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Link between Primate Lentiviral Coreceptor Usage and Nef Function

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SUMMARY

Simian immunodeficiency virus (SIVsmm) infection of sooty mangabeys (Cercocebus atys) is characterized by stable CD4⁺ T cell counts despite high plasma levels of CCR5-tropic viruses. However, in rare instances, SIVsmm acquires CXCR4 coreceptor tropism and causes severe CD4⁺ T cell depletion, albeit without clinical signs of immunodeficiency. Here, we show that CXCR4-tropic SIVsmm strains lost their ability to downmodulate TCR-CD3 by evolving unusual Nef mutations that initially reduced (I132V) and subsequently disrupted (I123L and L146F) interaction with the CD3 ξ chain. This coevolution of Env and Nef function suggests that CD3 downmodulation is advantageous for viral replication in activated CCR5⁺ memory T cells, but not in resting naive CXCR4⁺ T cells that have not yet undergone TCR-CD3-mediated stimulation. This may explain why HIV-1, which generally lacks the CD3 downmodulation function, commonly switches to CXCR4 usage, whereas this is extremely rare for SIV strains that have retained this Nef activity.

INTRODUCTION

The great majority of sooty mangabeys (SMs) that are naturally infected with simian immunodeficiency virus (SIVsmm) maintain stable CD4⁺ T cell counts despite high levels of viral replication (reviewed in Mir et al., 2011; Chahroudi et al., 2012). In very rare instances, however, SIVsmm infection can cause severe CD4⁺ T cell loss, albeit in the absence of clinical disease (Milush et al., 2007; Sumpter et al., 2007). In-depth analysis of two animals with almost complete loss of CD4⁺ T cells revealed the emergence of SIVsmm variants that switched from CCR5 (R5) to CXCR4 (X4) coreceptor usage (Milush et al., 2007, 2011). In HIV-1-infected humans, such an R5-to-X4 coreceptor switch is observed in about 50% of late-stage patients and associated with rapid CD4⁺ T cell depletion and accelerated disease progression (Connor et al., 1997; Xiao et al., 1998). In CD4low sooty mangabeys, CD4⁺ T cells are also lost, but there is no generalized chronic immune activation and no disease. These unusual CD4low SIVsmm infections are thus a useful model to study the interplay between viral X4 tropism, CD4⁺ T cell loss, and generalized immune activation.

In the present study, we examined the genetic and functional evolution of env and nef genes in two experimentally SIV-infected SMs with severe CD4⁺ T cell depletion and three additional SMs that were inoculated with plasma from one of these CD4low animals. We hypothesized that Env and Nef function might be linked, because coreceptor tropism determines the subsets of CD4⁺ T cells that become infected and the accessory Nef manipulates them in multiple ways to promote efficient viral replication (reviewed in Kirchhoff et al., 2008). In particular, we hypothesized that SIVsmm Nef, which unlike HIV-1 Nef is able to suppress T cell activation and apoptosis through TCR-CD3 downregulation (Schindler et al., 2006, 2008; Arhel et al., 2009), may contribute to the lack of immune activation and the absence of disease progression in these CD4low animals. Surprisingly, this was not the case. Instead, we found that many nef alleles of X4 SIVsmm strains were unable to downmodulate TCR-CD3 and to suppress T cell activation, while maintaining all other functions and even gaining activity in downmodulating CXCR4. Thus, the lack of the TCR-CD3 downregulation function, which is a hallmark of HIV-1 (Schindler et al., 2006, 2008; Schmökel et al., 2009), was not associated with increased immune activation in CD4low mangabeys. Instead, this adaptation may have been required for SIVsmm replication in naive CXCR4⁺ T cells that usually show a resting phenotype and (unlike memory CCR5⁺ T cells) have not undergone TCR-CD3 stimulation prior to virus infection. Thus, the loss of the CD3 modulation function of Nef may promote CXCR4 tropism and associated increased pathogenesis of HIV-1.
RESULTS

Sequence Evolution of env and nef in SIVsmm-Infected SMs with Severe CD4+ T Cell Loss

