VLPs is largely ineffective in eliciting a mucosal anti-MPER response in mice.

**Vaginal anti-Gag and anti-MPER response in mucosal immunizations**

Anti-p24 IgA Abs could not be detected in any vaginal secretions of the mice prior to boosting immunizations, and only three mice (one from Group 2 and two from Group 6) responded two weeks after the second boost.

In accordance with low levels of anti-MPER Abs in fecal samples, only 12/48 mice from all groups responded to MPER in vaginal secretions. Once again, two mice in Group 5 (the same mice with higher anti-MPER response in fecal samples) had two of the highest overall responses, while vaginal anti-MPER responses from other groups were minimal or undetectable. Consistent with fecal results, no anti-MPER Abs were detected in vaginal secretions from mice in Group 6. Overall, the results of anti-MPER response in vaginal secretions are congruent with results of anti-MPER response in fecal samples. These results suggest that while systemic priming and boosting with VLPs can be successful in eliciting antibody responses against Gag in serum and at mucosal sites, mucosal VLP priming may not be nearly as effective in this case, with minimal responses seen to Dgp41. Mucosal immunization elicits the highest antibody responses to gp41 at all sites in Group 5, which is primed and boosted with CTB-MPR.
Fig 6. Endpoint titers. Post-boosting (Week 10) antibody endpoint titers against Gag (A, C, E) and MPER (B, D, F) in serum (IgG, A and B), fecal (secretory IgA, C and D), and vaginal (secretory IgA, E and F) samples. Symbols indicate statistical significance as compared to week zero within the group evaluated by Kruskal-Wallis test and Dunn’s Multiple Comparison test: * p < 0.05, ** p < 0.01, *** p < 0.001.

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IFN-γ ELISPOT Assay

In addition to the humoral and mucosal immune responses, the cellular responses against peptides corresponding to Gag-specific CD4 and CD8 immunodominant epitopes were also assayed two weeks following the final immunization for the vaccination experiment described in Fig 3. Mice primed (either mucosally or systemically) and boosted with Gag/Dgp41 VLPs had expression of IFN-γ at levels over 40 sfu/10^6 cells in response to the CD8 epitope, suggesting a high CTL response against Gag in these groups (Fig 7). Overall, CD4 responses were lower with the highest response being mice primed and boosted systemically with VLPs at over 20 sfu per million splenocytes (Group 6, Fig 7). Minimal, but detectable, CD4 responses were seen with all groups primed with VLPs regardless of boosting regimen (Groups 1–3, Fig 6). In general, animals primed and boosted with VLPs induced higher T cell responses than those with mixed regimens (compare groups 2 and 6 to 1, 3, and 4). As expected, mice primed and boosted with CTB-MPR did not respond to either peptide (Group 5, Fig 7). These results suggest that both CD8 and CD4 T-cell responses against Gag were elicited with the VLPs with the most successful regimen being systemic administration.

Discussion

The MPER domain of gp41 in HIV-1, which plays multiple roles in early stages of the viral lifecycle, is targeted by Abs with strong, broad and diverse antiviral activities including transcytosis blockade (e.g. [3, 7, 9, 41, 44]), neutralization (e.g. [8, 15, 45, 46]), and Ab-dependent cell cytotoxicity [47]. Based on these findings, the MPER deservedly became the focus of intensifying vaccine development research.

Because the MPER is poorly immunogenic on its own, a platform with which to present the peptide is required and various carrier proteins have been tested (e.g. [48, 49]). Our lab has tested the immunogenicity of multiple MPER fusion proteins which utilized CTB [6, 7, 41], *Yersinia pestis* antigens [50], and hepatitis B core antigen (Cherni, Matoba, and Mor, unpublished). These studies demonstrated the success of heterologous prime-boost strategies with multiple routes of immunization and more than one vaccine construct in eliciting transcytosis-blocking Abs. However, an optimal presentation platform to display the MPER peptide that
would ensure a uniformly strong immune response against MPER, while overcoming the immune-dominance of the carrier proteins, is still elusive.

The ability of the MPER to elicit Abs with antiviral activities is context dependent. In particular, the proximity to the membrane plays a significant role. For example, recently it was demonstrated using a knock-in mouse model that 2F5 Abs could be elicited to very high titers \textit{in vivo} only when MPER was displayed in the context of a membrane [51]. Similarly, the MPER peptide (“P1”) attached to virosomes was shown to elicit transcytosis-blocking Abs in both serum and mucosal sites in macaques (where their elicitation correlated with protection from infection [52]) and humans [19]. Within the context of an intact Env, the MPER is available to bind neutralizing mAbs exclusively during the pre-hairpin intermediate stage of the infection process. Outside of this window, binding is hindered because the MPER is either sterically-obscured (e.g., on the surface of a virion), or strongly interacting with other gp41 domains [53]. These conformational considerations, among others (such as clonal anergy, [54]) contribute to the low propensity for induction of 2F5-like Abs during the acute phase of HIV-1 infection [53].

