Full title: *Mamu-B*17+ rhesus macaques vaccinated with *env*, *vif*, and *nef* manifest early control of SIVmac239 replication

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Short title: **Early control of SIV in env-vaccinated B*17+ macaques**

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Certain major histocompatibility complex class-I (MHC-I) alleles are associated with spontaneous control of viral replication in human immunodeficiency virus (HIV)-infected people and simian immunodeficiency virus (SIV)-infected rhesus macaques (RM). These cases of “elite” control of HIV/SIV replication are often immune-mediated, thereby providing a framework for studying anti-lentiviral immunity. Here we examined how vaccination impacts SIV replication in RM expressing the MHC-I allele Mamu-B*17. Approximately 21% of Mamu-B*17+ and 50% of Mamu-B*08+ RMS control chronic phase viremia after SIVmac239 infection. Because CD8+ T-cells targeting Mamu-B*08-restricted SIV epitopes have been implicated in virologic suppression in Mamu-B*08+ RMS, we investigated whether this might also be true for Mamu-B*17+ RMS. Two groups of Mamu-B*17+ RMS were vaccinated with genes encoding Mamu-B*17-restricted epitopes in Vif and Nef. These genes were delivered by themselves (Group 1) or together with env (Group 2). Group 3 included MHC-I-matched RMS and served as the control group. Surprisingly, the Group 1 vaccine regimen had little effect on viral replication compared to Group 3, suggesting that, unlike Mamu-B*08+ RMS, pre-existing SIV-specific CD8+ T-cells alone do not facilitate long term virologic suppression in Mamu-B*17+ RMS. Remarkably, however, 5/8 Group 2 vaccinees controlled viremia to <15 viral RNA copies/mL soon after infection. No serological neutralizing activity against SIVmac239 was detected in Group 2, although vaccine-elicited gp140-binding antibodies correlated inversely with nadir viral loads. Collectively, these data shed new light into the unique mechanism of elite control in Mamu-B*17+ RMS and implicate vaccine-induced, non-neutralizing anti-Env antibodies in the containment of immunodeficiency virus infection.
A better understanding of the immune correlates of protection against HIV might facilitate the development of a prophylactic vaccine. Therefore, we investigated simian immunodeficiency virus (SIV) infection outcomes in rhesus macaques expressing the major histocompatibility complex class I allele *Mamu*-B*17*. Approximately 21% of *Mamu*-B*17*+ macaques spontaneously control chronic phase viremia after SIV infection, an effect that may involve CD8+ T-cells targeting Mamu-B*17*-restricted SIV epitopes. We vaccinated *Mamu*-B*17*+ macaques with genes encoding immunodominant epitopes in Vif and Nef alone (Group 1) or together with env (Group 2). Although neither vaccine regimen prevented SIV infection, 5/8 Group 2 vaccinees controlled viremia to below detection limits shortly after infection. This outcome, which was not observed in Group 1, was associated with vaccine-induced, non-neutralizing Env-binding antibodies. Together, these findings suggest a limited contribution of Vif- and Nef-specific CD8+ T-cells for virologic control in *Mamu*-B*17*+ macaques and implicate anti-Env antibodies in containment of SIV infection.
Introduction

Despite improvements in prevention strategies and antiretroviral therapy (ART) coverage, thousands of new human immunodeficiency virus (HIV) infections are still occurring every day, highlighting the need for an effective HIV vaccine (1). Eliciting robust protection against HIV infection has not been straightforward, as seen by the failure of most HIV vaccines tested in humans to date (2–6). Although the RV144 trial remains the only report of vaccine-mediated reduction in HIV infection rates (7), the observed results were modest, short-lived, and continue to be contested (8, 9). The unsatisfactory performance of mainstream HIV vaccine regimens underscores the need to better understand the nature of effective anti-lentiviral immune responses.

Elite controllers (ECs) are a small fraction of HIV-infected individuals who spontaneously control chronic phase viremia in the absence of ART (10). Certain major histocompatibility complex class I (MHC-I) alleles, such as HLA-B*27 and HLA-B*57, are associated with elite control of HIV-1 infection (11), implying an immunological basis for this phenotype. Indeed, CD8+ T-cells targeting viral epitopes restricted by “protective” MHC-I molecules and natural killer (NK) cells are widely thought to be important mediators of antiviral activity in ECs (12, 13). The study of ECs thus provides a useful framework to investigate the basis for immune containment of lentivirus replication. Similar to human ECs, certain rhesus macaque (RM) MHC-I alleles are also associated with elite control of SIV infection. Indeed, approximately 21% of unvaccinated RMs expressing Mamu-B*17 control chronic phase viral replication after infection with SIVmac239 (14). The incidence of elite control in Mamu-B*08+ RMs is higher, reaching 50% of infected animals (15). Curiously, the peptide binding motifs of the Mamu-B*08 and
Mamu-B*17 molecules resemble those of HLA-B*27 and HLA-B*57, respectively (16, 17). This similarity is not explained by sequence homology between the human and RM MHC-I alleles, thereby implicating the presented peptide as an important determinant of elite control.

The immunodominant SIV epitopes restricted by Mamu-B*08 and Mamu-B*17 share a common feature, that is, their location in the accessory proteins Vif and Nef (18–20). While CD8+ T-cell responses targeting these Vif and Nef epitopes are crucial for virologic control in Mamu-B*08+ RMs (21), it is not clear to what extent Vif- and Nef-specific CD8+ T-cells contribute to the EC phenotype of Mamu-B*17+ RMs. We set out to clarify this issue by conducting an SIV vaccine trial in Mamu-B*17+ RMs. We hypothesized that vaccine-induced CD8+ T-cells targeting the immunodominant Mamu-B*17-restricted Vif HW8 (amino acids 66-73) and Nef IW9 (amino acids 165-173) epitopes would increase the incidence of elite control in Mamu-B*17+ RMs following infection with SIVmac239. Because vaccine-elicited anti-Env antibodies (Abs) have been linked to delayed acquisition of immunodeficiency virus infection following repeated mucosal challenges (22–24), we also evaluated whether these humoral responses would increase the protective efficacy of Vif HW8- and Nef IW9-specific CD8+ T-cells induced by vaccination. To this end, we used a heterologous prime/boost/boost/boost/boost (PBBBB) immunization regimen to vaccinate two groups of Mamu-B*17+ RMs with genes encoding the Vif HW8 and Nef IW9 epitopes. These epitopes were delivered by themselves (Group 1) or together with env (Group 2). As a result, vaccinees in both groups mounted Vif and Nef-specific CD8+ T-cells but only the ones in Group 2 developed Env-specific Abs. We assessed the efficacy of both regimens by repeatedly challenging vaccinees along with sham-vaccinated MHC-I-matched control RMs (Group 3) intrarectally with a marginal dose of SIVmac239. The challenge outcomes varied greatly between Groups 1 and 2, thereby revealing
important aspects of immune containment of lentivirus replication. Here we discuss the relevance of these findings for HIV vaccine development and for understanding the basis of elite control of SIV replication in Mamu-B*17+ RMs.

**Results**

Twenty-three RMs expressing the MHC-I allele Mamu-B*17 were used in this study. These animals were divided among three groups, depending on which immunogens they received. RMs in Group 1 (n = 7) were vaccinated with genes encoding the immunodominant Vif HW8 and Nef IW9 epitopes, whereas those in Group 2 (n = 8) received the same inserts with the addition of env (Fig. 1). The SIV sequences were delivered by a recombinant (r) yellow fever virus 17D (rYF17D) prime followed by three boosts with rDNA plasmids delivered by intramuscular electroporation (EP rDNA). Subsequently, vaccinees in Groups 1 and 2 were boosted once with each of the following viral vectors: adenovirus type-5 (rAd5), vesicular stomatitis virus (rVSV), and rhesus monkey rhadinovirus (rRRV) (Fig. 1). We used this PBBBB vaccine regimen because recurrent antigen stimulation is thought to facilitate the induction of effective cellular and humoral immune responses against lentiviruses (25, 26). It should be noted that some of the vaccine vectors employed in this study encoded segments of vif and nef, or full-length vif and nef fused with other genes, such as tat and rev (see Materials and Methods). As a result, RMs in Groups 1 and 2 also developed cellular immune responses against Tat and Rev. Additionally, some of the vaccine-encoded immunogens also included the subdominant Mamu-B*17-restricted epitopes Nef MW9 (amino acids 195-203) and, in the case of Group 2, Env FW9 (amino acids 830-838). Finally, eight Mamu-B17+ RMs were immunized
with vectors encoding irrelevant antigens or lacking any inserts (“empty” vectors) and served as
the controls for this experiment (Group 3; Fig. 1).

