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ABSTRACT

Mucosal surfaces are vulnerable to human immunodeficiency virus (HIV)/simian immunodeficiency virus (SIV) infection and thus are key sites for eliciting vaccine-mediated protection. Vaccine protocols carried out at the Yerkes Primate Research Center utilized SIVmac239-based immunization strategies with intrarectal and intravaginal SIVsmE660 challenge of rhesus macaques. We investigated whether there were genetic signatures associated with SIVsmE660 intrarectal and intravaginal transmissions in vaccinated and unvaccinated monkeys. When transmitted/founder (T/F) envelope (Env) sequences from 49 vaccinated and 15 unvaccinated macaques were compared to each other, we were unable to identify any vaccine breakthrough signatures. In contrast, when the vaccinated and control T/F Envs were combined and compared to the challenge stock, residues at gp120 positions 23, 45, 47, and 70 (Ile-Ala-Lys-Asn [I-A-K-N]) emerged as signatures of mucosal transmission. However, T/F Envs derived from intrarectal and intravaginal infections were not different. Our data suggest that the vaginal and rectal mucosal environments both imposed a strong selection bias for SIVsmE660 variants carrying I-A-K-N that was not further enhanced by immunization. These findings, combined with the strong conservation of A-K-N in most HIV-2/SIVmm isolates and the analogous residues in HIV-1/SIVcpz isolates, suggest that these residues confer increased transmission fitness to SIVsmE660.

IMPORTANCE

Most HIV-1 infections occur across a mucosal barrier, and it is therefore important to understand why these sites are vulnerable and how to protect them with a vaccine. To gain insight into these questions, we studied rhesus macaques that were vaccinated with SIVmac239 and unvaccinated controls to determine whether the SIVsmE660 viral variants that infected these two groups were different. We did not find differences between viral variants in the absence versus presence of vaccination-induced immunity, but we did find that the SIVsmE660 viral variants that infected the monkeys, regardless of vaccination, were different from the dominant population found in the viral challenge inoculum. Our data suggest that the mucosal environments of the vagina and rectum both impose a strong selection for the SIVsmE660 variants in the challenge inoculum that are most like SIV and HIVs that circulate in nature.

The majority of human immunodeficiency virus type 1 (HIV-1) transmission events occur across the genital or rectal mucosa and involve a reduction in the diversity of the infecting viral quasispecies (1). This genetic bottleneck of HIV-1 transmission has been attributed to various features of the HIV-1 envelope (Env) glycoproteins (2–8), resistance to interferon (7,9), and a selection bias toward consensus residues that confer increased in vivo fitness (10). These studies notwithstanding, it has been difficult to pinpoint the attributes that directly facilitate transmission of a particular HIV-1 variant from a genetically diverse quasispecies. A primary focus of nonhuman primate simian immunodeficiency virus (SIV) challenge models is therefore to recapitulate the critical events that occur during HIV-1 transmission so that any observed effect, including protection, is accurately reflected by the model. An approach in which rhesus macaques are inoculated intravaginally or intrarectally at regular intervals with a relatively low dose of SIV has been shown to recapitulate the hallmark reduction in viral diversity characteristic of HIV-1 transmission (11–14). As a result, the repeated, low-dose challenge method has become widely accepted and commonly utilized to model protection against mucosal SIV infection (14–32). Despite these numerous studies, the characteristics of the virus that influence intravaginal and intrarectal transmission and protection are incompletely understood.

We previously demonstrated that an SIVmac239-based regimen consisting of two DNA primes followed by two modified vaccinia virus Ankara (MVA) boosts provided significant protection against repeated, low-dose SIVsmE660 intrarectal challenge. Including granulocyte-macrophage colony-stimulating factor (GM-CSF) or CD40 ligand (CD40L) as an adjuvant with the DNA
primes further enhanced this protection (16–18, 33). We followed up on these results with a trial testing the protective capacity of DNA/MVA vaccination regimens and a novel protein-based prime boost regimen against intravaginal SIVsmE660 challenge. These studies were conducted at a single primate center using standard methodology, parallel immunization schedules, and Env immunogens all derived from SIVmac239. In addition, the challenge stock utilized for the follow-up trials (NM2010) was derived from the same SIVsmE660 seed stock as the challenge stock used in the initial trials (VH2000) and was expanded in vitro using the same protocol. The samples collected from these trials therefore provided a unique opportunity to explore genetic bottlenecks in SIVsmE660 Env that could be associated with SIVmac239-based vaccine protection and/or mucosal transmission.