To study the genetic and functional evolution of Env and Nef in two CD4low sooty mangabeys (SM1 and SM2), we amplified a 3.3 kb SIVsmm env-nef-LTR fragment (Figure 1A) by single-genome amplification (SGA) from 17 plasma samples obtained prior to, during, and after CD4 depletion (Figure 1B). In addition, we amplified the same fragment from nine plasma samples obtained at 2, 32, and 154 weeks postinfection (wpi) from three sooty mangabeys (SM7, SM8, and SM9) that were inoculated intravenously with 1.5 ml of plasma from one of the CD4low mangabey (SM2) at 304 wpi and also exhibited rapid CD4+ T cell loss (Milush et al., 2007). SGA was employed to eliminate PCR-induced artifacts and to allow assessment of linkages between Env and Nef function. In all five animals, severe CD4+ T cell loss was associated with a reduction in viral RNA loads by at least two orders of magnitude (Figure 1C).

A total of 211 env-nef-LTR sequences were analyzed from the five SMs. Animals SM1 and SM2 were infected in October 2001 with plasma from a chronically infected sooty mangabey (FQi), which had acquired its infection in the wild (Milush et al., 2007). Sequences obtained early after infection (12 wpi) from these SMs were closely related (Figure 2, bottom) but subsequently diversified, forming animal and time-point-specific clusters (Figure S1). The 31 sequences obtained from SM2 at 304 wpi (the time point of plasma sampling for i.v. injection of SM7,

Figure 1. Longitudinal Analysis of SIVsmm env-nef Sequences
(A) Single-genome amplification was used to generate env-nef fragments from plasma samples of the SIVsmm-infected SMs indicated.
(B and C) Peripheral CD4+ T cell levels (B) and viral loads (C) during the course of SIVsmm infection of the indicated animals. In SM1 and SM2, the CD4 levels declined to below 80 per µl of blood at 63 (SM1) and 79 (SM2) wpi, respectively. The loss of CD4+ T cells was associated with declining plasma viral loads in all five animals. The arrows above the curves indicate the time points of plasma sampling for SGA analysis. The numbers next to the arrows indicate the predominant charge of the V3 loop amino acid sequences amplified at the corresponding time points. The time point of plasma sampling for passage of the X4/multitropic SIVsmm from SM2 to SM7, SM8, and SM9 is indicated.
See also Figure S1.
Figure 2. Evolutionary Analysis of SIVsmm env-nef Sequences

A phylogenetic tree was constructed using the neighbor-joining (NJ) method. The scale bar represents 0.01 substitutions per site. Sequences are color-coded based on their animal origin. Two clusters that include nef alleles with amino acid substitutions I123L and L146F are indicated. Stars indicate SGA sequences used to examine the function of Env and Nef (green) or Nef only (black). The length of the amplicons was about 3.7 kb. See also Table S1.
SM8, and SM9) formed two lineages, referred to as “A” and “B” herein after (Figure 2, upper). Notably, all 30 sequences from SM7, SM8, and SM9 at 2 wpi were closely related to one another and interspersed with the cluster A SM2-304 sequences. In contrast, sequences derived from SM7 and SM8 at 32 or 154 wpi were more closely related to cluster B SM2-304 sequences (Figure 2). Thus, following passage of SIVsmm from SM2 to three additional animals, different members of the viral quasispecies predominated during early (cluster A) and chronic (cluster B) infection.

**CD4⁺ T Cell Loss Correlates with Increased CXCR4 Coreceptor Usage**

Previous studies showed that concomitant with the CD4⁺ T cell depletion in SM1 and SM2 viral variants emerged that exhibited an expanded coreceptor tropism, using CCR5, CXCR4, and CCR8 for entry (Milush et al., 2007). However, coreceptor usage was examined only for three time points and only in a high sensitive cell–cell fusion assay. We thus examined the coreceptor tropism of viruses infecting SM1 and SM2 in greater detail. A total of 30 env alleles from eight different time points (indicated in Figures 1 and 2) were selected for functional analyses. To generate virions containing these SIVsmm Envvs, we cotransfected 293T cells with vectors expressing the respective Env proteins and an env-defective HIV-1 luciferase reporter construct. Infection of TZM-bl cells, which express CD4 as well as R5 and X4, in the presence of X4- and R5-specific entry inhibitors (AMD3100 and SCH-D, respectively) confirmed that at 12 wpi viruses from both SM1 and SM2 used mainly R5 (Figure 3A). In contrast, Envvs obtained after the loss of CD4⁺ T cells (i.e., at 107, 304, and 340 wpi) preferentially used X4. Interestingly, Envvs obtained during CD4⁺ T cell decline (i.e., at 47 and 51 wpi) were blocked only poorly by both R5 and X4 inhibitors, suggesting alternative coreceptor usage (Figure 3A; data not shown).