We monitored vaccine-induced CD8+ T-cell responses against Mamu-B*17-restricted epitopes by staining PBMC with fluorochrome-labeled MHC-I tetramers. This analysis revealed that the PBPPP regimen generated high-frequency SIV-specific CD8+ T-cell responses in the majority of the Group 1 and Group 2 vaccinees (Fig. 2). Consistent with the immunodominance of Vif HW8 and Nef IW9 during SIV infection (20), vaccine-elicited CD8+ T-cells in Groups 1 and 2 were primarily directed against these two epitopes (Fig. 2A-D). Curiously, while Nef MW9-specific CD8+ T-cells were undetectable or at borderline levels in all Group 2 vaccinees, low to modest frequencies of these responses were observed in several RMs in Group 1 (Fig. 2E&F). A few macaques in Group 2 developed Env FW9-specific CD8+ T-cells but these responses remained scarce throughout the vaccine phase (Fig. 2G).

We also determined the magnitude of vaccine-induced SIV-specific T-cell responses in Groups 1 and 2 by performing intracellular cytokine staining (ICS) in PBMC at the time of the first IR SIV challenge (Fig. 3). In accordance with the MHC-I tetramer analysis, vaccine-induced CD8+ T-cell responses in Groups 1 and 2 were mainly directed against Vif and Nef, although a few animals in each group developed responses against Rev, Tat, and Env (Group 2 only) as well (Fig. 3A&B). Vaccine-elicited CD4+ T-cell responses in Group 2 were detected at higher frequencies than those in Group 1 and focused primarily on Env (Fig. 3A-C). There was no significant difference in the total magnitude of vaccine-elicited SIV-specific CD8+ or CD4+ T-cell responses between Groups 1 and 2 (Fig. 3C).

Vaccine-elicited Env-specific humoral responses in Group 2 were also evaluated at multiple time points during the vaccine phase. A longitudinal analysis of gp140-binding Abs
showed substantial increases in these responses after the EP rDNA and rAd5 vaccinations (Fig. 4A). Subsequent boosting with rVSV, but not with rRRV, resulted in a modest rise in these responses (Fig. 4A). We also quantified vaccine-elicited gp140-binding Abs on the day of the 1st SIV challenge (Fig. 4B). The median endpoint titer of gp140-binding Abs in Group 2 was 4,800 (Fig. 4B)–a value that was twice as high as that induced by an EP rDNA/rAd5/rVSV/rRRV vaccine regimen recently tested by our group (27). However, these responses were still about 100-fold lower than those generated by live-attenuated SIV vaccination (Fig. 4B). Low levels of NK cell-mediated, Ab-dependent cellular cytotoxicity (ADCC) against SIVmac239-infected cells were also detected in plasma from r08047 and r09062 collected on the day of the 1st SIV challenge, but ADCC activity was either absent or at borderline levels in the other animals (Fig. 4C). No neutralizing antibodies (nAbs) against SIVmac239 were detected on the day of the 1st SIV challenge or after SIV infection (Table 1).

To assess vaccine efficacy, all RMs in Groups 1-3 were subjected to repeated IR challenges with a marginal dose of SIVmac239 every two weeks, starting at study week 101 (Fig. 1). The challenge inoculum consisted of 200 50% tissue culture infective doses (TCID<sub>50</sub>) of an <em>in vivo</em>-titrated SIVmac239 stock. As a reference, the challenge dose employed here typically infects 80% of SIV naïve RMs after six IR exposures. RMs in Groups 1-3 became infected at similar rates (Fig. 5), indicating that vaccine-induced immune responses did not block acquisition of SIVmac239 infection.

Contrary to our expectations based on control of SIVmac239 replication in <em>Mamu-B*08+</em> RMs vaccinated with Vif and Nef (21), vaccine-induced CD8+ T-cell responses targeting the immunodominant Mamu-B*17-restricted Vif HW8 and Nef IW9 epitopes resulted only in a modicum of virologic control in Group 1 (Fig. 6A). Although two vaccinees (r09001 and r08034)
fared well in the chronic phase, none of the Group 1 RMs suppressed viremia to <15 viral RNA copies/mL and the latter RM lost control of viral replication after week 32 PI (Fig. 6A). The number of infected monkeys with chronic phase VLs below 1,000 vRNA copies/mL, even if transiently as in the case of r08034, was similar to that in Group 3, where r10018 developed a viral setpoint of approximately 1,000 vRNA copies/mL while r09083 controlled viral replication to <15 vRNA copies/mL at week 20 PI (Fig. 6A&C).

Surprisingly, however, Group 2 exhibited an entirely unexpected outcome after infection. Despite experiencing peak VLs in excess of 8.5×10^5 vRNA copies/mL of plasma, five Group 2 vaccinees (r05007, r08046, r08062, r08047, and r09062) controlled viral replication to <15 vRNA copies/mL by weeks 4-8 post infection (PI) (Fig. 6B). This corresponded to an average reduction in VLs of 5.3 logs, which was effected within only 2-6 weeks after peak viremia. Four of the five Group 2 vaccinees that showed early viral suppression maintained control of viral replication throughout the chronic phase. RM r08047 was the exception, as its VLs progressively increased after week 20 PI (Fig. 6B). Of note, after decreasing viremia to <15 vRNA copies/mL, the Group 2 controllers experienced occasional VLs blips in the ensuing weeks, similar to those reported by Hansen et al. (28) and Winstone et al. (29). RM r08062 began to deviate from this pattern at week 16 PI, after which viremia became persistent, albeit at low levels (140 vRNA copies/mL at week 75 PI; Fig. 6B).

We compared VLs among Groups 1-3 and made several observations about post-infection viral control in these groups. First, five Group 2 vaccinees had VLs that were either at or below the limit of detection (15 vRNA copies/mL) on day 6 PI (Fig. 7A), implying early control of viral replication. By comparison, only one vaccinee in Group 1 (r09090) and two monkeys in Group 3 (r08051 and r09038) had VLs below 15 vRNA copies/mL at this time point (Fig. 7A).
These differences were not, however, statistically significant (Fig. 7A). Second, peak VLs were significantly lower in Groups 1 and 2 than in Group 3, although these reductions were modest (Fig. 7B). Curiously, the two RMs (r03139 and r09090) with the lowest peak VLs were in Group 1 (Fig. 7B). Third, although setpoint VLs were not significantly decreased in either vaccinated group, Group 2 (but not Group 1) exhibited a significant reduction in nadir VLs, consistent with stringent control of post-acute viremia being the most distinctive feature of Group 2 (Fig. 7C&D). Fourth, it took a median of 1.46 weeks for the Group 1 macaques to experience peak viremia after SIV infection, compared to medians of 2 and 2.5 weeks in Groups 2 and 3, respectively (Fig. 7E). This difference between Group 1 and the other groups was statistically significant, although it is not clear why viral replication peaked earlier in the Group 1 vaccinees. There was no statistically significant difference in the time-to-peak VLs between Groups 2 and 3, despite a few outliers in the latter group (Fig. 7E).

It is important to emphasize that the level of control of chronic phase SIVmac239 replication manifested by the five controller Group 2 vaccinees is exceedingly rare. In fact, of the 197 RMs that have been rectally infected with SIVmac239 as part of eight previous and ongoing SIV vaccine trials conducted by our group, none exhibited the 5.3-log reduction in VLs within 2-6 weeks of peak viremia observed in the five Group 2 controllers (Fig. 8). Of note, this cumulative past experience also included RMs that expressed MHC-I alleles associated with elite control of SIV infection (Fig. 8D-H). These animals were vaccinated with various heterologous prime boost regimens encoding \textit{vif}, \textit{nef}, \textit{rev}, and \textit{tat} (Fig. 8D), \textit{nef} only (Fig. 8E), \textit{rev}, \textit{tat}, and \textit{nef} (Fig. 8F), and \textit{vif} only (Fig. 8G). The \textit{Mamu-B*08+} vaccinees in Fig. 8H were vaccinated with \textit{vif} and \textit{nef} minigenes containing Mamu-B*08-restricted epitopes or with minigenes of other regions of the SIV proteome that lack CD8+ T-cell determinants restricted by Mamu-B*08. As expected, a
fraction of these SIVmac239-infected RMs became ECs but, even in these cases, viremia was rarely reduced to <15 vRNA copies/mL by weeks 4-8 PI (Fig. 8D-H). These different kinetics of virologic control after SIVmac239 infection underscore the uniqueness of the five Group 2 controllers.