MATERIALS AND METHODS

Ethics statement. The Emory University Institutional Animal Care and Use Committee (AWA no. A3180-01) approved these studies of nonhuman primates under protocol YER-2000936-061014GA. This study was conducted in strict accordance with U.S. Department of Agriculture regulations and the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health (34). SIV-infected animals were housed in standard nonhuman primate cages, received standard primate feed as well as fresh fruit and enrichment daily, and had continual access to water. Animals had continuous access to enrichment resources, including objects for perching and other manipulanda. Animal welfare was monitored daily. Appropriate procedures were performed to ensure that potential distress, pain, or discomfort was alleviated. The sedative ketamine (10 mg/kg of body weight) or telazol (4 mg/kg) was used for blood draws. Euthanasia using pentobarbital (100 mg/kg) under anesthesia was performed only when deemed clinically necessary by veterinary medical staff and according to IACUC endpoint guidelines.

Rhesus macaque vaccine trials. Plasma samples were obtained from vaccinated or unvaccinated rhesus macaques following SIVsmE660 challenge during four SIV vaccine trials carried out previously at the Yerkes National Primate Research Center (designated M2, M11, M12, and M15). All trials utilized SIVmac239 immunogens and a repeated, low-dose SIVsmE660 mucosal challenge using equivalent animal infectious doses in rhesus macaques. Immunized and control animals in the M11 and M15 trials were challenged 12 times intrarectally, while animals in the M2 and M12 trials were challenged 12 times intravaginally. The M11 trial was conducted by Rama Amara and consisted of three vaccine groups: DNA prime/MVA boost (DDMM), DNA prime with GM-CSF adjuvant/MVA boost (D1D2/MM), and three MVA immunizations (MMM). An additional arm was later included, designated M12, which consisted of DNA prime with CD40 ligand adjuvant followed by MVA boost (D1D2/MM/MM). The M2 trial was led by Bali Pulendran and consisted of three gp140 protein or virus-like particle (VLP) immunizations delivered with alum (protein+alum or VLP+alum) or nanoparticles (NP) (monophosphoryl lipid A and resiquimod) (protein+NP or VLP+NP) as adjuvants. The M15 trial was conducted by Rama Amara and was similar to M11 in that the vaccination arms were DNA prime, with or without the GM-CSF adjuvant, with two MVA boosts.

