ART influences HIV persistence in the female reproductive tract and cervicovaginal secretions

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The recently completed HIV prevention trials network study 052 is a landmark collaboration demonstrating that HIV transmission in discordant couples can be dramatically reduced by treating the infected individual with antiretroviral therapy (ART). However, the cellular and virological events that occur in the female reproductive tract (FRT) during ART that result in such a drastic decrease in transmission were not studied and remain unknown. Here, we implemented an in vivo model of ART in BM/liver/thymus (BLT) humanized mice in order to better understand the ability of ART to prevent secondary HIV transmission. We demonstrated that the entire FRT of BLT mice is reconstituted with human CD4+ cells that are shed into cervicovaginal secretions (CVS). A high percentage of the CD4+ T cells in the FRT and CVS expressed CCR5 and therefore are potential HIV target cells. Infection with HIV increased the numbers of CD4+ and CD8+ T cells in CVS of BLT mice. Furthermore, HIV was present in CVS during infection. Finally, we evaluated the effect of ART on HIV levels in the FRT and CVS and demonstrated that ART can efficiently suppress cell-free HIV-RNA in CVS, despite residual levels of HIV-RNA+ cells in both the FRT and CVS.

Introduction
Most clinical trials of HIV prevention have aimed at preventing HIV acquisition by topical or systemic administration of preventative antiretroviral drugs to uninfected individuals (1–10). Results from these clinical trials have shown either partial or no protection. The basis for these discordant results are not yet clear and have been postulated to be due to a combination of a lack of adherence and inadequate drug levels at the site of exposure (5, 7, 11). In contrast, the HIV prevention trials network study 052 (HPTN 052) demonstrated 93% protection against secondary heterosexual transmission when infected individuals received early antiretroviral therapy (ART) (12). Importantly, no linked partner infections were observed when the HIV-infected participant was stably suppressed by ART. The prevailing hypothesis for the success of HPTN 052 is that ART reduces genital cell-free and/or genital cell–associated HIV to levels that are too low to support HIV transmission (12). This hypothesis is supported by observational studies suggesting a strong correlation between plasma/genital HIV-RNA levels and risk of heterosexual transmission (13, 14); it is also supported by the ability of ART to decrease the genital levels of HIV in both men and women (15–17). There is very limited data in the literature to determine whether transmission occurs from cell-free virus only or if productively infected cells themselves can transmit HIV in the absence of cell-free virions (18).

In order to better understand the ability of ART to prevent secondary transmission of HIV, we used a small animal model of HIV infection to further characterize key virological and immunological events that occur in the female reproductive tract (FRT) during ART. We designed the following experiments using BM/liver/thymus humanized mice (BLT mice). First, we performed a detailed and comprehensive phenotypic characterization of the human lymphocyte subsets present in the FRT and cervicovaginal secretions (CVS). Next, we analyzed HIV levels and cellular dynamics in CVS during HIV infection. Finally, we evaluated virological suppression and cellular dynamics in the FRT and CVS during ART. We provide data demonstrating that HIV replication occurs in CVS soon after exposure and continues during the course of infection. This is followed by an increase of CD4+ T cells in CVS, providing additional target cells for infection. This CD4+ T cell increase is followed by a delayed increase of CD8+ T cells in CVS. Surprisingly, despite the strong suppressive effect of ART on the viral load in CVS, HIV-RNA+ cells were still present in both the FRT and CVS. However, when analyzed ex vivo, cells isolated from the FRT and CVS of ART-suppressed BLT mice did not transmit HIV in a coculture assay. Thus, our results provide in vivo evidence supporting the hypothesis behind the success of HPTN 052 (12) for limiting sexual transmission from HIV–infected women.

Results
Reconstitution of the FRT of BLT mice with human CD4+ cells. BLT mice were prepared as previously described (19–23) and were well reconstituted with human hematopoietic cells (CD45+) in peripheral blood (PB) (median 70%, range 22–95, interquartile

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range 56–78, n = 142). In addition, we used IHC to assess reconstitution and distribution of HIV target cells (human CD4+ cells, CD68+ myeloid/immature DC, and CD11c+ DCs) in the FRT of BLT mice (Figure 1 and Supplemental Figures 1 and 2; supplemental material available online with this article; doi:10.1172/JCI64212DS1). Human CD4+ cells were observed throughout the FRT. Specifically, in the vagina, human CD4+ cells were mainly observed in the lamina propria, while few CD4+ cells were present in the epithelium. Vaginal CD4+ cells were dispersed throughout the lamina propria both as single cells and as focal aggregates in close proximity to the epithelial layer, similar to their distribution in healthy women (24, 25). Cervical CD4+ cells were present as single cells close to the epithelium and distributed throughout the lamina propria. In the uterine endometrium, CD4+ cells were found in the stroma both as small clusters closely adjacent to the epithelial layer and scattered as single cells, resembling their distribution in women (24–26). Similarly, inspection of the FRT for the presence of human macrophages and DCs demonstrated that, like in humans, these cell types are dispersed throughout the lamina propria of the vagina, cervix, and uterus (refs. 24, 26–28, and Supplemental Figures 1 and 2). Furthermore, like in women, macrophages and DCs in BLT mice were more abundant in the cervical mucosa than the vaginal mucosa (24). Together, these results show the efficient repopulation and adequate distribution of the human target cells postulated to be involved in vaginal HIV transmission throughout the entire FRT of BLT mice (29).