We have recently reported an extraordinary case of control of SIVmac239 infection in a vaccinated RM (Fig. 8E and (30)). However, the virus replicating in that animal contained a deletion in nef as early as week 2 PI, which likely compromised its replicative fitness. Given this precedent, we explored the possibility that the Group 2 controllers harbored attenuated viruses, even though we have previously shown that the SIVmac239 challenge stock employed here is made almost entirely of wild-type genomes (30). We conducted this analysis in acute phase (weeks 2-4 PI) plasma from the four RM manifesting the earliest control of viral replication (r08046, r05007, r08062, and r08047) and found >99% homogeneity in nef sequences amplified at week 2 PI (Fig. 9). Viral sequence variation was more prevalent in the later time points, especially in r08062 and r08047 (Fig. 9C&D), but none of the nef sequences analyzed contained insertions or deletions (Fig. 9). Of note, both the Nef IW9 and Nef MW9 epitopes were essentially intact in these acute phase samples (Table 2). Thus, gross genetic defects in nef were not detected in the acute phase virus of 4/5 Group 2 controllers, corroborating the interpretation that the outstanding control of SIVmac239 replication manifested by these RMs was vaccine-mediated.

We also analyzed env sequence evolution in the aforementioned acute phase samples and found no overlap in the diversity patterns detected in r08046, r05007, r08062, and r08047 (Fig. 10). In spite of many low frequency mutations scattered throughout env in all four RMs, only a few variants were present at ≥10% frequencies (Fig. 10). RM r08046 exhibited a R751G
substitution at week 4 PI that was present in 81% of sequence reads (Fig. 10A). This polymorphism is frequently observed in virus isolated from SIVmac239-infected RMs and does not by itself significantly alter virus infectivity in vitro (31). One of the env changes in r08062 resulted in a stop codon in gp120 (W225Stop) and comprised 40% of the circulating viral quasispecies at week 3 PI (Fig. 10C). However, viral variants harboring this stop codon were no longer detected in week 8 plasma from r08062 (Suppl. Fig. 1). This RM also exhibited a synonymous mutation in the Leu codon at position 656 (CTA\>TTA) in 100% of sequence reads (Fig. 10C). The other mutations in r08062 and in r08047 modified amino acids at various positions in gp120 and gp41 (Fig. 10C&D). Variants of the Env FW9 epitope were detected at weeks 2-4 PI but these mutants never reached frequencies >6% (Table 2). A summary of amino acid substitutions in Env detected at ≥10% frequencies in these acute phase samples is shown in Fig. 11. Thus, a shared pattern of env sequence evolution did not explain the rapid post-acute phase virologic control manifested by 4/5 Group 2 controllers.

We then searched for vaccine-induced immune signatures that might explain the distinct virologic outcomes observed in Groups 1 and 2. Eight immunological variables determined either at the time of the first IR SIV challenge or at week 2.4 PI were selected for this analysis (Table 3). These variables were then compared with nadir VLs—the virologic marker most affected by vaccine-induced immune responses (Table 3). The only immunological predictor of virologic control that emerged from this analysis was the endpoint titer of vaccine-induced gp140-binding Abs in Group 2 at the time of the first SIV challenge, which inversely correlated with nadir VLs (Table 3; Fig. 12). This association was not, however, statistically significant after correcting for multiple comparisons. Collectively, these data suggest that in the context of potent
vaccine-elicited SIV-specific T-cell responses, increasing titers of gp140-binding Abs may result in substantial control of viral replication in *Mamu-B*\(^{*}17\)* RMs.

**Discussion**

Here we explored whether vaccinating *Mamu-B*\(^{*}17\)* RMs with inserts encoding Mamu-B*\(^{*}17\)-restricted SIV epitopes would increase the frequency of elite control after SIVmac239 infection, as is the case with vaccination of *Mamu-B*\(^{*}08\)* RMs with Mamu-B*\(^{*}08\)-restricted SIV epitopes (21). We also examined whether the addition of *env* to the vaccine regimen would confer protection from mucosal acquisition of the challenge virus, as has been reported previously (22–24). To our surprise, none of the *vif* and *nef*-vaccinated Group 1 animals controlled viral replication to <15 vRNA copies/mL of plasma, suggesting that the underlying mechanisms of elite control differ between *Mamu-B*\(^{*}17\)* and *Mamu-B*\(^{*}08\)* RMs. Even though the inclusion of *env* in the vaccine did not prevent acquisition of SIVmac239, we identified remarkable levels of virologic control in most of the Group 2 vaccinees. Indeed, 5/8 Group 2 RMs (vaccinated with *vif*, *nef*, and *env*) suppressed viral replication to <15 vRNA copies/mL of plasma by 4-8 weeks PI. This virologic control did not correlate with vaccine-induced SIV-specific T-cells or NK cell-mediated ADCC activity, nor did it depend on anti-SIVmac239 nAbs because serological neutralizing activity against SIVmac239 was not detected at the time of the 1\(^{st}\) SIV challenge or even shortly after infection. Vaccine-elicited gp140-binding Abs at the time of challenge were the only immune predictor of virologic control in Group 2.

Approximately 50% of *Mamu-B*\(^{*}08\)* and 21% of *Mamu-B*\(^{*}17\)* SIV naïve RMs become ECs after SIVmac239 infection (14, 15, 20). Given the virulence of SIVmac239, elite control in
this context is defined as having a setpoint VL ≤1,000 vRNA copies/mL of plasma. Of note, expression of these alleles does not affect acute phase viremia after primary SIVmac239 infection. Indeed, peak VLs are often indistinguishable between unvaccinated SIVmac239-infected RMs expressing *Mamu-B*08 or *Mamu-B*17 and animals lacking these protective MHC-I alleles (14, 15). Despite the predisposition of *Mamu-B*17+ RMs to control SIV replication, the kinetics with which the five Group 2 controllers suppressed viremia in the present experiment was completely different from that seen in typical EC RMs. Our historical VL analysis of SIVmac239-infected animals shows that only a few RMs expressing MHC-I alleles associated with elite control of SIV infection controlled viremia to <15 vRNA copies/mL by week 20 PI (Fig. 8D-H). Even in those few cases, it took on average 12 weeks for their plasma VLs to reach <15 vRNA copies/mL. By comparison, the five Group 2 controllers in the present experiment controlled viremia to <15 vRNA copies/mL after an average of 6 weeks of infection. Vaccine-induced Env-specific immune responses appeared to be required for this impressive outcome considering they were the only feature that distinguished Group 2 from Group 1. This conclusion is also supported by the inverse association between vaccine-elicited gp140-binding Ab titers and nadir VLs in Group 2. Nevertheless, assuming that Abs were involved in control, it is not clear how these vaccine-elicited gp140-specific Abs contributed to virologic control considering that they lacked neutralizing activity against SIVmac239 and their ability to recruit NK cell-mediated ADCC activity against SIVmac239-infected cells did not predict post-infection VLs. One caveat to these analyses is the small number of animals in Group 2, which limited our ability to accurately identify immune correlates of virologic control. Additionally, given the wide range of non-neutralizing Ab effector functions reported for HIV-specific Abs (32), it is possible that antiviral activities other than NK-cell mediated ADCC might have explained the rapid control
of viral replication observed in some of the Group 2 vaccinees. Furthermore, the absence of a
group of MHC-I-matched RMs vaccinated with env only also precludes any definitive
conclusions regarding the role of vaccine-induced Env-specific immune responses in the
challenge outcome. Such a group would have revealed whether or not vaccine-induced Env-
specific immune responses are sufficient for rapid post-acute phase control of viral replication in
*Mamu-B*17+ RMs and also to what extent vaccine-elicited T-cell responses against Vif and Nef
contributed to virologic containment in the five Group 2 controllers. Additional experiments will
be needed to elucidate these issues.