SGA and sequencing of SIVsmE660 env genes. Plasma collected at the second positive viral load test result during weekly challenges was used for 384-well single genome amplification (SGA) of SIVsmE660 env genes. Viral RNA was extracted from plasma using a QIAmp viral RNA kit according to the manufacturer’s instructions (Qiagen). Reverse transcription of viral RNA into cDNA was performed using a SuperScript III kit according to the manufacturer’s instructions (Invitrogen), with reverse primer SIVsm/macEnvR1 (5′-TGTAATAATCTCCCTTCCGTCGATCCC-3′). cDNA was diluted to result in <30% positive wells for SGA. First-round PCR was performed in a 15-μl volume using Phusion Hotstart II high-fidelity DNA polymerase (Thermo Scientific) with SIVsm/macEnvF1 (5′-CCTTCCCCCCATCCAGTACCAG-3′) and SIVsm/macEnvR1 (5′-TGTAATAATCTCCCTTCCGTCGATCCC-3′). Cycling conditions were 98°C for 2 min; 10 cycles of 95°C for 15 s, 54°C for 40 s, and 68°C for 4 min; 25 cycles of 95°C for 15 s, 54°C for 60 s, and 68°C for 4 min, adding 5 s to the extension per cycle; 72°C for 30 min; and 4°C hold. Second-round PCR was performed with the same enzyme in a 10-μl volume with 1 μl from the first round of PCR and SIVEnvF2 (5′-CACCTTATGACTAGAGGGACACCCCTTGGAAAGAGG-3′) and SIVEnvR2 (5′-ATGACATCTCTTTGCAATTGTAATAAATCCCTTCCAGTCCCCCC-3′). Cycling conditions were 98°C for 2 min; 30 cycles of 95°C for 15 s, 54°C for 60 s, and 72°C for 2.5 min; 72°C for 10 min; and 4°C hold. PCR amplicons were purified using a Qiagen PCR clean-up kit following the manufacturer’s instructions (Qiagen). Amplicons were sequenced with Beckman Coulter Genomics. The following primers were used: F1 (5′-ATCCATTTCAAGGATGATTGAGCCACTCC-3′), F2 (5′-AGGTTAAAAAGGGGACAAAAGGATAGAATA-3′), F3 (5′-TTTAGAGGAAGAGTTTCTTATGGCAAAAT-3′), and F4 (5′-AACACATTGCTGCTATTGTGGGACAAATAGA-3′) and R1 (5′-GCAACAGATACCCGCTTCCGGTGACCAATATC-3′), R2 (5′-CTCCTTCCGAGGGGAAATATACCATTCATT-3′), R3N (5′-TTGGAAATCTACATATTTTCTTGCTG-3′), and R4 (5′-TCATCTCTGCATCCACCATCAGTATT-3′).

Sequence analyses. Sequencer v5 was used to generate nucleotide sequence contigs, and sequences with evidence of mixed peaks were omitted from the analysis. Geneious v6.1.7 was used to translate nucleotide sequences, create alignments, and calculate pairwise diversity. Nucleotide alignments were exported from Geneious in FASTA format and used to generate PhyML (http://www.hiv.lanl.gov/content/sequence PHYML/ interface.html) and Rainbow (http://www.hiv.lanl.gov/content/seq uence/RAINBOWTREE/rainbowtree.html) trees. GenBank accession numbers for the VH2000 challenge stock sequences are FJ579014 to FJ579055, and those for the transmitted/founder (T/F) Env and NM2010 challenge stock sequences are KP734047 to KP735110 and KT288894.

Statistical analyses. Amino acid alignments were exported from Geneious in FASTA format for Sequence Harmony comparisons (http://www.ibi.vu.nl/programs/shmrwww/) and for generation of probability Weblogos (http://weblogo.threeplusone.com/create.cgi). Fisher’s exact tests were calculated via Prism 6 for Mac OS X.

Nucleotide sequence accession numbers. The NM2010 challenge stock sequences, M2 and M15 T/F Env sequences, and additional M11 and M12 T/F Env sequences are available in GenBank under accession numbers KT288416 to KT288894.

RESULTS

SIVsmE660 challenge stocks used for intrarectal and intravagi nal challenges were genetically indistinguishable. The M11, M12, M15, and M2 trials were preclinical SIVmac239-based immunization studies carried out at the Yerkes National Primate Research Center in rhesus macaques (16–18; unpublished data). M11, M12, and M15 vaccine regimens delivered immunogens via two DNA primes followed by two heterologous MVA boosts (DDMM) at 8-week intervals, followed by SIVsmE660 challenge at 48 to 52 weeks (Fig. 1). These trials also tested GM-CSF (DNA+GM-CSF) or CD40L (DNA+CD40L) adjuvanted DNA primes or a homologous prime boost using only MVA. The M2 trial delivered SIVmac239 Gag p55 and Env gp140 (protein) or virus-like particle (VLP) immunizations with alum or nanoparticles (NP) containing Toll-like receptor ligands (Fig. 1), followed by challenge at 38 weeks. Two SIVsmE660 challenge stocks were used; VH2000 was provided by Vanessa Hirsch and used for intrarectal challenges in M11 and M12, and NM2010 was provided by Nancy Miller and used for intravaginal challenges in M15 and M2. Ranajit Pal at ABL prepared NM2010 using the same seed stock (E660/PT71) and culture procedures (growth in pigtail ma-
Caque peripheral blood mononuclear cells [PBMC]) used to derive VH2000. To determine the genetic similarity of the two SIVsmE660 challenge stocks, we analyzed 37 SGA Env sequences (provided by Brandon Keele) from the VH2000 challenge stock and 59 SGA Env sequences from the NM2010 challenge stock. The sequences from the two stocks cameled on a PhyML phylogenetic tree (Fig. 2), and the maximum pairwise nucleotide difference was 3%. At the amino acid level, there were no signatures detected by Sequence Harmony analysis that could differentiate the two stocks using a Z-score of $-3$ as the threshold for significance (data not shown) (35). Based on the common lineage of the two stocks and their high genetic similarity, we combined the sequences to investigate viral determinants of vaccine breakthrough or transmission.