To determine which alternative coreceptors were utilized, we exposed GHOST cells stably expressing CD4 and different G protein-coupled receptor HIV-1 luciferase reporter constructs pseudotyped with the SIVsmm Env proteins of interest. We found that at 12 wpi both SM1 and SM2 Envvs used predominantly R5 and GPR15 (BOB) and to a lesser extent CXCR6 (STRL33, Bonzo) for viral entry (Figures 3B and 3C). At subsequent time points, SIVsmm SM1 evolved to utilize mainly X4 (≥ 107 wpi; Figures 3B and 3D), whereas SIVsmm SM2 broadened its coreceptor usage to efficiently utilize X4 as well as other coreceptors (Figures 3B, 3D, and 3E). Notably, this expanded coreceptor tropism was maintained after passage of virus from SM2 to SM7, SM8, and SM9 (Figures 3B and 3F), although Envvs from these newly infected SMs mainly used CXCR4 at 154 wpi (data not shown). The coreceptor switch/ expansion phase in both SM1 and SM2 (47–79 wpi in SM1 and 47 to 107 wpi in SM2) coincided with the acquisition of several amino acid changes (Figure S2), including changes in the V3 loop region of Env. However, amino acid substitutions that increased its net charge (from +4 to +7) emerged only later during infection and were associated with increased efficacy of X4-mediated viral entry (Figure 3; Table S1). The transitional Envvs were generally less infectious than Envvs obtained early or later during infection (Figure 3A; data not shown). Thus, the coreceptor switch from R5 to X4 in both SM1 and SM2 required complex changes in Env that were associated with reduced infectivity compared to the early R5- and late X4-tropic SIVsmm strains.

**Loss of Nef-Mediated Modulation of TCR-CD3 and Suppression of T Cell Activation**

To determine whether specific features of Nef were associated with the loss of CD4⁺ T cells, we analyzed the function of a total of 61 nef alleles obtained at different time points from all five animals (Figure S3). As reported previously (Schindler et al., 2006), nef alleles were cloned into an HIV-1 NL4-3-based IRES-eGFP proviral vector coexpressing Nef and eGFP from a bicistronic RNA. Virus stocks were generated by cotransfection of 293T cells with the proviral constructs and a vector expressing the VSV-G envelope protein to transduce peripheral blood mononuclear cells (PBMCs) with high efficacy for flow cytometric analyses. These analyses showed that Nef-mediated downmodulation of CD4 and MHC-I did not change significantly throughout the course of infection (Figures 4A and 4B). In contrast, nef alleles derived from SM1 and SM2 after the loss of CD4⁺ T cells exhibited a significant decline in CD3 downmodulation activity compared to those derived early during infection (Figure 4C). Four of eight nef alleles derived from SM2 at 304 wpi and all three nef genes obtained at 340 and 365 wpi were completely inactive in downmodulating CD3. This Nef function was also significantly reduced in viruses derived from SM1 at 340 and 365 wpi, although some marginal activity was retained (Figure 4C). The efficiency of Nef-mediated modulation of CD28 was higher in SM2 than in SM1, but most SM1 nef alleles from later time points (107–365 wpi) exhibited only marginal activity. Interestingly, Nef-mediated downmodulation of X4 increased significantly in viruses that also utilized this coreceptor (Figure 4E). When we grouped the SIVsmm constructs based on their coreceptor tropism, we noted that in both SM1 and SM2 X4 tropism was significantly associated with a loss of Nef-mediated downmodulation of TCR-CD3 and a gain of the CXCR4 modulation activity (Figure 4F). In SM1, X4 SIVsmm strains also lost the CD28 downmodulation function of Nef. Taken together, SIVsmm strains that were present early during infection and capable of TCR-CD3 modulation utilized mainly R5 in both SM1 and SM2, whereas those emerging late during infections and lacking this Nef function used predominantly X4 for entry into target cells (Figure 4G).