The observation that the Group 1 vaccine regimen had little effect on viral replication
compared to the control group suggests that, unlike *Mamu-B*08+ RMs (21), pre-existing SIV-
specific CD8+ T-cells alone do not facilitate long term virologic suppression in *Mamu-B*17+
RMs. These discordant outcomes imply a differential dependence on Vif- and Nef-specific CD8+
T-cells for elite control of SIV replication in *Mamu-B*17+ and *Mamu-B*08+ animals. In support
of this notion, it is possible to predict which unvaccinated *Mamu-B*08+ RMs will become ECs
after SIVmac239 infection based on patterns of viral sequence evolution in *Mamu-B*08-
restricted epitopes shortly after infection (33). In contrast, the emergence of CD8+ T-cell
“escape” viral variants does not dictate the ability of certain *Mamu-B*17+ animals to contain SIV
viremia (20). In this regard, it is important to mention that other host genetic factors have also
been associated with control of immunodeficiency virus replication (34). Thus, variability in such
factors may explain the different SIV infection outcomes observed in *vif- and nef-vaccinated
*Mamu-B*08+ versus *Mamu-B*17+ RMs. Additional studies will be needed to elucidate the
differences in elite control between *Mamu-B*08+ and *Mamu-B*17+ animals.
Importantly, the ability of gp140-binding Ab titers to predict vaccine-mediated control of SIVmac239 replication is not limited to Mamu-B*17+ RMs. We have recently reported that an EP rDNA/rAd5/rVSV/rRRV vaccine regimen encoding env, gag, vif, rev, tat, and nef resulted in substantial reductions in plasma VLs in SIVmac239-infected RMs lacking MHC-I alleles associated with elite control (27). Similar to the present study, vaccine-elicited Env-specific Abs lacked detectable neutralizing activity against SIVmac239 but their gp140-binding titers correlated with virologic control. Although vaccinees were not protected from SIV infection in either case, the fact that vaccine-induced gp140-binding Abs predicted control of viremia in these two independent settings is instructive for two reasons. First, the observation that high levels of vaccine-induced Env-specific Ab responses might improve control of viral replication in RMs regardless of their MHC-I genotype broadens the applicability of the present findings. Second, any immunological signature associated with containment of SIVmac239 infection warrants further investigation given the stringency of this molecular clone as a challenge virus (35). Indeed, most unvaccinated or drug-naïve SIVmac239-infected Indian RMs experience high peak (10^6-10^9 vRNA copies/mL) and setpoint (10^6 vRNA copies/mL) VLs and are euthanized due to AIDS-defining illnesses by two years after infection (36). Additionally, the SIVmac239 Env glycoprotein is exceptionally difficult to neutralize (37–41), likely due to its closed conformation. Innumerable vaccine regimens have been evaluated using SIVmac239 challenges, but relatively few have resulted in substantial reductions in viremia after infection (27, 29, 42–45) or prevented virus spread beyond the initial foci of infected cells (28, 46). However, except for live-attenuated SIV vaccines (47, 48), no active immunization strategy has consistently afforded sterilizing protection against challenge with SIVmac239. While certain features of the antiviral T-cell response elicited by live-attenuated SIV vaccines have been...
identified as correlates of protection (48–51), vaccine efficacy in this case also relies heavily on Env-specific humoral immunity (52). The fact that only a fraction of SIVmac239Δnef-vaccinated RMs develops detectable anti-SIVmac239 nAb responses imply that Ab-mediated effector functions other than neutralization (e.g., ADCC) can also contribute to the efficacy of live-attenuated SIV vaccines (52, 53). In this light, it is noteworthy that the titers of gp140-binding Abs elicited by SIVmac239Δnef vaccination exceeded those generated by our mixed modality vaccine regimens by approximately 100-fold (Fig. 4). Group 2 had greater titers of these responses than the aforementioned EP rDNA/rAd5/rVSV/rRRV group but this difference was not statistically significant. Although it is impossible at this point to establish a causal link between these anti-Env humoral responses and containment of viral replication, future studies should evaluate if the robust efficacy of live-attenuated SIV vaccination can be replicated by matching its titers of gp140-binding Abs.

In conclusion, here we show that a rYF17D/EP rDNA/rAd5/rVSV/rRRV vaccine regimen encoding env and genes expressing Mamu-B*17-restricted CD8+ T-cell epitopes resulted in stringent control of viral replication in Mamu-B*17+ RMs shortly after SIVmac239 infection. We present evidence suggesting that vaccine-elicited Env-specific Abs devoid of detectable neutralizing activity against SIVmac239 were critical mediators of virologic control and that similar outcomes may be achieved in RMs that do not express protective MHC-I alleles. However, the exact mechanism by which these humoral responses exerted antiviral activity and to what extent cellular immune responses contributed to viral suppression remain undefined. These findings advance our understanding of vaccine-mediated control of lentivirus infection and establish a new benchmark for evaluating HIV vaccine candidates in the SIVmac239 challenge model.
Materials and Methods

Research Animals and Ethics Statement

Twenty-four RMs expressing the Mamu-B*17 MHC-I allele were originally enrolled in this vaccine trial. Unfortunately, RM r04105 had to be euthanized due to recurrent diarrhea and weight loss, thus reducing the size of Group 1 to seven RMs. Because r04105 was euthanized at study week 39—before it was boosted with rAd5, this animal was not included in the immunological comparisons of vaccine-induced T-cell responses between Groups 1 and 2.

The details regarding animal welfare described herein are either similar or identical to those published in one of our previous experiments. “The Indian RMs (Macaca mulatta) utilized in this study were housed at the Wisconsin National Primate Research Center (WNPRC). All animals were cared for in accordance with the guidelines of the Weatherall report and the principles described in the National Research Council’s Guide for the Care and Use of Laboratory Animals under a protocol approved by the University of Wisconsin Graduate School Animal Care and Use Committee” (animal welfare assurance no. A3368-01; protocol no. G00696) (54). “Furthermore, the RMs in this study were managed according to the animal husbandry program of the WNPRC, which aims at providing consistent and excellent care to nonhuman primates at the center. This program is employed by the Colony Management Unit and is based on the laws, regulations, and guidelines promulgated by the United States Department of Agriculture (e.g., the Animal Welfare Act and its regulations, and the Animal Care Policy Manual), Institute for Laboratory Animal Research (e.g., Guide for the Care and Use of Laboratory Animals, 8th edition), Public Health Service, National Research Council, Centers for
Disease Control, and the Association for Assessment and Accreditation of Laboratory Animal Care International. The nutritional plan utilized by the WNPRC is based on recommendations published by the National Research Council. Specifically, RMs were fed twice daily with Teklad Global 20% Protein Primate Diet and food intake was closely monitored by Animal Research Technicians. This diet was also supplemented with a variety of fruits, vegetables, and other edible objects as part of the environmental enrichment program established by the Behavioral Management Unit. Paired/grouped animals exhibiting stereotypical and/or incompatible behaviors were reported to the Behavioral Management staff and managed accordingly. All primary enclosures (i.e., stationary cages, mobile racks, and pens) and animal rooms were cleaned daily with water and sanitized at least once every two weeks. Lights were on a 12:12 diurnal schedule. Vaccinations were performed under anesthesia (Ketamine administered at 5-12 mg/kg depending on the animal) and all efforts were made to minimize suffering. Euthanasia was performed at the end of the study or whenever an animal experienced conditions deemed distressful by one of the veterinarians at the WNPRC. All euthanasia were performed in accordance with the recommendations of the Panel on Euthanasia of the American Veterinary Medical Association and consisted of an IV overdose (greater than or equal to 50 mg/kg or to effect) of sodium pentobarbital or equivalent, as approved by a clinical veterinarian, preceded by ketamine (at least 15 mg/kg body weight) given by the intramuscular (IM) route.”

Additional animal information, including MHC-I alleles, age at the beginning of study, and sex, is shown in Table 4.

**Vaccinations**
The parental live-attenuated YF17DD vaccine strain and four rYF17D vectors expressing individual SIVmac239 inserts were used in this study. The SIVmac239 inserts encoded (1) Vif amino acids 1-110; (2) Nef amino acids 45-210; and (3) the amino terminus of gp41 (Env amino acids 526-690). In order to increase the stability of the latter rYF17D vector, the gp41 fusion peptide was inactivated and the Cys residues at positions 611 and 617 were mutated to Ala. The last SIVmac239 insert (4) consisted of a fusion of gp120 segments corresponding to amino acids 64-109, 209-315, and 343-502. The codon usage of the aforementioned SIVmac239 inserts matched that of YF17D. RMs in Group 2 were vaccinated subcutaneously with 300,000 plaque-forming units (PFU) of each of rYF17D vectors 1-4. These constructs were delivered in separate injections, each containing a final volume of 0.5 mLs. RMs in Group 1 were vaccinated in the same way, except that they received only vectors 1 and 2. RMs in Group 3 were sham-vaccinated with the same dose of the parental YF17DD vaccine. The construction of some of the rYF17D vectors used here has been published elsewhere (55, 56).