The majority of human T cells in the FRT and CVS of BLT mice express a memory phenotype and the CCR5 HIV coreceptor. After establishing the presence and distribution of human CD4+ cells in the FRT of BLT mice (Figure 1), we proceeded to compare their cell surface phenotype to those from PB and CVS (Figure 2). CVS samples from BLT mice were collected, and cells were isolated for analysis. Cells were also isolated from the entire FRT. CVS and FRT samples were then analyzed for the presence of human CD4+ and CD8+ T cells and compared with samples obtained from PB (Figure 2, A and B). Our results show that, in all 3 compartments, the majority of human CD4+ T cells in the FRT of BLT mice (Figure 1) and the level of CD4+ T cells in all 3 compartments was the same prior to infection with HIV (Figure 2B) (PB vs. FRT, P = 0.22; PB vs. CVS, P = 0.22; and CVS vs. FRT, P = 0.22). In addition, when compared with PB T cells, a significantly higher percentage of cells in the FRT and CVS expressed CCR5, the primary HIV coreceptor involved in mucosal HIV transmission (ref. 30 and Figure 2C) (CD4+ T cells: FRT vs. PB, P = 0.01, and CVS vs. PB, P < 0.0001; CD8+ T cells: FRT vs. PB, P = 0.01, and CVS vs. PB, P < 0.0001). This is consistent with what has been observed in women and with the preferential mucosal transmission of CCR5-tropic HIV (31, 32).

To confirm that cells in the FRT of BLT mice have a similar memory phenotype as in women (24, 33, 34), we characterized the naive-memory phenotype of T cell subsets in PB, the FRT, and CVS (naive: CD45RA+CD27+; central memory (CM): CD45RA CD27+; effector memory (EM): CD45RA CD27+) (Figure 2, D and E). The majority of CD4+ and CD8+ human T cells in PB of BLT mice had a naive phenotype (CD45RA/CD27+) (CD4+ T cells: naive vs. CM, P < 0.0001, and naive vs. EM, P < 0.0001; CD8+ T cells: naive vs. CM, P < 0.0001, and naive vs. EM, P < 0.0001). In contrast, the majority of human T cells in CVS had a CM phenotype (CD4+ and CD8+ T cells: CM vs. naive, P < 0.0001, and CM vs. EM, P < 0.0001). Also notable were significantly higher levels of CD4+ EM T cells (CD45RA CD27) in CVS and the FRT compared with PB (Figure 2D) (EM cells in FRT vs. PB, P = 0.022 and CVS vs. PB, P < 0.0001) and of CD8+ EM T cells in CVS compared with PB (P < 0.0001) (Figure 2E). Increasing evidence suggests that CD4+ T cells expressing the integrin αβ7 heterodimer (a mucosal homing receptor) are especially susceptible to HIV infection and may play a central role in the earliest steps of transmission (35, 36). Therefore, we investigated the expression of αβ7 on memory CD4+ T cells in PB of humans and PB, FRT, and CVS of BLT mice. The percentage of memory CD4+ T cells in PB expressing αβ7 was similar in humans and BLT mice (Supplemental Figure 3). Furthermore, memory CD4+ T cell expression of αβ7 in the FRT and CVS of BLT mice was significantly lower than in PB (PB vs. FRT, P = 0.003, and PB vs. CVS, P < 0.0001) (Supplemental Figure 3), which is similar to what is observed in women, where fewer FRT memory CD4+ T cells express αβ7 when compared with PB (32). We also observed no difference in CD4+ memory T cell αβ7 expression between the FRT and CVS of BLT mice (P = 0.13) (Supplemental Figure 3). In summary, the majority of lymphocytes in the FRT and CVS of BLT mice were memory T cells expressing CCR5, which is remarkably similar to what is observed in the FRT of humans (32, 33, 37).

HIV and HIV-infected cells are present in the FRT and CVS of BLT mice. We established that HIV-infected cells are present in the FRT of HIV-infected BLT mice by analyzing different regions of the FRT from a BLT mouse infected vaginally with HIV-1JR-CSF, a well-characterized T cell tropic isolate that utilizes CCR5 as a coreceptor for entry (ref. 38 and Figure 3A). HIV-RNA was also readily detected in CVS from infected mice (Figure 3, B–D). As seen in humans, there was an overall significant positive linear correlation between the viral load in plasma and CVS (Spearman’s correlation coefficient = 0.55 [95% CI, 0.36–0.69, P < 0.001, n = 51]) (Figure 3E and refs. 31, 39–42).