The SM2 304 wpi plasma sample used for inoculation of SM7, SM8, and SM9 contained a mixture of SIVsmm nef variants that did (cluster A, n = 17/31) or did not (cluster B, n = 14/31) downmodulate TCR-CD3. Interestingly, all nef alleles obtained from these three animals during acute infection (2 wpi) were active against TCR-CD3 (Figure 4C). In contrast, most nef alleles derived from SM7 at 32 and 154 lacked this nef function, whereas those obtained from SM8 and SM9 remained active at these later time points (Figure 4C). Other Nef functions, such as modulation of CD4, CXCR4, and MHC-I did not change significantly in the three newly infected animals during follow-up, whereas the efficiency of CD28 modulation generally decreased from 2 to 154 wpi (Figures 4A–4E). These data indicate that
Nef-mediated modulation of TCR-CD3 is required early in SIVsmm infection, but in certain cases becomes dispensable, or is even selected against, when the number of CD4+ target cells is low.

To determine whether the loss of Nef-mediated downmodulation of CD3 in SM1 and SM2 was associated with an increased responsiveness of infected T cells to stimulation, we measured the surface expression of CD69, a marker of early T cell activation in HIV-1-infected PHA-treated PBMC cultures. These analyses showed that infected T cells expressing Nef alleles obtained after the loss of CD4+ T cells also expressed increased levels of CD69 (Figure 4H). Next, we transduced Jurkat T cells containing the luciferase reporter gene under the control of an NFAT-dependent promoter (Fortin et al., 2004) with the proviral HIV-1 eGFP/Nef constructs and examined their responsiveness to activation. SIVsmm nef alleles obtained from SM1 and SM2
up to 107 wpi suppressed NFAT induction. In contrast, nef alleles obtained from both animals after 300 wpi rendered the T cells hyperresponsive to stimulation, similar to HIV-1 Nefs (Figure 4I). In agreement with previous data (Schindler et al., 2006; Khalid et al., 2012), suppression of CD69 expression and NFAT activation correlated significantly with the efficiency of Nef-mediated downmodulation of TCR-CD3 (Figure 4J). Finally, we also examined the ability of these nef alleles to enhance virion infectivity and to promote virus release by counteracting SM tetherin. We found that both of these Nef functions were preserved throughout the course of CD4<sup>+</sup> SIVsmm infection (Figure S4). Altogether, these results show that nef genes from late-stage X4 SIVsmm strains allowed efficient stimulation of virally infected T cells and thus had functional properties that were more reminiscent of nef alleles derived from HIV-1 than from R5 SIVsmm strains.

**Residues I123 and L146 Are Critical for CD3 Downmodulation by SIVsmm Nef**

We next examined which amino acid changes were responsible for the selective loss of the CD3 downmodulation function. Sequence alignments showed that active and inactive Nefs from SM2 differed by amino acid changes of I123L and L146F (Figure S3). These two substitutions were preceded by a substitution of I132V in SM2, and this change was also observed in most Nef alleles derived from SM1 at later time points (≥107 wpi). Notably, a naturally occurring amino acid variation at this position (I132T) specifically disrupts the CD3 downmodulation activity of HIV-2 Nef (Khalid et al., 2012). Comparison of the functional activity of SIVsmm Nefs differing in these residues showed that Nefs containing the I123L and L146F substitutions were inactive in CD3 downmodulation, whereas those containing the I132V change showed an intermediate phenotype (Figure S4). Notably, the I123L and L146F changes always occurred in combination with one another, but never together with the I132V substitution (Figure S3).

The temporal appearance of different nef substitutions suggested a timeline where Nef-mediated CD3 downmodulation was initially reduced by the I132V substitution and later disrupted entirely by the I123L and L146F substitutions. To confirm the role of the latter changes in TCR-CD3 downmodulation, we analyzed a set of mutant SIVsmm Nefs differing specifically in these two residues. Combined changes of I123L and L146F disrupted the CD3 downmodulation activity in early SIVsmm SM2 Nefs, whereas the reciprocal revertant changes restored this activity in late inactive Nefs (Figure 5B). The I123L and L146F changes in Nef also affected the responsiveness of virally infected cells to stimulation and the induction of apoptosis but had no significant effect on other Nef activities, such as modulation of CD4, CD28, CXCR4, and MHC-I or enhancement of virion infectivity (data not shown).