A series of three EP rDNA vaccinations, given at 3-week intervals, started at study week 12. Four pCMVkan plasmids encoding SIVmac239 minigenes were used in the EP rDNA vaccinations (57). These minigenes were identical to the ones delivered by the rYF17D vectors, except that the amino acid substitutions described above were reverted to wild-type and the SIV open reading frames were optimized for mammalian expression. RMs in Groups 1 and 2 were vaccinated with pCMVkan constructs encoding the aforementioned vif and nef minigenes. In addition to these constructs, RMs in Group 2 also received two pCMVkan plasmids encoding env minigenes described above. RMs in Group 3 were sham-vaccinated with “empty” pCMCkan plasmid. One milligram of each pCMVkan construct and 0.1 mg of the rhesus interleukin-12-expressing AG157 plasmid were resuspended in 0.5 mL PBS and loaded into separate...
injections. These DNA formulations were administered intramuscularly by the TriGrid in vivo electroporation system (Ichor Medical Systems, Inc., San Diego, CA). Muscles in the thighs and forearms were used for these vaccinations and these anatomical sites were rotated in subsequent immunizations so that each location did not receive vectors encoding the same SIV insert twice.

The rAd5 boost occurred at study week 54. RMs in Groups 1 and 2 were boosted with two rAd5 vectors produced by Viraquest, Inc., each encoding the same vif or nef minigenes described above that were also optimized for mammalian expression. RMs in Group 2 also received a rAd5 vector encoding the full-length SIVmac239 gp160 glycoprotein. RMs in Group 3 were sham-vaccinated with “empty” Ad5 vectors lacking any inserts. The two latter Ad5 constructs were produced by the International AIDS Vaccine Initiative. A dose of $10^{11}$ viral particles of each rAd5 vector was administered intramuscularly to sites in the forearms and thighs.

The rVSV boost occurred at study week 78. The rVSV vector was based on a modified attenuated virus strain developed by Profectus Biosciences (58). A rVSV vector encoding a fusion of the SIVmac239 Nef, Tat, and Vif proteins was used in RMs in both Groups 1 and 2. A separate rVSV vector encoding the SIVmac239 Env protein was administered to the Group 2 vaccinees only. SIV Env was modified by replacing the cytoplasmic tail with the corresponding sequence from VSV G (59). RMs in Group 3 were sham-vaccinated with a rVSV construct encoding the malaria CSP antigen. All three rVSV vectors were provided by Profectus Biosciences. A dose of $10^7$ PFU of each rVSV vector was administered intramuscularly to separate sites in the forearms and thighs.
The rRRV boost occurred at study week 84. The Group 1 RMs were vaccinated with a mixture of three rRRV vectors, each expressing SIVmac239 *vif*, *net*, and a fusion of the *rev-tat-net* genes. The Group 2 RMs received the same rRRV vectors with the addition of a rRRV construct expressing full-length *env*. The Group 3 RMs were sham-vaccinated with a rRRV vector encoding enhanced fluorescent green protein. One mL of PBS containing $7.1 \times 10^7$ genome copies of each appropriate rRRV vector was administered via both intravenous and IR routes. Details about the generation of these rRRV vectors have been described previously (42).

**SIVmac239 challenges**

Seventeen weeks after the rRRV boost, RMs in Groups 1-3 were subjected to repeated IR inoculations of 200 TCID$_{50}$ ($4.8 \times 10^5$ vRNA copies) of the same SIVmac239 stock described in (30). These IR challenges occurred every two weeks. Plasma VLs were assessed six and ten days after each exposure. Once an animal experienced a positive VL at either one of these time points, it was no longer challenged. Only RMs that remained aviremic at both time points were re-challenged on day 14.

**SIV RNA viral load measurements**

The following description of plasma SIV RNA viral loads is identical to that published in one of our recent manuscripts (27). "VLs were measured using 0.5 mL of EDTA-anticoagulated rhesus macaque plasma based on a modification of a previously published (60). Total RNA was extracted from plasma samples using QIAgen DSP virus/pathogen Midi kits, on a
QIASymphonyXP laboratory automation instrument platform. Six replicate two step RT-PCR reactions were performed per sample using a random primed reverse transcription reaction, followed by 45 cycles of PCR using the following primers and probe: forward primer: SGAG21: 5′-GTCTGCGTCAT(dP)TGGTGCA TTC-3′; reverse primer SGAG22: 5′-CACTAG(dK)TGTCCTGCAGT(dP)TGTTTTG-3′; probe: PSGAG23: 5′-FAM-CTTC(dP)TCAGT(dK)TGTTTCACTTTCTCTGCG-BHQ1-3′. The limit of reliable quantitation on an input volume of 0.5 mL of plasma was 15 vRNA copies/mL.

**Quantification of Mamu-B*17 tetramer+ CD8+ T-cells in PBMC**

The following description is either identical or similar to that published in one of our recent manuscripts (27). Fluorochrome-labeled Mamu-B*17 tetrarmers produced at the NIH Tetramer Core Facility were used to quantify SIV-specific CD8+ T-cells in peripheral blood mononuclear cells (PBMC) according to a recently published protocol (61). Approximately 80,000 PBMC were incubated in R10 medium (RPMI 1640 medium supplemented with GlutaMax [Life Technologies], 10% FBS [VWR], and 1% antibiotic/antimycotic [VWR]) with titrated amounts of each tetramer at room temperature (RT) for 45 min. The cells were then stained with fluorochrome-labeled monoclonal antibodies (mAbs) directed against the surface molecules CD3 (clone SP34-2), CD8α (clone RPA-T8), CD14 (clone M5E2), CD16 (clone 3G8), and CD20 (clone 2H7) for 25 min. This step also included an amine-reactive dye (ARD; Live/DEAD Fixable Aqua Dead Cell Stain; Life Technologies). The cells were then washed with Wash Buffer (Dulbecco's PBS with 0.1% bovine serum albumin and 0.45 g/L NaN3) and fixed with PBS containing 2% of paraformaldehyde (PFA) for 20 min at 4°C. The cells were washed one more
time before they were acquired in a Special-Order Product BD LSR II cytometer. The gating strategy used to analyze the data has been described elsewhere (61). Briefly, we used FlowJo 9.6 to determine the percentages of live CD14- CD16- CD20- CD3+ CD8+ tetramer+ lymphocytes in PBMC.

**Intracellular cytokine staining (ICS) assays**

The following description is either identical or similar to that published in one of our recent manuscripts (27). Freshly isolated PBMC were cultured in R10 medium containing unlabeled co-stimulatory mAbs against CD28 and CD49d and a phycoerythrin-conjugated mAb specific for CD107a. The tubes were kept in a 5.0% CO₂ incubator for 9 hours (h) at 37 °C. One hour into the incubation period, Brefeldin A (Biolegend, Inc.) and GolgiStop (BD Biosciences) were added to all tubes in order to inhibit protein transport. The antigen stimuli consisted of three or five pools of SIVmac239 peptides (15mers overlapping by 11 aa) corresponding to (1) the entire Vif protein (amino acids 1-214), (2) the entire Nef protein (amino acids 1-263), (3) both Rev (amino acids 1-107) and Tat (amino acids 1-130) proteins, (4) Env gp120 (amino acids 1-531) and (5) Env gp41 (amino acids 516-879). The final concentration of each 15mer in the ICS tubes was 1.0 μM. The same steps outlined above were used to stain the cells with mAbs against surface markers and subsequently fix them. The surface staining master mix included mAbs against CD4 (clone OKT4; Biolegend, Inc.) and CD8 (clone RPA-T8; Biolegend, Inc.), in addition to the same mAbs against CD14, CD16, and CD20 and the ARD reagent described above. For 10 min, cells were permeabilized by homogenization in Perm Buffer (1X BD FACS lysing solution 2 [Beckton Dickinson] and 0.05% Tween 20 [Sigma-Aldrich]), and then were washed with Wash
Buffer. Lastly, cells were incubated for 1 h in the dark at RT with mAbs against CD3 (clone SP34-2; BD Biosciences), IFN-γ (clone 4S. B3; Biolegend, Inc.), TNF-α (clone Mab11; BD Biosciences), and CD69 (clone FN50; Biolegend, Inc.). Once this incubation was completed, cells were washed and stored at 4 °C until acquisition.