No vaccination breakthrough signature was identified in Env. To investigate potential sequence signatures in vaccination breakthrough, we analyzed an average of 10 ± 4 SGA T/F Env sequences from 49 immunized monkeys and 15 unvaccinated control monkeys (Fig. 1). The T/F Env sequences did not cluster according to vaccination group or status or with particular challenge stock sequences when aligned and analyzed phylogenetically (see Fig. S1 in the supplemental material), which is consistent with our previous analysis (14). The sequence analysis also revealed that a small minority of the animals (3 of 64) was clearly infected with more than one variant; however, we did not perform exhaustive sequencing to identify minor variants, as this was not the focus of our analysis. To explore the possibility of a vaccine sieving effect, the T/F Env sequences from the 49 vaccinated animals were compared against those from the 15 unvaccinated controls using Sequence Harmony (35). This analysis did not reveal significant differences (Z-score $<-3$) at any amino acid position that could be associated with vaccine breakthrough (data not shown).

Signature residues in gp120 C1 are associated with mucosal transmission. In the absence of a vaccine breakthrough signature,
indicating that these evidence for enhanced Z-score of less consistent with numbering in references 15, 36, and 37) achieved a the gp120 C1 region (based on SIVmac239 numbering and con- 67) to search for signatures that might favor mucosal transmission against all T/F Env sequences regardless of vaccination status (see Table S1 in the supplemental material), as illustrated in Fig. 4A and B, and thus no evidence for enhanced selection of I-A-K-N variants during vac- cination breakthrough in the experiments analyzed here.

we compared the combined challenge stock sequences (n = 96) against all T/F Env sequences regardless of vaccination status (n = 67) to search for signatures that might favor mucosal transmission of SIVsmE660. Four amino acid positions (23, 45, 47, and 70) in the gp120 C1 region (based on SIVmac239 numbering and consistent with numbering in references 15, 36, and 37) achieved a Z-score of less than −3 in Sequence Harmony (Fig. 3A and B), indicating that these residues could be associated with a selective advantage during mucosal transmission of SIVsmE660. The four signature positions were dominated by Val-Thr-Arg-Ser (V-T-R-S) in the SIVsmE660 challenge stocks (Fig. 3A). Using an independent Fisher exact test for each of the four positions, we found significantly higher frequencies of A-K-N at these positions in the T/F Envs than in the challenge stocks (Fig. 3B; also see Table S1 in the supplemental material) (P ≤ 0.002), while enrichment of I at position 23 was less significant (P = 0.017). This result is consistent with a recent meta-analysis performed by Gonzalez et al. (37).

As described above, Sequence Harmony analysis did not identify any signature residues associated with vaccination breakthrough; however, another study had reported that I-A-K-N was favored over V-T-R-S in breakthrough infections of SIVmac239 Env-vaccinated rhesus macaques (15). We therefore compared the frequencies of amino acids at these positions in the 51 vaccinated and 16 control T/F Envs using a Fisher exact test. There was no significant difference between the two (see Table S1 in the supplemental material), as illustrated in Fig. 4A and B, and thus no evidence for enhanced selection of I-A-K-N variants during vaccination breakthrough in the experiments analyzed here.