To further define the role of residues I123 and L146 in Nef-mediated CD3 modulation, we performed fluorescence-activated cell sorting (FACS)-based internalization assays in Jurkat T cells. Predictably (Swigut et al., 2003), a control SIVmac239 Nef was highly active (Figure 5B). The SIVsmm SM2 75w-L4 and mutant 304w-H1 L123I/F146L Nefs also accelerated internalization of TCR-CD3, whereas the 75w-L4 I123L/L146F and wild-type 304w-H1 Nefs had no significant effect (Figure 5C). To examine whether residues I123 and L146 directly affected the interaction of Nef with CD3ζ, we performed FRET and coimmunoprecipitation analyses. Our results showed that the SIVmac239 Nef as well as SIVsmm 75w-L4 and 304w-H1 L123I/F146L Nefs were effective in pulling down CD3ζ (Figure 5D) and resulted in significant FRET signals (Figure 5E). Consistent with their inability to downmodulate TCR-CD3, no interaction was detected for the 75w-L4 I123L/L146F and wild-type 304w-H1 Nefs (Figures 5D and 5E). Thus, changes in residues I123 and L146 affect the ability of SIVsmm Nef to interact with the CD3ζ chain.

Next, we examined the effect of the I123L and L146F substitutions on the ability of Nef to modulate the subcellular localization of TCR-CD3. For these experiments, we utilized a previously described fusion construct between the cytoplasmic tail of CD3ζ-GFP.
fluorescence was detected exclusively in intracellular perinuclear compartments in the presence of the SIVmac239, SIVsmm SM2 75w-L4, and 304-H1 L123I/F146L Nefs. Thus, the microscopy analyses confirmed that the I123L and L146F changes disrupt the ability of Nef to remove TCR-CD3 from the cells surface. These substitutions were present in a total of 35 nef alleles (14/31 SM2 304 wpi, 7/7 SM2 340/365 wpi; 5/5 SM7 32 wpi and 9/13 SM7 154 wpi sequences) that formed a distinct phylogenetic cluster (Figure 2).

To better understand how the I123L/L146F changes affected Nef-mediated modulation of TCR-CD3, we examined the localization of these residues in the crystal structure of the SIVmac239 Nef core domain complexed with the first ITAM motif of the CD3ζ chain (Kim et al., 2010). SIVmac originated from SIVsmm and Nef is highly conserved between these viruses. We found that residues I123 and L146 are located on two central α helices, which are located within the core domain of Nef (Figure 6A). These two helices are kept in place by the two gatekeeper residues F122 and Y145 (F94 and W117 in HIV-1 Nef), which divide the intermediate groove into a recognition site for SH3 domains and a binding site for endocytic sorting motifs (Horenkamp et al., 2011). I123 and L146 oppose each other on these two helices at a minimal distance of 4.2 Å but are not directly surface accessible (Figures 6B and 6C). Mutation to 123L and 146F, which are similarly hydrophobic but much larger, are predicted to result in a much tighter interaction between these residues (Figure 6C). Thus, the I123L and L146F changes may shift the position of the two helices relative to each other and thus alter the size and structure of the associated binding pocket in such a way that it becomes inaccessible for the CD3ζ chain but still accommodates the cytoplasmic domain of CD4, which binds the same hydrophobic crevice (Grzesiek et al., 1996).

**DISCUSSION**

In the present study, we show that SIVsmm strains that switch from R5 to X4 coreceptor usage and cause severe CD4+ T cell
depletion gain activity in Nef-mediated downmodulation of X4 but lose their ability to remove TCR-CD3 from the cell surface (summarized in Figure S5). In one animal studied in detail, this loss of function was due to two amino acid substitutions of I123L and L148F in the core region of Nef that specifically disrupted its interaction with the TCR-CD3-z chain. Like X4-tropic HIV-1 strains—but unlike to common SIVsmm strains—the emerging X4-using SIVsmm variants were unable to prevent T cell activation. However, none of the five animals infected with these SIVsmm strains showed generalized immune activation or signs of disease (Milush et al., 2007, 2011). Thus, in sooty mangabeys chronic inflammation and AIDS pathogenesis requires more than CD4+ T cell loss, CXCR4 tropism and loss of the Nef-mediated TCR-CD3 downregulation function. Unexpectedly, this animal model provides evidence that the latter represents an adaptation for efficient viral replication in CXCR4+ T cells.