After gating on live CD14- CD16- CD20- CD3+ lymphocytes, we selected either CD4+ or CD8+ cells for subsequent analyses. Cells were considered positive for IFN-γ, TNF-α, or CD107a only if these molecules were co-expressed with CD69, a marker of recent activation. Leukocyte activation cocktail (LAC; BD Pharmingen)–stimulated cells stained with fluorochrome-labeled control mAbs of the same isotypes as those against IFN-γ, TNF-α, and CD107a guided the identification of positive populations. Two criteria were used to determine a positive response. First, the frequency of events had to be ≥2-fold higher than their corresponding values in background-subtracted negative control tests. Second, the gates for each response had to contain ≥10 events. These calculations were performed with Microsoft Excel and results were presented as the percentage of responding CD4+ or CD8+ T-cells, that is, live CD14- CD16- CD20- CD3+ lymphocytes of either subset producing any combination of IFN-γ, TNF-α, or CD107a.

Anti-Env antibody measurements by ELISA

The following description is identical to that reported by our group in a recent publication (27). "Vaccine induced anti-Env responses were measured by ELISA. To begin, the ELISA plate was coated with 100 μL of purified SIVmac239 gp140 protein (Immune Technology Corp. #IT-001-140p) at a concentration of 0.5 μg/mL and incubated overnight at RT. On the following day,
the plate was washed with 1× PBS-Tween20 and wells were blocked with 300 μL of 5% powdered milk in PBS for 1 hr at 37 °C. Subsequently, the plate was washed and 100 μL of diluted plasma samples were added to the corresponding wells. After a 1-hr incubation at RT, the plate was washed and 100 μL of a 1:2,000 dilution of Goat Anti-Monkey IgG-HRP antibody (Santa Cruz Biotechnology, sc-2458) were added to all wells for 1 hr at 37 °C. Finally, the plate was washed before being developed with 100 μL of 3,3',5,5'-Tetramethylbenzidine (EMD Millipore, 613544-100ML). After a short incubation, the reaction was stopped with TMB Stop Solution (Southern Biotech, 0412-01) and the plate was read (Biotek Synergy 2) at 450 nm. The endpoint antibody titers of vaccine-induced anti-Env antibody responses were measured in serum collected at the time of SIV challenge. These titers were determined as the greatest dilution at which the absorbance in experimental wells was at least two-fold higher than that measured in pooled pre-vaccination serum from all animals in the experiment.”

**Antibody dependent cellular cytotoxicity (ADCC) assay**

The following description is identical to that reported by our group in a recent publication (27). “The SIVmac239 and SHIV\textsubscript{AD8-EO} stocks used in ADCC assays were produced by transfection of infectious molecular clones into HEK293T cells using GenJet transfection reagent (SignaGen). Virus-containing supernatants were collected 48 and 72 h post-transfection and stored at –80 °C. The SHIV\textsubscript{AD8-EO} clone was provided by Dr. Malcom Martin (NIAID, Bethesda, MD). After heat inactivation for 30 min at 56 °C, RM plasma samples were tested for non-specific ADCC due to the presence of antibodies to human cellular antigens by co-incubating uninfected CEM.NKR-CCR5-sLTR-Luc target cells (AIDS Research and Reference
Reagent Program, Division of AIDS, NIAID, NIH) with an NK cell line (KHYG-1 cells) expressing RM CD16 at a 10:1 effector-to-target ratio in the presence of serial dilutions of plasma (62). This NK cell line was developed in house, as described previously (59). Non-specific lysis was detected as a reduction in background luciferase activity (% RLU) for target cells incubated with NK cells in the presence compared to the absence of plasma. Plasma samples that directed ADCC against uninfected cells were depleted of anti-human antibodies by repeated cycles of incubation with CEM.NKR-CCR5-sLTR-Luc cells, followed by centrifugation and plasma transfer, until ADCC responses to uninfected cells were no longer detectable."

“To measure ADCC activity in plasma of vaccinated animals, CEM.NKR-CCR5-sLTR-Luc target cells were infected with SIVmac239 or SHIVAD8-EO (internal negative control) by spinoculation for 3 h at 1200 × g in the presence of 40 µg/mL polybrene (EMD Millipore). Four days post-infection, target cells were incubated with the NK cell line KHYG-1 at a 10:1 effector-to-target ratio in the presence of serial plasma dilutions. Luciferase activity was measured after 8 h using the britelite plus luciferase assay system (PerkinElmer). Triplicate wells were tested at each plasma dilution, and wells containing effector cells incubated with uninfected or infected target cells in the absence of plasma were used to determine background and maximal luciferase activity, respectively. ADCC responses were calculated from the dose-dependent loss of luciferase activity in the presence of plasma relative to background and maximal luciferase control wells.”

Pseudovirus neutralization assays
The following description is nearly identical to that reported by our group in a recent publication (27). “Replication incompetent SIVmac239 pseudovirus was produced by co-transfecting env plasmids with an env-deficient backbone plasmid (pSG3Δenv) in HEK293T cells in a 1:2 ratio, using the X-tremeGENE 9 transfection reagent (Roche). Pseudovirus was harvested after 72 h by sterile-filtration (0.22 µm) of cell culture supernatants, and neutralization was tested by incubating pseudovirus and serum for 1 h at 37 °C before transferring them onto TZM-bl cells (AIDS Research and Reference Reagent Program, Division of AIDS, NIAID, NIH) as previously described (63). Neutralization was measured in duplicate wells within each experiment. Neutralization was tested starting at 1:20 serum dilutions followed by nine serial 3-fold dilutions to ensure highest sensitivity and range of detection. Neutralization IC$_{50}$ titers were calculated using the ‘One site – Fit logIC50’ regression in Graphpad Prism v7.0. We could not detect vaccine-induced nAb titers against SIVmac239 pseudovirus in any of the RMs in Group 2 at the time of the first SIV challenge.”

Amplicon-based sequencing of SIV nef and env

Viral RNA was isolated from plasma using the QIAamp viral RNA mini kit (Qiagen) according to the manufacturer’s protocol and eluted in 60 µL of AVE buffer, aliquoted and stored at −80°C for future use. Viral RNA was reverse transcribed and amplified by using the SuperScript III One-Step RT-PCR system with High Fidelity Platinum Taq polymerase (Invitrogen). The 3’ half of the genome was amplified in two amplicons SIVmac239-Amp3-5660-F (5’-GGCATAGCCTCATAAAATATCTG-3’) and SIVmac239-Amp3-8487-R (5’-ATTGCAGAACCTGCCGTTG-3’); SIVmac239-Amp4-7821-F (5’-
CAGTCACCATTATGTCTGGATTG-3′) and SIVmac239-Amp4-10235-R (5′-GAATACAGAGCGAAATGCAGTG3′). All SIVmac239 primer positions are based on published SIVmac239 sequence present in GenBank (M33262). The RT-PCR conditions were as follows for 50°C for 30 min, 94 °C for 2 min; 40 cycles of 94 °C for 15 seconds (sec), 55 °C for 30 sec, and 68 °C for 3 min; and 68 °C for 5 min.

Amplicons were visualized on a 1.0% agarose gel and purified using the Purelink quick gel extraction kit (Invitrogen). RT–PCR products were quantified using a Promega quantiflor-ST fluorometer (Promega) and analyzed for quality using an Agilent 2100 bioanalyzer with high sensitivity DNA chips. PCR amplicons were fragmented and barcoded using NexteraXT DNA Library Prep Kit, as per manufacturer’s protocol. Samples were pooled and sequenced on an Illumina MiSeq platform, using a 2 x 250 bp V2 reagent kit. Paired-end reads obtained from Illumina MiSeq were then processed using our internal sequencing analysis pipeline which consists of removing PCR duplicate reads using FastUniq (64), quality trimming with trimmomatic (65) and de novo assembly using VICUNA (66) with finishing and annotation completed with V-FAT (https://www.broadinstitute.org/viral-genomics/v-fat). Reads were then aligned to the consensus assembly using Mosaik v2.1.73 with V-Phaser v2.0 used to call intra-host variants as described elsewhere (66–68). All raw sequence reads have been deposited to the NCBI Sequence Read Archive under the study accession number SRP016012 with the following experimental accession numbers SRX3797244 – SRX3797252.