We previously demonstrated that plants can efficiently make HIV-1 enveloped VLPs consisting of p55$^{\text{GAG}}$ and a membrane-anchored stripped-down Env that we call “deconstructed gp41” [40]. Devoid of the immunodominant gp120 and the N-terminal regions of gp41, we hypothesized that such VLPs would present the MPER in a conformation that approximate the pre-fusion intermediate stage [2]. Interestingly, recent structural analyses of a similar gp41 construct containing the MPER and the transmembrane domains of gp41 (MPR-TM) reveal that indeed, the neutralizing epitopes of 4E10 and 2F5 are accessible to these monoclonal antibodies when the protein is presented in a non-denatured trimeric state [2, 55]. The antibodies bind to the MPR-TM with sub-nanomolar affinities as determined by competitive ELISA and surface plasmon resonance [2, 55].

The plant-derived VLPs incorporate the Gag protein and the deconstructed envelope protein Dgp41 at Gag:Dgp41 ratios that vary slightly between 1:1 [40] and 2:1 (here). Previous studies by others aimed at producing recombinant VLPs reported lower levels of Env protein incorporation. For example, Hammonds and co-workers reported much higher Gag:Env ratios in VLPs produced in human cell lines (varying between 25:1 to 50:1) or through baculovirus infection of insect cells (10:1) [56]. The presence of more dgp41 molecules on the surface of plant-derived VLPs as compared to other recombinant VLPs is immunologically advantageous.

Here we tested the hypothesis that such plant-derived VLPs could serve to elicit immune responses against the presented antigens. As shown above, HIV-1 Gag/Dgp41 VLPs produced in \textit{N. benthamiana} were able to elicit substantial titers of Abs against both Gag and gp41 epitopes through a heterologous platform prime-boost regimens in conjunction with CTB-MPR. Intraperitoneally-administered VLPs were also effective as immunogens on their own (Figs 2–5) and significant levels of Abs were raised against the MPER after boosting with VLPs. Our ELISA detection system is not likely to enable us to detect potential conformational Abs and therefore potentially underestimates the humoral response to the VLPs. Although mucosal priming immunizations were not sufficient to elicit Abs against Gag or gp41, such responses were shown for animals that subsequently were boosted with systemically-delivered plant-produced VLPs (Fig 6). In conjunction with the VLP induction of strong Ab responses, we detected Gag-specific CD4 and CD8 T-cell responses, which were the highest in groups primed and boosted with VLPs regardless of immunization route (Fig 7).

The use of plants as production platforms for candidate subunit vaccines is now well established, achieving important milestones in the last couple of years including winning FDA approval for use in humans [39, 57]. As was recently demonstrated by stepping up plant production of a promising anti-Ebola virus mAb cocktail [39, 58, 59], plants offer a highly competitive alternative in terms of costs, scale-up and safety to traditional production of vaccines such
as fermentation (yeast and animal cell cultures), or eggs. In particular, plants were shown to express VLPs such as those of Hepatitis B virus [60, 61], Norwalk virus [62, 63], influenza [64], and even HIV-1 [65, 66].

Only few studies were published presenting immunogenicity data for plant-based HIV-1 full-length Gag or membrane-anchored gp41. Our co-expression system [40] was the first to produce enough full-length p55Gag together with Dgp41 to conduct a study like the one we describe here (Gag yield is ~ 22 mg/kg fresh leaf material). However, plant-expressed derivatives of Gag (i.e. p24, p17, and p41) were tested and were shown to elicit anti-Gag Abs as well as CTL responses [65, 67–70]. In addition, several studies demonstrated success in eliciting immunological responses against plant-based MPER constructs [71–75], however none were membrane anchored. While we acknowledge that further investigation is necessary to elicit a more robust and durable anti-MPER Ab response and demonstrate its protective efficacy in HIV-1 challenge models, our results here offer evidence that plant-based Gag/Dgp41 VLPs can induce relevant Ab and T cell responses and forms the basis for the development of an effective component for future HIV-1 prime/boost immunization trials.

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Author Contributions

Conceived and designed the experiments: SAK NM TSM. Performed the experiments: SAK MDL LRM JK. Analyzed the data: SAK LRM BGH PF NM TSM. Wrote the paper: SAK LRM BGH PF NM TSM.

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