Env and Nef Coevolution

CCR5 is mainly expressed on activated memory CD4+ T cells, particularly effector-memory CD4+ T cells, whereas CXCR4 is predominantly expressed on quiescent or resting naive CD4+ T cells. The viral Env glycoprotein determines which of these subsets of T cells are infected and the accessory Nef protein coreceptor utilization by rare SIVsmm variants also correlated with the efficiency of Nef-mediated downmodulation of this chemokine receptor further underscores this linkage.

Another change in Nef function that correlated with the acquisition of X4 tropism by SIVsmm was a loss of the CD3 downmodulation activity (Figure 4C). We have previously shown that this Nef function is highly conserved in HIV-2 and most lineages of SIV, but absent in HIV-1 and its vpu containing SIV precursors (Schindler et al., 2006, 2008; Schmokel et al., 2009; Khalid et al., 2012). CXCR4 is mainly present on naïve T cells that have not yet encountered their cognate antigen and usually exhibit a quiescent or resting phenotype that is not conducive to efficient viral replication. It is thus possible that SIVsmm tropism for X4+ T cells may select for nef alleles that lack the CD3 modulation function because TCR-CD3 stimulation of this largely resting T cell subset by APCs may be required for effective viral replication (schematically outlined in Figure S6).

In contrast, activated CCR5+ memory CD4+ T cells likely allow effective viral replication without further stimulation. Notably, in one animal (SM2) X4-tropic SIVsmm did not lose the TCR-CD3 downmodulation function entirely but may have compensated by also losing the CD28 downmodulation function (Figure 4F). The TCR-CD3 complex and the CD28 costimulatory factor modulates the interaction of these cells with antigen-presenting cells (APCs) and their responsiveness to stimulation by downmodulation of CD4, CD28, CXCR4, and (in the case of HIV-2 and most SIVs) TCR-CD3 surface expression. Thus, it is not surprising that Env and Nef function are linked, although this has not previously been reported. The fact that X4
subset of CD4+ T cells and to evolve the ability to replicate in X4+ naive CD4+ T cells because it generally lacks the CD3 downmodulation function of Nef. Instead, the great majority of naturally infected SMs maintain high levels of CD4+ T cells. In this stable host environment, activated R5+ effector-memory CD4+ T cells, which are the preferential targets for virus infection (Paiardini et al., 2011), are available throughout the course of infection and SIVsmm thus remains R5-tropic and active in CD3 downmodulation (schematically shown in Figure 7A). Even in the ~10% of SMs that eventually experience a significant loss of CD4+ T cells after decades of infection, this loss occurs gradually and is not associated with viral X4 coreceptor usage (Sumpter et al., 2007; Taaffe et al., 2010). It is thus surprising that after i.v. inoculation with plasma from SM FQi, SIVsmm variants capable of utilizing X4 emerged in two out of 14 animals (Silvestri et al., 2005; Milush et al., 2007; Gordon et al., 2008; Bosinger et al., 2009). Given that all EnvS obtained from SM1 and SM2 early during infection utilized R5 and GPR15 (Figure 3), it is unlikely that X4 SIVsmm variants were present in the virus inoculum. Furthermore, there was no evidence for X4 viral strains in the source animal FQi, which reached an age of 22 years and died of age-related causes. Instead, the emergence of these unusual variants may have been due to the virus dose and/or the route of infection. Like in X4-tropic HIV-1 infection, the ability of SIVsmm to infect the large pool of naive X4+ T cells was associated with rapid and severe CD4+ T cell depletion (Figure 1B). Nonetheless, SM1 and SM2 did not develop signs of clinical immunodeficiency throughout 13 years of follow-up. A possible explanation for this is that SMs are equipped with T cells that are capable of performing helper functions but lack the CD4 receptor and are thus resistant to SIVsmm infection (Milush et al., 2011; Sundaravaradan et al., 2013).