Statistics

Peak VLs were determined as the highest VL measurement within the first 4 weeks after infection. Nadir VLs were considered as the lowest VL measurement between peak and week 8.
PI. Setpoint VLs were calculated as the geometric mean of all VLs measured between week 8 PI and the last chronic phase time point available. The Mann-Whitney U test was used to compare the total magnitude of vaccine-induced SIV-specific T-cell responses between Groups 1 and 2. The Kaplan-Meier method and log-rank test were used to determine whether the Group 1 or Group 2 vaccine regimens affected acquisition of SIV infection. For this analysis, the time-to-productive infection was analyzed using the Kaplan-Meier method and the differences between Groups 1 and 3 and Groups 2 and 3 were evaluated using log-rank tests. The Mann-Whitney U test was also used to determine the efficacy of each vaccine regimen in reducing viral replication. Viral loads at multiple time points were compared between each of Groups 1 and 2 and the control Group 3. Lastly, the Spearman rank correlation was used to indicate immune correlates of protection. All significance tests were two-tailed.

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Figure legends

Figure 1. Experimental design. Twenty-three *Mamu-B*17+ RMs were vaccinated with a rYF17D/EP rDNA/rAd5/rVSV/rRRV regimen and divided among three groups depending on which vaccine inserts they received. Animals in Group 1 (n = 7) were vaccinated with *vif* and *nef* sequences. Animals in Group 2 (n = 8) were vaccinated with the same SIV inserts plus *env*. RMs in Group 3 were sham-vaccinated with vectors lacking SIV genes or expressing irrelevant inserts and served as the control group for this experiment. At study week 101, vaccine efficacy was assessed by subjecting all animals to repeated intrarectal (IR) challenges with a marginal dose of SIVmac239 (200 TCID<sub>50</sub>) every two weeks.

Figure 2. Development of vaccine-induced CD8+ T-cell responses against *Mamu-B*17-restricted SIV epitopes in Groups 1 and 2. Fluorochrome-labeled *Mamu-B*17 tetramers folded with peptides corresponding to SIV epitopes were used to track vaccine-elicited SIV-specific CD8+ T-cells in PBMC from RMs in Group 1 (A, C, E) and Group 2 (B, D, F, and G). The percentages of live tetramer+ CD8+ T-cells specific for Vif HW8 (A and B), Nef IW9 (C and D), Nef MW9 (E and F), and Env FW9 (G) are shown at multiple time points throughout the vaccine phase. No Env FW9-specific CD8+ T-cell data are available for the Group 1 RMs because those animals were not vaccinated with *env*. The times of each vaccination (black vertical dotted lines) and the day of the first IR SIV challenge (red vertical solid line) are indicated in each graph. RMs in Groups 1 and 2 are color-coded in blue and black, respectively.
Figure 3. Vaccine-induced SIV-specific CD4+ and CD8+ T-cell responses in Groups 1 and 2 at the time of the first SIV challenge. CD8+ and CD4+ T-cell responses were measured in PBMC by ICS using pools of peptides (15mers overlapping by 11 amino acids) spanning the appropriate SIVmac239 proteins. Peptides covering the Rev and Tat proteins were tested together. The percentages of responding CD4+ or CD8+ T-cells displayed in all panels were calculated by adding the frequencies of positive responses producing any combination of three immunological functions (IFN-γ, TNF-α, and CD107a). The magnitude and specificity of vaccine-induced CD8+ (left panels) and CD4+ (right panels) T-cell responses are shown for Group 1 (A) and Group 2 (B). C) Comparison of the total magnitude of vaccine-elicited SIV-specific CD8+ (left panel) and CD4+ (right panel) between Groups 1 and 2. The Mann-Whitney U test was used for these comparisons and no statistically significant difference was found. RMs in Groups 1 and 2 are color-coded in blue and black, respectively, and each symbol corresponds to one vaccinee. Lines represent medians. NA, not applicable. NS, not significant.

Figure 4. Vaccine-induced Env-specific humoral immune responses in Group 2. A) Env-binding antibodies (Abs) were measured by ELISA using plate-bound gp140 at multiple time points during the vaccine phase. Straight 1:200 dilutions of plasma from each RM in Group 2 were used for this analysis. B) Log-transformed endpoint titers of vaccine-induced gp140-binding Abs in sera from the Group 2 vaccinees collected at the time of the first SIV challenge. As a reference, these values were plotted alongside the endpoint titers of gp140-binding Abs in four RM that had been infected with SIVmac239Δnef for 28 weeks as part of a previous experiment (69). The endpoint titers of gp140-binding Abs in macaques vaccinated with an EP rDNA/rAd5/rVSV/rRRV regimen encoding env, gag, vif, tat, rev, and nef are also shown (27). C)
Ab-dependent cellular cytotoxicity (ADCC) activity was screened against SIVmac239-infected target cells using plasma collected from the Group 2 vaccinees at the time of the first IR SIV challenge (black lines). SHIV<sub>AD8-EO</sub>-infected target cells were used as internal controls (green lines) for non-specific killing. The decrease in relative light units indicates the loss of virus-infected cells in the presence of an NK cell line during the duration of the assay. Dotted lines denote 50% activity. Plasma from an SIV-infected RM (382-03) with a defined ADCC titer against SIVmac239-infected cells was used as a positive control for these measurements.

**Figure 5. Kaplan-Meier rate of infection after repeated IR challenges with SIVmac239.**
Macaques in Groups 1-3 were inoculated intrarectally with 200 TCID<sub>50</sub> of SIVmac239 every other week. The rate of SIV infection in Groups 1 and 2 was not significantly different than that of Group 3. The <i>P</i> value for the Group 1 versus Group 3 comparison was 0.96. The <i>P</i> value for the Group 2 versus Group 3 comparison was 0.28.

**Figure 6. Plasma viral RNA levels after SIVmac239 infection.** Viral load (VL) traces for individual animals in Group 1 (A), Group 2 (B), and Group 3 (C). VLs were log-transformed and correspond to the number of vRNA copies/mL of plasma. The dotted lines in all the graphs are for reference only and indicate a VL of 10<sup>3</sup> vRNA copies/mL. The dashed lines are also for reference only and denote a VL of 10<sup>6</sup> vRNA copies/mL. The pink rectangle in each graph frames the interval during which five Group 2 vaccinees controlled viremia to <15 vRNA copies/mL. Groups 1, 2, and 3 are color-coded in blue, black, and red, respectively.

**Figure 7. SIV plasma viral load comparisons among Groups 1, 2, and 3.** A) Plasma viral
loads (VLs) measured on day 6 post infection (PI). B) Peak VLs. C) Nadir VLs. D) Setpoint VLs, calculated as the geometric mean of VLs measured between week 8 PI and the last chronic phase time point available. E) Time to peak viremia, determined as the week PI when each animal experienced its peak VL. The dotted lines in graphs A-D are for reference only and indicate a VL of $10^3$ vRNA copies/mL. The dashed lines in graphs A-D are also for reference only and denote a VL of $10^6$ vRNA copies/mL. Lines represent medians and $P$-values were calculated using the Mann-Whitney U test. Groups 1, 2, and 3 are color-coded in blue, black, and red, respectively, and each symbol corresponds to one vaccinee.