HIV infection results in a dramatic reduction in the percentage of CD4+ T cells in the FRT and CVS. Following HIV infection, mucosal tissues like the gut undergo a rapid depletion of CD4+ T cells (43), and this hallmark of HIV infection is faithfully recapitulated in BLT mice (20, 21). A significant decrease in the percentage of cervical CD4+ T cells during chronic HIV infection in women has also been observed (31–33, 37, 44). In order to investigate these phenomena in the FRT and CVS of BLT mice, we analyzed CD4+ T cell levels longitudinally after HIV vaginal exposure. Prior to HIV infection, the levels of CD4+ T cells were similar in the FRT, CVS, and PB (Figure 2B and Figure 4, A–D). After vaginal infection, there was a decline in the levels of human CD4+ T cells in all 3 compartments (Figure 4, A and D). Specifically, a modest decline to approximately 70% was observed in the PB, whereas CD4+ T cell levels in CVS incurred a significantly more dramatic decrease to approximately 5% (for weeks 3–7, P = 0.0001–0.0042). Interestingly, the levels of CD4+ T cells in the FRT also declined but stabilized at about 30%, an intermediate level between PB and CVS (Figure 4, A and D).

HIV infection after mucosal exposure occurs by 3 main routes: rectal, vaginal, and oral. BLT mice are an outstanding model for each of these individual modes of transmission (20–23, 45–48). In
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in humans and BLT mice results in a dramatic decrease in the levels of CD4+ T cells in the FRT and CVS that is not reflected in PB. In previous studies, no determination was made about whether the observed decrease in CD4+ T cells in the FRT was due to a reduction in the number of CD4+ T cells and/or an increase in the number of CD8+ T cells (33, 49, 50). To address this important question, we performed a longitudinal analysis of CD4+ and CD8+ T cell numbers in CVS following HIV infection. Regardless of the route of exposure (vaginal, rectal, or oral) or type of inoculum, we observed significant increases in the numbers of both CD4+ and CD8+ T cells in CVS, albeit with different kinetics (Figure 5, A–C). Specifically, the increase in CD4+ T cells occurred during weeks 1–2 after infection and was somewhat modest (5- to 13-fold for each of the 3 exposure routes). In sharp contrast, the increase in the number of CD8+ T cells occurred later at weeks 2–7 and was much greater (231-, 218-, and 56-fold after vaginal, oral, and rectal infections, respectively). These results demonstrate that the reduction in the percentages of CD4+ T cells observed in CVS of

order to understand the cellular dynamics that occur in the FRT after local versus distal HIV infection, we evaluated all 3 modes of mucosal HIV acquisition following exposure to cell-free and/or cell-associated HIV (Supplemental Table 1). Specifically, to complement the data presented above obtained after vaginal exposure, we analyzed the levels of human CD4+ T cells in PB, FRT, and CVS following rectal and oral HIV exposures (Figure 4, B–D). Longitudinal analysis demonstrated a decrease in the percentage of CD4+ T cells in the FRT and CVS following rectal and oral infection (Figure 4, B–D). Moreover, as in the case of vaginal infection, there was a significant decrease in the levels of CD4+ T cells in CVS when compared with PB (rectal infection weeks 2–9, P = 0.049–0.002; oral infection weeks 1–6, P = 0.0016–0.0001). Regardless of the route of infection (vaginal, rectal, or oral) or type of inoculum (cell-free or cell-associated virus), there was a greater decrease in the levels of CD4+ T cells in the FRT and CVS in comparison to PB.

HIV infection results in increased numbers of human CD4+ and CD8+ T cells in CVS of BLT mice. As indicated above, HIV infection

Figure 1. Human CD4+ T cells are present throughout the FRT of BLT mice. Immunohistochemical analysis of the entire FRT of a HIV–BLT mouse demonstrates the presence of CD4+ cells in the vagina, cervix, and uterus. CD4+ cells are stained brown. Scale bars: 100 μm.
decrease in the percentage of CD4+ T cells in CVS described above and the less pronounced decrease observed in PB. During ART, there was a dramatic increase in the percentage of CD4+ T cells in CVS and an increase in the percentage of CD4 + T cells in PB (Figure 6B). These findings are in agreement with the fact that HIV + women on ART have a significantly higher percentage of cervical CD4+ T cells than infected women not receiving treatment (57). In addition — as indicated above — prior to treatment, there was a dramatic increase in total CD8+ T cells in CVS (Figure 6C). However, in response to ART, there was a rapid and substantial decrease in the numbers of CD8+ T cells in this compartment. In contrast, while the numbers of CD8 + cells decreased, the numbers of CD4 + T cells increased, resulting in the eventual return to near-normal levels (Figure 6C). These results reveal that the observed increase in the percentage of CD4+ T cell in CVS during ART (Figure 6B) was caused by a concurrent decrease in the numbers of CD8+ T cells and an increase in the numbers of CD4+ T cells.