Whereas X4 coreceptor usage is very rare in SIVsmm infection, it is observed in about 50% of late stage HIV-1-infected individuals and associated with rapid CD4+ T cell decline and accelerated progression to AIDS (Moore et al., 2004). It is currently not understood which mechanisms drive X4 coreceptor usage and why this phenomenon is more common in HIV-1 than in SIV infections. Our results provide a potential explanation for the latter. Nef alleles lacking the CD3 downmodulation function emerged only after the drastic loss of CD4+ T cells and only in the context of X4-tropic SIVsmm variants (Figure S5). Thus, lack of CD3 modulation was clearly not a cause of impaired CD4+ T cell homeostasis but may have emerged as a consequence of the coreceptor switch to X4. The specific loss of the CD3 downmodulation function during the later stages of X4 SIVsmm infection in both CD4low SMs suggests that lack of this Nef function may be advantageous for viral replication in X4+ naive T cells that have not yet undergone TCR-CD3 stimulation. We have previously shown that modulation of CD3 by Nef disrupts TCR-CD3-mediated stimulation of virally infected CD4+ T cells by APCs (Arhel et al., 2009). Thus, HIV-1 and the late stage X4 SIVsmm strains analyzed in the present study may be better equipped for replication in resting naive CXCR4+ T cells because they do not prevent activation of their target cells to levels required for efficient viral gene expression. It has previously been suggested that an increased number of activated CXCR4+ T cells renders HIV-1-infected patients more
likely to switch their coreceptor usage from R5 to X4 (Weinberger and Perelson, 2011). Our results are consistent with this and suggest that X4-tropic SIVsmm strains may evolve to support the generation of activated T cells by rendering them responsive to TCR-CD3-mediated stimulation.

HIV-2 originated from SIVsmm and both are closely related. HIV-2 also shows promiscuity of coreceptor utilization but uses CXCR4 more frequently than SIVsmm, albeit less commonly than HIV-1 (Mörner et al., 1999; Calado et al., 2010). Similarly to HIV-1 infection, the acquisition of CXCR4 coreceptor usage by HIV-2 is only observed in symptomatic AIDS patients with low numbers of CD4+ T cells (Guillon et al., 1998; Calado et al., 2010). Furthermore, low CD4+ T cell counts are associated with reduced efficacy of TCR-CD3 downmodulation by Nef in viremic HIV-2-infected individuals (Khalid et al., 2012). These results are in agreement with a link between CXCR4 usage and Nef-mediated downmodulation of TCR-CD3 in HIV-2 infection but further studies are required to obtain definitive proof.

Nef-Mediated Suppression of T Cell Activation Is Advantageous during Acute Infection

The virus stock used to infect animals SM7, SM8, and SM9 contained two different viral lineages (“A” and “B,” Figure 2) that exhibited the same coreceptor tropism (Figure 2E) but differed in their TCR-CD3 modulation function of Nef. Approximately half of the viral sequences (14/31) amplified from the viral inoculum contained the I123L and L146M substitutions in Nef that disrupt this function. Nonetheless, all 31 nef alleles derived from SM7, SM8, and SM9 during acute infection at 2 wpi lacked these changes and modulated CD3. These data strongly suggest that downmodulation of CD3 is required during the earliest stages of infection. Although X4 tropism was maintained in all three animals after viral passage, the transmitted SIVsmm strains had a broad coreceptor tropism that included R5 (Figure 3). Activated CD4+ T cells are readily available during acute infection and Nef-mediated downmodulation of TCR-CD3 may thus be beneficial for initial viral spread in lymphatic tissues. The finding that SIVsmm variants lacking the CD3 downmodulation function did not emerge in all SMs after severe CD4+ T cell depletion suggests that infection of resting naive CXCR4+ T cells may represent a suboptimal niche for SIVsmm replication in the absence of other CD4+ target T cell populations.

Two Types of Changes in the Core Domain of Nef Disrupt Downmodulation of TCR-CD3

Previous mutagenesis experiments have failed to selectively eliminate CD3 downmodulation without impairing other Nef functions. In fact, the only other mutation (I132T) known to selectively disrupt the interaction of Nef with the CD3ζ chain was identified in a variant of HIV-2 (Khalid et al., 2012). Interestingly, a change at the same position (I132V) preceded the emergence of the I123L and L146F substitutions in SM2 and also became predominant in SM1 later during infection (Figure S3). The I132V change reduced the effect of Nef on CD3 but did not disrupt it entirely (Figure 5A). I132 is located at the bottom of a hydrophobic crevice in Nef (Kim et al., 2010), and the change to a polar threonine disrupts the hydrophobic contacts formed between Nef L75 of the YxxL ITAM motif in the ζ chain of CD3 (Khalid et al., 2012). In contrast, residues I123 and L146 do not interact directly with the ζ chain of CD3. However, they determine the relative position of the two large alpha helices that border the ζ chain recognition site. It is thus possible that the big, γ-branched head group of L123 and the large aromatic F146 reposition the hydrophobic crevice of the recognition site in such way that the ζ chain of CD3 is no longer accommodated, whereas binding to the endocytic sorting motif of CD4 as well as all other Nef functions remain intact. Thus, the virus may fine-tune Nef function and thus the responsiveness of virally infected T cells to stimulation by the acquisition of subtle changes that cooperate to alter the interaction of Nef with specific cellular targets in an indirect allosteric manner and keep all essential surface properties intact.