Figure 8. Viral loads in the five Group 2 controllers compared to those in RMs historically infected with SIVmac239. The outcome of SIVmac239 infection in the five Group 2 vaccinees that manifested early control of viral replication was compared to those of 197 RMs that were rectally infected with SIVmac239 as part of eight SIV vaccine trials conducted by our group. The historical viral loads (VLs) for each independent SIV vaccine trial are plotted in panels A-H and include both vaccinated (gray lines) and control (salmon lines) RMs. VLs from RMs that expressed MHC-I alleles associated with elite control of SIV infection (i.e., Mamu-B*08 and Mamu-B*17) are shown in dashed lines. VLs from RMs that did not express these protective MHC-I alleles are shown in solid lines. To better visualize the early control of viral replication manifested by the five Group 2 vaccinees, their VLs (black lines) were also plotted in each panel. Additionally, only the first 20 weeks of SIV infection are shown in each graph. A) In trial #1, twenty-four RMs were vaccinated with env, gag, vif, rev, tat, and nef delivered by either rRRV alone or rRRV followed by two boosts with rAd5 and rVSV. Eight RMs served as the controls for the challenge phase (Martins et al., unpublished data). B) The details of this
experiment were published recently (27). Briefly, thirty-two RMs were vaccinated with an EP rDNA/rAd5/rVSV/rRRV regimen encoding four different sets of SIV inserts. Eight RMs served as the controls for this experiment. One vaccinee did not get infected, so VL traces for 39 RMs are shown in the graph. C) The details of this experiment were published recently (70). Briefly, four different mixed modality vaccine regimens were used to deliver minigenes of SIV gag, vif, and nef to 32 RMs. Eight RMs served as the controls for this experiment. D) Ten Mamu-B*08+ RMs were vaccinated with a rAd5/rVSV/rRRV regimen encoding vif, rev, tat, and nef and six MHC-I-matched RMs served as the controls for the challenge phase (Martins et al., unpublished data). Three vaccinees did not acquire SIV infection in this experiment so VL traces for 13 RMs are shown in the graph. E) The details of this experiment were published recently (71). Briefly, sixteen Mamu-B*08+ RMs were vaccinated with an EP rDNA/rYF17D/rAd5 regimen encoding nef and two MHC-I-matched macaques served as the controls for the challenge phase. F) The details of this experiment were published recently (30). Four RMs, two of which were Mamu-B*08+, were vaccinated with an EP rDNA/rYF17D/rRRV regimen encoding either gag or nef. For unknown reasons, the animal highlighted in pink harbored a nef-deleted SIV variant as early as week 2 PI. The replicative fitness cost imposed by this nef deletion likely underlies the stringent control of viral replication manifested by this animal. G) Twenty RMs were vaccinated with an EP rDNA/recombinant vaccinia/rVSV/rAd5/rRRV regimen encoding vif only. Twenty MHC-I-matched RMs served as the controls for the challenge phase. Eight vaccinees and nine control RMs expressed either Mamu-B*08 or Mamu-B*17 (Martins et al., unpublished data). One vaccinee did not acquire SIV infection so VL traces for 39 RMs are shown in the graph. H) The details of this experiment were published elsewhere (21). Briefly, sixteen Mamu-B*08+ RMs were vaccinated with a rYF17D/rAd5 regimen encoding either vif and nef minigenes containing...
Mamu-B*08-restricted epitopes or regions of the SIV proteome lacking epitopes restricted by Mamu-B*08. Four MHC-I-matched RMs served as the controls for the challenge phase.

**Figure 9. nef sequence diversity in acute phase plasma from four Group 2 controllers.**

Heat maps illustrating the levels of sequence diversity at each codon in the nef open reading frame. The range of amino acids covered by each row is indicated on the right-hand side of the figure and should be used only as a reference because both synonymous and non-synonymous mutations are considered in these heat maps. The codon corresponding to the first amino acid in Nef is located in the top left corner of each grid. Each row spans 17 amino acids of the Nef protein, except for the bottom row, which covers 9 amino acids and includes the last residue at position 264. The panels on the left correspond to plasma samples collected at week 2 post infection (PI). The panels on the right correspond to plasma samples collected at week 3 PI, or week 4 PI in the case of r08046. Data from r08046, r05007, r08062, and r08047 are shown in A, B, C, and D, respectively.

**Figure 10. env sequence diversity in acute phase plasma from four Group 2 controllers.**

Heat maps illustrating the levels of sequence diversity at each codon in the env open reading frame. The range of amino acids covered by each row is indicated on the right-hand side of the figure only as a reference because both synonymous and non-synonymous mutations are considered in these heat maps. When applicable, the frequency and location of amino acid substitutions are described in the text of the manuscript and in Figure 11. The codon corresponding to the first amino acid in Env is located in the top left corner of each grid. Each row spans 30 amino acids of the Env protein, except for the bottom row, which covers 9 amino acids.
acids and includes the last residue at position 879. The panels on the left correspond to plasma samples collected at week 2 post infection (PI). The panels on the right correspond to plasma samples collected at week 3 PI, or week 4 PI in the case of r08046. Data from r08046, r05007, r08062, and r08047 are shown in A, B, C, and D, respectively.

**Figure 11.** Summary of amino acid substitutions in Env present at ≥10% frequencies in acute phase samples from four Group 2 controllers. Each substitution is enclosed by a square and its details are listed in a text box outside the Env main sequence. The details provided for each amino acid substitution include the animal ID, time point post infection, position in the Env protein, and the frequency of sequence reads displaying the wild-type (boldface type) and mutant residues. As a reference, several topological features of the SIV Env protein are shown in the figure, including the five variable loops (V) in gp120, the gp120/gp41 cleavage site, the membrane spanning domain and highly immunogenic region of gp41. The location of the Mamu-B*17-restricted Env FW9 epitope is also indicated in the carboxyl terminus of gp41.

**Figure 12.** Association between titers of vaccine-induced gp140-binding Abs at the time of the 1st SIV challenge and nadir viral loads in Group 2. This is a graphical representation of the comparison between the log_{10} endpoint titers of vaccine-induced gp140-binding antibodies at the time of the first IR SIV challenge and nadir viral loads in Group 2 listed in Table 3. The symbol for macaque r09062 (open black square) is masked by the symbol of r05007 (filled black square) because both animals had the same titer of gp140-binding antibodies.
(25,600) and nadir VL (15 vRNA copies/mL of plasma). The Bonferroni-corrected $P$ value is shown in the figure. Each symbol corresponds to one Group 2 vaccinee.
Table 1. Serum neutralization of SIVmac239 in Group 2*

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<td>rh1982</td>
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*Because none of the samples resulted in a 50% reduction in SIV infection *in vitro* at the lowest reciprocal serum dilution (1:20), the 50% inhibitory dose for each animal is shown as <20.
Table 2. Percentage of viral sequences encoding intact Mamu-B*17-restricted CD8+ T-cell epitopes in acute phase samples from four Group 2 controllers.

<table>
<thead>
<tr>
<th>Animal ID</th>
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<th>Nef</th>
<th>Env</th>
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<td>IW9</td>
<td>MW9</td>
<td>FW9</td>
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<td>100%</td>
<td>99.7%</td>
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<td>100%</td>
<td>100%</td>
<td>100%</td>
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Week 3 or 4 PI

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</table>

N.A., not available. *No week 3 PI plasma was available from r08046 so the sample collected at week 4 PI was used instead.

Downloaded from http://jvi.asm.org on September 27, 2018 by guest
Table 3. Correlations between immunological variables measured in vaccinees in Groups 1 and 2 and their respective nadir VLs.

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<tr>
<th>Immunological variable</th>
<th>Groups included in the analysis</th>
<th>Time point</th>
<th>Correlation coefficient ($r$)</th>
<th>Unadjusted $P$-value</th>
<th>Bonferroni-corrected $P$-value</th>
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<td>Time of 1st SIV challenge</td>
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Table 4. Animal characteristics.

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SIVmac239 inserts

**Group 1**  
\[\text{vif} \quad \text{rev} \quad \text{tat} \quad \text{nef} \]  
\(n = 7\)

**Group 2**  
\[\text{env} \quad \text{vif} \quad \text{rev} \quad \text{tat} \quad \text{nef} \]  
\(n = 8\)

**Group 3**  
Sham vaccinated controls  
\(n = 8\)

\[\text{rYF17D} \quad \text{EP rDNA} \quad \text{rAd5} \quad \text{rSV} \quad \text{rRRV} \quad \text{iR challenges with SIVmac239 every 2 weeks}\]

\(\text{Week 0} \quad 12 \quad 15 \quad 18 \quad 54 \quad 78 \quad 84 \quad 101\)
A) Vaccine-elicited gp140-binding Abs

Absorbance at 450 nm

Weeks after rYF17D prime

0 10 20 30 50 60 70 80 90 100

EP rDNA rAd5 rVSV rRRV 1st SIV challenge

B) Titer of gp140-binding Abs on the day of 1st SIV challenge

Log₁₀ Ab titer

Group 2 SIVmac239Δnef

EP rDNA/rAd5/rVSV/rRRV

P = 0.004
P = 0.12

C) Vaccine-induced ADCC activity in Group 2

r08046

% Relative light units

Reciprocal plasma dilution

ADCC against SIVmac239-infected target cells

ADCC against SHIVΔΔE0-infected target cells