ART efficiently suppresses HIV in CVS and restores CD4+ and CD8+ T cell numbers. In order to examine the effect of ART on HIV levels and CD4+ and CD8+ T cell numbers in CVS, we treated infected mice with an ART regimen that consisted of TDF, emtricitabine (FTC), and raltegravir (RAL). This triple-drug combination has been shown to strongly suppress viral load in both humans and BLT mice with continuous dosing (51–53). However, ART discontinuation in patients results in a rapid rebound of plasma viremia as well as a loss of PB CD4+ T cells (54, 55). These phenomena are also fully recapitulated in BLT mice (53, 56). In BLT mice infected vaginally with cell-associated virus (Supplemental Table 1), ART administration resulted in a dramatic and sustained decrease in viral load in both plasma and CVS as early as 2 weeks after ART initiation (Figure 6A). Analysis of CD4+ T cells in PB and CVS from these mice prior to ART demonstrated the characteristic steep decrease in the percentage of CD4+ T cells in CVS described above and the less pronounced decrease observed in PB. During ART, there was a dramatic increase in the percentage of CD4+ T cells in CVS and an increase in the percentage of CD4+ T cells in PB (Figure 6B). These findings are in agreement with the fact that HIV+ women on ART have a significantly higher percentage of cervical CD4+ T cells than infected women not receiving treatment (57). In addition — as indicated above — prior to treatment, there was a dramatic increase in total CD8+ T cells in CVS (Figure 6C). However, in response to ART, there was a rapid and substantial decrease in the numbers of CD8+ T cells in this compartment. In contrast, while the numbers of CD8+ cells decreased, the numbers of CD4+ T cells increased, resulting in the eventual return to near-normal levels (Figure 6C). These results reveal that the observed increase in the percentage of CD4+ T cell in CVS during ART (Figure 6B) was caused by a concurrent decrease in the numbers of CD8+ T cells and an increase in the numbers of CD4+ T cells.
cells constitute an important source of infectious virus, we established a sensitive coculture assay to measure the number of infectious cells in PB, CVS, and the FRT of infected mice receiving ART (Figure 7, C and D). Our results showed (i) that infectious cells were readily detected in PB, CVS, and the FRT of infected BLT mice; (ii) that ART suppresses the number of infectious cells in each of these compartments; and (iii) that this reduction was statistically significant ($P = 0.04$) (Figure 7D). Thus, our results demonstrate that ART efficiently suppresses levels of cell-free HIV and infectious cells in PB, CVS, and the FRT. Most importantly, we then determined if ART suppresses the number of HIV-RNA$^+$ cells in the PB, CVS, and FRT of BLT mice to levels below those needed to prevent secondary mucosal HIV transmission. We assessed the ability of HIV-infected cells to establish infection in vivo by vaginally exposing BLT mice to 2 different doses of HIV-infected PB mononuclear cells (PBMCs) and monitoring their plasma level of HIV-RNA for levels observed in plasma and CVS (Figure 7A) parallels a similar decrease in the levels of cell-associated HIV in PB, CVS, and the FRT. For this purpose, we isolated cells from PB, FRT, and CVS and analyzed their levels of cell-associated HIV-RNA (Figure 7B). Our results show that ART significantly decreased the amount of cell-associated HIV-RNA in all 3 compartments (PB ART vs. No ART, $P = 0.0009$; FRT ART vs. No ART, $P = 0.0002$; and CVS ART vs. No ART, $P = 0.015$) (Figure 7B). However, cell-associated HIV-RNA remained readily detectable in the FRT and/or CVS of the majority of mice undergoing ART. Therefore, despite the strong reduction in the levels of cell-free HIV-RNA in mice receiving ART, our analysis reveals an important dichotomy between the suppression of cell-free HIV-RNA levels observed in CVS and the continued presence of residual levels of cell-associated HIV-RNA in the FRT/CVS. To determine whether these residual HIV-RNA$^+$ cells constitute an important source of infectious virus, we established a sensitive coculture assay to measure the number of infectious cells in PB, CVS, and the FRT of infected mice receiving ART (Figure 7C and D). Our results showed (i) that infectious cells were readily detected in PB, CVS, and the FRT of infected BLT mice; (ii) that ART suppresses the number of infectious cells in each of these 3 compartments; and (iii) that this reduction was statistically significant ($P = 0.04$) (Figure 7D). Thus, our results demonstrate that ART efficiently suppresses levels of cell-free HIV and infectious cells in PB, CVS, and the FRT. Most importantly, we then determined if ART suppresses the number of HIV-RNA$^+$ cells in the PB, CVS, and FRT of BLT mice to levels below those needed to prevent secondary mucosal HIV transmission. We assessed the ability of HIV-infected cells to establish infection in vivo by vaginally exposing BLT mice to 2 different doses of HIV-infected PB mononuclear cells (PBMCs) and monitoring their plasma level of HIV-RNA for...
8 weeks. As shown in Figure 7E, all BLT mice exposed to 5,000 HIV-infected PBMCs remained negative for HIV-RNA. However, 50% of mice exposed to 10,000 HIV-infected cells became positive for plasma HIV-RNA. Our results revealed that the number of HIV-RNA+ cells in all 3 compartments analyzed from ART-suppressed BLT mice is well below what is needed to establish vaginal HIV transmission in this model.

Discussion

In this manuscript, we provide data demonstrating (i) that regardless of the route of infection (vaginal, rectal, or oral) and whether the inoculum is cell-free or cell-associated HIV, local viral replication occurs in the FRT and CVS during acute HIV infection, the time when secondary transmission is most likely to occur (Figure 3); (ii) that this is followed by a transient increase in CD4+ T cell levels in CVS during acute HIV infection, the time when secondary transmission is most likely to occur (Figure 3); (ii) that this is followed by a somewhat delayed increase in CD8+ T cell levels in CVS (Figure 5); and (iii) that this is itself followed by a somewhat delayed increase in CD8+ T cell levels in CVS (Figure 5). In addition, our study provides in vivo evidence supporting the hypothesis behind the success of HPTN 052 (12): that initiating ART can reduce the risk of secondary HIV transmission by efficiently suppressing HIV levels in the genital tract (Figures 6 and 7).