CONCLUSIONS

In this paper, we show that the rare acquisition of X4 tropism by SIVsmm and the concomitant severe depletion of CD4+ T cells are associated with changes in Nef function, i.e., increased downmodulation of CD4 and loss of the ability to remove TCR-CD3 from the cell surface to suppress T cell activation. This coevolution suggests a functional linkage between Env and Nef that has previously gone unrecognized. The lack of Nef-mediated downmodulation of CD3 may enhance viral replication fitness in naive resting CXCR4+ T cells, because it allows their TCR-CD3-mediated stimulation by APCs to increase virus production. Thus, lack of CD3 downmodulation may be one reason why X4 tropism is more common in HIV-1 than in SIVsmm infection.

EXPERIMENTAL PROCEDURES

Animals and Viral Loads

Animals were infected and plasma viral loads and CD4+ T cell counts were determined as previously described (Milush et al., 2007). Local animal care and use committee and National Institutes of Health protocols were strictly followed in the maintenance of the animals at the Yerkes National Primate Center.

Single Genome Amplification

Single genome amplification and sequencing was performed as specified in the Supplemental Experimental Procedures.

Proviral Constructs

Proviral constructs were generated as described in the Supplemental Experimental Procedures.

Envelope Expression Constructs

Full-length env genes were PCR amplified from the single genome amplicons as outlined in the Supplemental Experimental Procedures.

Cell Culture

Cells were cultured as described in the Supplemental Experimental Procedures.

Virus Stocks and Transductions

Virus stocks were generated and used for transduction of various cell lines as specified in the Supplemental Experimental Procedures.
Infectivity Assays

Virus infectivity was determined using TZM-bl, P4-CCR5, and GHOST cells as outlined in the Supplemental Experimental Procedures.

Flow Cytometric Analysis

CD4, TCR-CD3, MHC-I, CD28, CXCR-4 (BD Pharmingen, Clone 12G5), and eGFP reporter expression in human PBMCs transduced with HIV-1 (NL4-3) constructs coexpressing Nef and eGFP were measured as described (Schindler et al., 2008). IL-2R, CD69 expression was measured by standard FACS staining as outlined in the Supplemental Experimental Procedures.

PBMC Activation and Apoptosis

The effect of various nef alleles on the responsiveness of virally infected PBMCs to stimulation and subsequent induction of apoptosis was examined as described (Schindler et al., 2008). See the Supplemental Experimental Procedures for detail.

NFAT Induction

Jurkat cells stably transfected with an NFAT-dependent reporter gene vector (Fortin et al., 2004) were either left uninfected or transduced with HIV-1 Nef/eGFP constructs expressing various nef alleles. Except for those cells used as controls, cultures were treated with PHA (1 μg/ml; Murex). Luciferase activity was measured and n-fold induction was determined by calculating the ratio between measured relative light units of treated samples over untreated samples as described previously (Schindler et al., 2006).

Tetherin Antagonism

The ability of SIVsmm Nef proteins to counteract SM tetherin was determined as described in the Supplemental Experimental Procedures.

Microscopic Analyses

Confocal microscopy was performed as outlined in the Supplemental Experimental Procedures.

FACS-Based FRET assay

293T cells were transfected with constructs expressing CD3-ζ and Nef fusions with YFP and CFP, respectively, and interactions were determined as described previously (Banning, et al., 2010).

Nef Model Building

Mutations of I123L and L146F were inserted into the SIVmac239 Nef structure and displayed using the program PyMOL (http://www.pymol.org). Sequence polarity was calculated with the program UCSF Chimera as described (Lüft et al., 2011).

Statistical Analysis

The PRISM package version 5.0 (Abacus Concepts) was used for all calculations.

ACCESSION NUMBERS

The GenBank accession numbers for the SIVsmm env-nef SGA sequences reported in this paper are KF477983–KF478193.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2013.10.028.

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