A Fisher exact test was also used to compare amino acid frequencies at positions 23, 45, 47, and 70 between animals infected intrarectally and intravaginally. There were no statistically significant differences at any of the signature positions between T/F Envs from these two groups (Fig. 4C and D; also see Table S1 in the supplemental material). A similar approach was used to investigate whether an association of amino acids at positions 23, 45, 47, and 70 with the number of challenges required to transmit infection (low, 1 to 6 doses; high, 7 to 12 doses) or TRIM5α encoded host susceptibility to infection. Neither of these was statistically significantly different in terms of the frequencies of I-A-K-N (see Table S1).

I-A-K-N restores transmission fitness to SIVsmE660. Overall, the data suggest that while V-T-R-S variants transmit across the mucosa, they do so at a lower frequency than expected by their dominance in the challenge stocks. Gonzalez et al. recently hypothe- sized that I-A-K-N variants of SIVsmE660 have higher transmission fitness than those carrying V-T-R-S (37). To explore this hypothesis with our data, we used challenge stock and T/F Env sequences to calculate the relative transmission fitness (W) of the V-T-R-S and I-A-K-N genotypes and all other genotypes combined (38). W_{IAKN} (Transmitted_{IAKN}/Stock_{IAKN} 2.5) was approximately 5-fold greater than W_{VTRS} (Transmitted_{VTRS}/Stock_{VTRS} 0.48) and 2-fold greater than W_{Others} (Transmitted_{Others}/Stock_{Others} 1.2) by these calculations. We used these values to extrapolate population compositions after an additional nine transmission cycles (Fig. 5A). By transmission cycle 3, I-A-K-N variants are predicted to overtake V-T-R-S and other variants as the majority genotype (60%). By transmission cycle 10, I-A-K-N virtually extinguished V-T-R-S and the other genotypes, making up >99% of the population. Thus, according to our extrapolations from a single infection cycle, I-A-K-N provides a strong advantage for in vivo SIVsmE660 mucosal transmission.
To gain further insight into the advantageous nature of the I-A-K-N genotype and determine whether it is also more fit for aspects of viral replication other than transmission, we investigated the amino acid compositions at positions 23, 45, 47, and 70 in the HIV-2/SIVsmm lineage, excluding SIVsmE660-derived sequences. One hundred twelve HIV-2/SIVsmm reference Env sequences were downloaded from the LANL HIV sequence database (http://www.hiv.lanl.gov/content/sequence/NEWALIGN/ign.html) and examined alongside the challenge stock and T/F Env sequences. Interestingly, V and I, the only residues found at position 23 in SIVsmE660, occurred in only a small minority of the HIV-2/SIVsmm reference Env sequences, which displayed high entropy at this position (Fig. 5B). For the other three positions, A-K-N were highly conserved in the HIV-2/SIVsmm reference Env sequences, reflecting their increased frequency in the T/F Env. An analysis of the analogous residues in more than 5,000 Env sequences from the HIV-1/SIVcpz lineage downloaded from the LANL database followed a pattern similar to that of the HIV-2/SIVsmm sequences (Fig. 6), in this case, with A-E-N dominating. Thus, diversifying selection at position 23 and high conservation at positions 45, 47, and 70 occur in the great majority of natural SIV and HIV variants.

**DISCUSSION**

This study and two other recent publications have identified amino acids 23, 45, 47, and 70 in the C1 region of gp120 as important during mucosal SIVsmE660 challenge of rhesus macaques (15, 37). Roederer et al. attributed an enrichment of I-A-K-N to neutralization resistance (emphasizing A-K) that facilitated vaccine breakthrough infection, while a retrospective analysis of multiple independent trials carried out by Gonzalez et al. hypothesized that the enrichment of I-A-K-N was the result of increased transmission fitness and was independent of preexisting immunity. Due to the purposeful design of the M11, M12, M2, and M15 vaccine trials, we recognized a unique opportunity to explore both of these hypotheses.