Interestingly, our results also highlighted a potentially important dichotomy between the levels of cell-free virus and cell-associated HIV-RNA in the FRT and CVS of ART-suppressed mice. Specifically, in animals with undetectable cell-free HIV-RNA in plasma and CVS, significant levels of HIV-infected cells producing viral RNA remain in the FRT and CVS of some ART-treated mice (Figure 7). The presence of cell-associated HIV-RNA has been demonstrated in other secretions of ART-suppressed women. Valea et al. demonstrated the presence of cell-associated HIV-RNA in breast milk obtained from ART-treated mothers with undetectable levels of cell-free HIV-RNA in plasma and breast milk (58). These results are in agreement with our observations in BLT mice demonstrating that cell-associated HIV-RNA can persist in mucosal secretions despite ART. Collectively, our results and those of Valea et al. may have important implications for the design of effective HIV prevention and curative approaches.

Worldwide, the majority of new HIV infections occur after heterosexual exposure (59). In vaginally exposed women, the primary ports of HIV entry are the mucosal surfaces of the vagina, cervix, and uterus (60–62). The identity and the location of the initial cells involved in HIV-1 transmission are a subject of great debate (29). The DC-T cell milieu is a highly permissive site for virus growth, and DCs likely contribute to driving the productive infection in CD4+ T cells (63–67). Hence, both intraepithelial Langerhans cells and DCs have potential important roles in vaginal HIV transmission (65–69). However, the mucosa of the human FRT contains an abundance of CD4+ T cells (24, 25, 34, 68), and experiments in both NHPs (non-human primates) and human explant models suggest that the first productively infected cells are likely T cells (61, 68, 70–73).

Figure 4. After vaginal, rectal, or oral HIV infection, there is a decrease in the percentage of CD4+ T cells in CVS and the FRT. (A–C) The percentage of CD4+ T cells in PB (filled diamond, solid line) and CVS (open triangle, dashed line) was measured in HIV− (n = 28) and vaginally (A) (n = 18), rectally (B) (n = 11), or orally (C) (n = 12) infected BLT mice. Data is shown as mean ± SEM. A Mann-Whitney U test was used to compare CD4+ T cell levels between the PB and CVS (*P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001). (D) Additional analysis of the percentage of CD4+ T cells in the FRT was performed in HIV− BLT mice (n = 7); vaginally infected BLT mice (filled circle) at weeks 1 (n = 1), 3 (n = 1), 5 (n = 1), and 7 (n = 1); rectally infected mice (asterisk) at weeks 6 (n = 2) and 10 (n = 1); and an orally infected mouse (filled square) at week 8 (n = 1) after exposure.
Regardless, each of the most relevant human HIV target cells (CD4+ T cells, macrophages, and DCs) are present throughout the entire FRT of BLT mice (Figure 1, Supplemental Figures 1 and 2, and ref. 21). Reconstitution of the FRT of BLT mice with the appropriate human hematopoietic cells renders BLT mice susceptible to vaginal HIV transmission (21). The susceptibility of BLT mice to vaginal HIV infection has allowed this model to be used to evaluate novel approaches of HIV prevention (21, 23, 74).

In this manuscript, we utilized BLT mice to elucidate and study critical events occurring in the FRT and CVS during HIV infection. Specifically, by performing comprehensive analyses of the T cell subsets present in the FRT and CVS, we have gained insight into the human immune cell populations in this organ. Consistent with observations made in healthy women, the majority of the human lymphocytes present in both the FRT and CVS of BLT mice are memory T cells (Figure 2D and refs. 31, 32). Also, consistent with the preferential viral transmission of CCR5-tropic viruses, a high percentage of CD4+ T cells present in the FRT and CVS express CCR5 (Figure 2C and refs. 31, 32). Furthermore, consistent with humans, a significant number of the memory CD4+ T cells present in the FRT and CVS of BLT mice expressed α4β7 (Supplemental Figure 3). The similarities between the phenotypes of hematopoietic cells present in the FRT of humans and BLT mice emphasize the utility of BLT mice as an in vivo model for the study of events occurring at the site where HIV exposure occurs.

Our results demonstrating parallel reductions in the percentage of CD4+ T cells in the FRT and CVS of BLT mice indicate that the cell populations are closely linked throughout the course of infection (Figure 4). These striking similarities between the dynamics of T cells present in CVS and the FRT after HIV infection suggest that cells from CVS could be potentially used as a surrogate for monitoring some of the changes that occur in the FRT. Thus, these results may have significant implications that could facilitate and simplify future studies of transmission and prevention in both humans and in NHP models by minimizing the need to harvest or biopsy the FRT.

In order to study the dynamics of human CD4+ and CD8+ T cells and viral replication that occur in the FRT after vaginal HIV infection, we vaginally exposed BLT mice to HIV. Our results demonstrate an increase of CD4+ T cells in CVS during the first 2 weeks after infection (Figure 5A), providing additional HIV target cells to sustain and potentially spread the initial infection. These results are consistent with HIV-RNA being present in CVS within 1 week after exposure (Figure 3B), suggesting that local HIV replication occurs in the FRT and/or CVS followed by the establishment of systemic infection in all mice by 2 weeks after exposure (Figure 3B). Especially noteworthy is the timing of viral shedding into CVS after vaginal exposure, which is characterized by an early peak in viremia followed by a gradual decline (Figure 3B) that mimics the HIV genital shedding profile observed in CVS of women during acute HIV infection (39). Notably, we also observed an increase in CD8+ T cells in CVS after vaginal infection (Figure 5A). However, in contrast to the increase of CD4+ cells occurring within one to 2 weeks after exposure, the increase of CD8+ T cells was first detected 2 weeks after exposure and did not peak until week 5-7 (Figure 5A). These results are consistent with reports from vaginal-infection studies in NHPs demonstrating that initial SIV infection takes place in a few CD4+ T cells, resulting in local inflammation and recruitment of additional CD4+ T cells (61, 72, 75). Clusters of SIV-infected cells are present within inflammatory infiltrates, which increase in size during days 4-10 after exposure (61, 72, 75). In addition, genital CD8+ T cell influx was detected 2-3 weeks after vaginal SIV infection of NHPs, a time by which all animals had become systemically infected (72, 76). Thus, the timing of CD4+ and CD8+ T cell increase in CVS, as well as the timing of local viral replication and systemic infection that we have observed in BLT mice, is strikingly similar to NHPs. Together, these results support the hypothesis that the increase of CD8+ T cells in the CVS/FRT is delayed after exposure, potentially preventing effective suppression of HIV replication at early stages after exposure (75, 76).

We next investigated the effect of ART on HIV levels in CVS of BLT mice. Consistent with results obtained in humans, ART treatment of infected BLT mice resulted in a significant decrease in the levels of HIV in both PB and CVS (15, 16). However, our finding showing the absence of cell-free HIV in CVS during ART, concurrent with the continued presence of infected cells producing HIV-RNA, could have important implications for HIV prevention and eradication strategies. Consistent with the lack of transmission observed in heterosexual couples where the infected partner is undergoing

Figure 5. After vaginal, rectal, or oral HIV infection, there is an increase of CD8+ and CD4+ T cells in CVS. (A–C) The CD8+ (filled upward triangle, dashed red line) and CD4+ T cell numbers (filled downward triangle, blue dashed line) in CVS were measured in HIV– (n = 28) and vaginally (A) (n = 18), rectally (B) (n = 4), or orally (C) (n = 4) infected BLT mice. Data presented as mean ± SEM. Statistical analysis represents comparisons of the weeks indicated vs. the numbers of CD4+ or CD8+ T cells in naive mice. A Mann-Whitney U test was used to compare the numbers of CD4+ and CD8+ T cells in the CVS of naive and HIV-infected BLT mice (**P < 0.01, ***P < 0.001, ****P < 0.0001).
ART (12), our analysis showed that the residual levels of HIV-RNA cells present in mice receiving ART were too low to transmit HIV in vitro. This lack of HIV transmission could have been due to too few infected cells for cell-to-cell transmission and/or too little cell-free replication–competent virus produced from the residual HIV-RNA cells for in vitro infection of target cells. The residual levels of HIV-RNA cells detected in the CVS and FRT of ART-suppressed mice were well below the number of HIV-infected cells required for HIV transmission in BLT mice. The availability of a small animal model that so accurately recapitulates key aspects of the human condition represents a unique tool for the in vivo study of the intricate cellular dynamics occurring during HIV infection in the FRT and to address critical questions in the field, such as whether cell-to-cell transmission contributes sexual transmission of HIV or whether all transmission is via cell-free, replication-competent virions. In addition, this model could prove helpful in the evaluation of novel approaches to prevent cell-free and cell-associated HIV transmission.

Methods

Generation of BLT mice
BLT mice were prepared essentially as previously described (19–23). Briefly, BLT mice were individually bioengineered by transplanting (BM transplant) human fetal liver–derived CD34+ hematopoietic stem cells into NOD/SCID/γc−/− mice (stock number 00557, NSG mice; The Jackson Laboratory) implanted with autologous human liver and thymus tissue (Advanced Bioscience Resources). A sandwich of 1- to 2-mm pieces of human thymus-liver-thymus tissue was implanted under the kidney capsule of 6- to 14-week-old female NSG mice. Mice were exposed to sublethal γ irradiation (300 cGy) 4–24 hours prior to the transplantation of CD34+ stem cells (2.5–8 × 10^6 CD34+ cells) to facilitate BM engraftment.