Regarding the association between positions 45 and 47 and resistance to neutralizing antibody, in-depth analysis of prechallenge antibody responses against the autologous breakthrough T/F Env variants in the M11 and M12 trials yielded results that were not consistent with those of Roederer et al. in that both A-K and T-R variants were susceptible to heterologous and autologous neutralization (14). In the M11/M12 study, neither T-R nor A-K predicted susceptibility to heterologous or vaccine-induced neutralizing antibody responses, and breakthrough variants from vaccinated animals were not enriched for A-K over the T/F Env from unvaccinated control animals. The Sequence Harmony signature analysis presented here, performed on a larger number of sequences from the M2, M11, M12, and M15 trials, also failed to identify any amino acid signatures in T/F Envs from vaccinated versus control animals. Furthermore, results from direct Fisher’s exact tests of amino acids at positions 23, 45, 47, and 70 between these two groups were not significant. However, the results of our current study are in agreement with those of Gonzalez et al. and suggest that I-A-K-N is advantageous for mucosal transmission regardless of whether vaccine-induced antibodies are present or absent (37). It is important to note that the trials analyzed here used entirely different vaccination strategies and challenge stocks from the Roederer et al. study, which could explain the inconsistencies between our observations. Nevertheless, our findings do argue against positions 45 and 47 being a universal marker for neutralization resistance (14, 36, 37, 39, 40).

Even if amino acid changes at positions 45 and 47 alter SIVsmE660 neutralization sensitivity, the ability to evade host antibody responses cannot be the dominant selective pressure on transmission. Roederer et al. suggested that stepwise mutations from T-R to T-K, A-R, and A-K at positions 45 to 47 resulted in increased neutralization resistance (T-R < T-K < A-R < A-K) (15). However, we did not isolate any sequences with T-K at positions 45 and 47 in the 96 challenge stock sequences or among the 67 T/F Env sequences. This could be due to the fact that SGA, not deep sequencing, was used here. Even so, a search of all SIVsmm Env sequences available in the LANL HIV sequence database for those that contain the T-K motif CTTKN (amino acid positions 44 to 48) identified only 22 sequences, 21 of which were from a single SIVsmE660-infected monkey (15). Thus, even if the T-K intermediate confers an increase in neutralization resistance, there ap-
pears to be selective pressure against this amino acid combination during in vitro propagation of viral stocks, in animal transmission models, and in the natural animal host environment. The reasons underlying selection for or against certain residues in the SIVsmE660 gp120 C1 region could shed light upon this region’s potential contributions to viral fitness.

The question remains as to why I-A-K-N variants of SIVsmE660 are favored during mucosal transmission and what they can teach us about HIV-1 transmission and protection. These variants could be more efficient than others at any number of steps, including crossing the mucosal epithelium, infecting a susceptible CD4+ target cell, evading innate immunity, expanding locally to infect other CD4+ target cells, or disseminating out of the mucosa (41). However, residues A-K-N are so highly conserved in HIV-2/SIVsm isolates that only the atypical amino acid composition of the SIVsmE660 challenge stock Env quasispecies allowed this selection event to be observed. It might then be tempting to alter the genetic composition of SIV or simian-human immunodeficiency virus (SHIV) challenge stocks beyond what is found in nature to enhance our ability to detect informative sites and learn more about the transmission process. However, this approach runs the risk of generating a viral quasispecies that is no longer representative of circulating HIV-1 variants. If these populations deviate too far from nature, the dominant selective pressure would be adaptation back to the animal host, obfuscating the vaccine-mediated protection that these models are set up to test. Indeed, V-T-R-S, despite being the dominant genotype in the SIVsmE660 quasispecies, is disadvantageous not only for transmission but also during natural SIV infection. In considering the nearly invariant nature of the analogous residues in HIV-1 (A-E-N) (Fig. 6), it becomes clear that the chance of a vaccinated individual encountering an HIV-1 variant as atypical SIVsmE660 V-T-R-S during a transmission event in the human host is extremely low. It is also improbable that this signature would be easily identified in acute versus chronic or donor versus recipient comparisons of HIV-1 Env sequences, due to the invariant nature of the residues. Our results, combined with those of Gonzalez et al. (37), should prompt a careful consideration of the authenticity of nonhuman primate challenge models, wherein protection may be influenced by the effects of reinstating in vivo fitness, a phenomenon which does not occur in natural HIV-1 infection.

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