Immunohistochemical analyses
Immunohistochemical analyses were performed on paraffin-embedded FRT sections. FRTs for IHC were harvested from BLT mice, fixed in 4% paraformaldehyde overnight at 4°C, embedded in paraffin, cut into 5-μm sections, and mounted onto poly-L-lysine-coated glass slides. Following paraffin removal, antigen retrieval (Diva Decloaker, Biocare Medical), and blocking of nonspecific Ig-binding sites (Background Sniper, Biocare Medical), tissue sections were stained with primary antibodies overnight at 4°C and developed with a biotin-free HRP-polymer system (MACH3 Mouse HRP-Polymer Detection, Biocare Medical). All tissue sections were treated with HRP substrate (ImmPACT DAB, Vector Laboratories) and then counterstained with hematoxylin. Primary antibodies specific for human CD4 (clone 1F6, Leica Biosystems), human CD68 (clone DAK-P1, Dako), and human CD11c (clone D511, Leica Biosystems) were used to identify human cells (24, 77–81). HIV-infected cells were detected with an antibody directed against HIV p24 gag (clone Kal-1, Dako). As a control, tissue sections were stained with the following isotype control antibodies: mouse IgG1 (clone DAK-G01, Dako) and mouse IgG2a (clone DAL-G02, Dako). The stained sections were scanned by ScanScope CS (Aperio, Leica Biosystems) and the images were then analyzed by ImageScope (Aperio, Leica Biosystems).

Flow cytometric analyses
Flow cytometric analyses were performed on PB, FRT, and CVS cells. CVS samples were obtained by performing a cervicovaginal lavage with sterile PBS (3 washes of 20 μl each, ~60 μl total volume). To ensure that the procedure wasatraumatic, cervicovaginal lavages were performed with 20 μl or 200 μl sterile filter pipet tips that were inserted no more than 3–5 mm into the vaginal cavity. Following centrifugation in a microcentrifuge (300 g for 5 minutes), CVS cell-free supernatants were used for HIV-RNA analyses and the cell pellets were used for flow cytometric analyses. The CVS samples contained a highly variable number of total cells (due to variable cell shedding). Therefore, the human CD3+ T cell numbers in CVS were normalized to 100,000 total cells. FRTs were harvested and digested, and single cell suspensions were isolated as previously described (23). BLT mice were monitored for levels of human hematopoietic cells (CD45+) in PB by flow cytometry, as we have previously described (19–23).
The antibody clones included in the flow cytometry panels that were used to analyze PB and cells isolated from the FRT and CVS of BLT mice are as follows: Panel A – CD3 FITC (HIT3a), CD4 PE (RPA-T4), CD8 PerCP (SK1), and CD45 APC (H130); Panel B – CD8 FITC (SK1), CXCR4 PE (12G5) or IgG2a PE (G155-178), CD4 PerCP (SK3), CD3 PE-Cy7 (SK7), CCR5 APC (3A9) or IgG2a PE (G155-178), and CD45 APC-Cy7 (2D1); Panel C – CD45RA FITC (H100) or IgG2 FITC (27-35), CD27 PE (M-T271) or IgG1k PE (MOPC-21), CD8 PerCP (SK1), CD3 PE-Cy7 (SK7), CD45 APC (H130), and CD4 APC-H7 (RPA-T4); Panel D – CD27 FITC (M-T271) or IgG1k FITC (MOPC-21), α4β7 PE (ACT-1) or IgG1k PE (P3.6.2.8.1), CD8 PerCP (SK1), CD3 PECy7 (SK7), CD45 APC (H130), CD4 APC-H7 (RPA-T4), and CD45RA Pacific Blue (F8-11-13) or IgG1k Pacific Blue (MOPC-21). All antibodies were purchased from BD Biosciences, except for CD45RA Pacific Blue (AbD Serotec), unconjugated IgG1k (eBioscience), and unconjugated α4β7 (ACT-1, catalog 11718) (NIH AIDS Research and Reference Reagent program) (82). The unconjugated α4β7 (ACT-1) and isotype IgG1k antibodies were labeled with PE using the LYNX Rapid RPE antibody conjugation kit (AbD Serotec). Gates defining CD27, CD45RA, CCR5, and α4β7 expression were set with isotype-matched fluorophore-conjugated antibodies. In Figure 2B, to account for the differential presence of NKT cells
that are CD3+ but lack expression of CD4 and CD8, the percentage of CD4+ and CD8+ T cells in PB, CVS, and FRT was calculated by dividing the number of CD3+ T cells that expressed CD4 or CD8 by the total number of CD3+ T cells that expressed either CD4 or CD8. Flow cytometric data collection and analyses were performed using a BD FACSCanto or Fortessa cytometer and FACSDiva software.

**Exposure of BLT mice to HIV and treatment with ART**

BLT mice were infected vaginally, rectally, or orally with cell-free or cell-associated CCR5 tropic HIV-1 essentially as previously described (19–23, 45, 47).

**Cell-free HIV exposures.** Stocks of HIV-1JR-CSF or HIV-1NL4-3 were prepared and titered using TZM-bl cells (AIDS Research and Reference Reagent Program, Division of AIDS, National Institute of Allergy and Infectious Diseases) as previously described (19–23). BLT mice were exposed vaginally to 3 × 10^5–3.5 × 10^6 tissue culture infectious units (TCIU), rectally to 3.5 × 10^5–4.8 × 10^6 TCIU, or orally to 1.4 × 10^6 TCIU of HIV-1.

**Cell-associated HIV exposures.** Human PBMCs were infected in vitro with HIV-1JR-CSF. PBMCs were cultured in the presence of IL-2 and PHA for 3 days and then inoculated at a MOI of 0.1 with HIV-1JR-CSF by a 2-hour spin infection. Next, PBMCs were washed 3 times, resuspended in IMDM medium, and — 3–4 days after inoculation — the percentage of HIV-infected cells was determined by intracellular staining for p24 gag (clone KC57-FITC, Beckman-Coulter) using the Fix and Perm kit (Invitrogen). BLT mice were vaginally exposed to 1 × 10^5 HIV p24+ PBMCs (up to 1.9 × 10^7 total PBMCs) or orally to 3.75 × 10^5 HIV p24+ PBMCs (up to 1.6 × 10^7 total PBMCs). Vaginal exposure to cell-associated HIV was performed by administering 5,000 or 10,000 HIV-1 p24+ cells in a volume of 10 μl to anesthetized BLT mice. We did not observe any phenotypic differences when using HIV-1JR-CSF or HIV-1NL4-3 for exposures or when using cell-free virus versus cell-associated virus. HIV-infected BLT mice designated for treatment received daily ART (by i.p. injection) consisting of FTC (140-200 mg/kg body weight), tenofovir disoproxil fumarate (TDF; 146–208 mg/kg), and RAL (56–80 mg/kg), consisting of FTC (140–200 mg/kg body weight), tenofovir disoproxil fumarate (TDF; 146–208 mg/kg), and RAL (56–80 mg/kg), as we have previously described (53). ART was administered for an average of 5.6 weeks (range 4–8 weeks). No overt drug-associated toxicity was observed.

**Real-time PCR for HIV-RNA**

Infection of BLT mice with HIV was monitored in plasma and CVS supernatant by determining levels of cell-free HIV-RNA using one-step quantitative PCR (qPCR) (ABI custom TaqMan Assays-by-Design) according to the manufacturer’s instructions (primers 5′-CAATTGCTGTTCGACATTACGAAAGA-3′ and 5′-TCGTTGATGTCCCCCCACT-3′) and MGB-probe 5′FAM-CCACCCACACGATTGAAAA GCACCTGCTAA-Q 5′ (83)) as previously described, with a limit of detection of 750 HIV-RNA copies per ml (53). The lower sensitivity of our standard assay compared with that of the standard clinical assay (50 copies per ml) is due to the lower volume of plasma that can be obtained routinely from mice. The levels of cell-associated HIV-RNA were analyzed in mononuclear cells isolated from PB, FRT, and CVS. HIV-RNA was extracted from mononuclear cells (RNeasy Mini Kit, Qiagen), and a one-step qPCR was performed as described above. For all analyses, cell-free and cell-associated HIV-RNA measures were log_{10} transformed.

**Replication competence of HIV in cells from the PB, CVS, or FRT of infected BLT mice**

The replication competence of virus present in cells obtained from infected BLT mice was determined using a coculture assay. For this purpose, cells isolated from PB, CVS, or the FRT of infected animals were cocultured with TZM-bl indicator cells in the presence of DEAE-dextran (Sigma-Aldrich) (2 μg/ml) for 24 hours. The culture medium was then removed and replaced with fresh medium. Cells were incubated for an additional 48 hours, fixed, and stained for β-galactosidase activity. Individually infected cells were manually counted under an inverted microscope. The limit of detection was two infectious units per 1 × 10^6 cells.

**Statistics**

Bivariate statistical comparisons were performed using the 2-sample Mann-Whitney U test. The Bonferroni step-down (Holm) correction was used to adjust for multiple testing for sets of related hypothesis tests in Figure 2 and Supplemental Figure 3. Data in Figure 2 and Supplemental Figure 3 were analyzed as independent observations. Some of the PB and CVS data in Figure 2 represent multiple longitudinal time points from the same animal (Figure 2B, D, and E) and/or paired observations (Figure 2, B–E). In addition, some of the PB data in Supplemental Figure 3 represent multiple longitudinal time points from the same animal and/or are paired with CVS and/or FRT data. Collectively, PB and CVS data were obtained from 30 mice for Figure 2B (n = 30, PB; n = 30, CVS), 20 mice for Figure 2C (n = 9, PB; n = 19, CVS), 29 mice for Figure 2D (n = 9, PB; n = 26, CVS), and 18 mice for Figure 2E (n = 9, PB; n = 15, CVS). The data in Supplemental Figure 3 represents a total of 21 BLT mice (n = 16, PB; n = 13, CVS; and n = 5, FRT). Associations between viral load in plasma and CVS supernatant were estimated with Spearman’s correlation coefficient. Correlation estimates were calculated at each time point, and an average correlation was computed by taking a weight average of time-specific estimates (84). The bootstrap (85) was used to calculate CIs for the average correlation and to assess whether the average correlation was nonzero. Analyses were conducted using R version 2.13, GraphPad Prism version 5.0, and SAS version 9.4.

**Study approval**

Mice were maintained at the Division of Laboratory Animal Medicine at the UNC Chapel Hill in accordance with protocols approved by the Institutional Animal Care and Use Committee.

**Author contributions**

JVG conceived, designed, and coordinated the study. RO, AW, and JVG designed the research experimental studies. RO, AW, and JVG designed the research experimental studies with additional support from PWD, MT, and LØ. RO, AW, MGH, AR, and JVG analyzed the data and wrote the manuscript.

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