Donor and Recipient Envs from Heterosexual Human Immunodeficiency Virus Subtype C Transmission Pairs Require High Receptor Levels for Entry

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Previously, our laboratory demonstrated that a severe genetic bottleneck selects for viruses with compact, glycan-restricted envelope (Env) glycoproteins during heterosexual transmission of subtype C HIV-1 (4). A similar bottleneck has since been demonstrated in other subtypes (3, 7, 12, 16). In our studies, chronically infected donor and acutely infected recipient Envs differed primarily in the first, second, and fourth variable loops (V1/V2 and V4), regions previously demonstrated to affect the structure and function of the envelope as well as binding and entry of target cells (1, 10, 11, 17). These biological properties could be important for transmission across a mucosal surface, which has been shown to play a critical but uncharacterized role in the restriction of entry (7).

One level of this restriction could be entry into potential target cells expressing low levels of CD4 receptors compared to that of T lymphocytes (6, 13, 15). One of several possible target populations is thought to be tissue macrophages. However, a recent study has demonstrated equivalent macrophage infection among these donor and recipient Envs (9). Studies presented here were designed to further investigate whether differences in transmissibility between the donor and recipient Envs can be related to their abilities to interact with the CD4 receptor and CCR5 coreceptor during the viral entry process.

For experiments described in this study, pseudotyped virus was generated by cotransfection of pcDNA3.1 or pCR3.1 expression plasmids containing individual env sequences from six donor-recipient pairs along with an HXB2 env luc proviral plasmid into 293T cells. These six pairs included four pairs in which transmission occurred from female to male (pairs 153, 221, 135, and 53) and two pairs in which transmission occurred from male to female (pairs 205 and 109). A more detailed characterization of the transmission pairs has been described previously by Derdeyn et al. (4) (pairs 53, 109, 135, and 153) and Haaland et al. (7) (pairs 205 and 215). For a majority of the transmission pairs, we analyzed five independent donor and recipient env clones. Donor env clones were selected to reflect a broad representation of the donor phylogenetic tree and V1-V4 region length (see Fig. 1). Recipient clones were generally homogeneous, differing in only a limited number of amino acids (see below). In preparation for transfection, a total of 1 × 10⁸ 293T cells were seeded in 1 ml Dulbecco’s modified Eagle’s medium containing 10% fetal bovine serum and 1% penicillin and streptomycin (DMEM-FBS-Pen-Strep) in 24-well tissue culture-treated plates. The following day, transfection was performed using 1,200 ng HXB2 backbone, 600 ng donor or recipient env plasmid, and 5.4 μg Fugene 6 according to the manufacturer’s instructions. Transfected cells were then incubated in tissue culture at 37°C for 2.5 days, after which supernatant was harvested and filtered through 0.45-μm filters. The day prior to virus harvesting, 8 × 10⁴ target cells for each of the eight HeLa cell lines (described below) were seeded into 96-well tissue culture-treated plates. Infection was performed by aspiration of media from the target cells, followed by the addition of 25 μl 80 μg/ml DEAE in DMEM with 1% FBS along with 25 μl of filtered 293T cell supernatant. Target cells and virus containing supernatant were then incubated for 2 h at 37°C, after which supernatant was harvested and filtered through 0.45-μm filters. Luciferase substrate (100 μl; Promega, Madison, WI) was added to 40 μl of each cell lysate, and light emission was quantified using a Synergy multidetector microplate reader (Biotek, Winooski, VT). Target HeLa cells pre-
FIG. 1. High-CD4-expressing cell lines. Panels present results from cells expressing high levels of CD4 with decreasing levels of CCR5 (JC-6 > JC-10 > HI-J). (A to C) Infectivity for donor (closed circles) and recipient (open triangles) Envs for each transmission pair (indicated on the x axis) were plotted as a percentage of infectivity in JC-53 cells, which express high levels of CD4 and CCR5. Each symbol represents the mean of the results for three independent experiments performed in duplicate. The horizontal line for each set indicates the median with the interquartile range. (D to F) All donor and all recipient Envs were combined and plotted as in panels A to C. A horizontal line represents the median for each group. Controls for all panels included results from two NL4-3 Env clones, two BA-L clones, and one CCR5-dependent control Env clone, R14.11. The range of V1-V4 loop lengths for each donor:recipient pair was as follows: ZM53, 269 to 292:268; ZM109, 265 to 286:266; ZM135, 285 to 290:278; ZM153, 269 to 283:278; ZM205, 265 to 284:277; ZM215, 285 to 308:288.
viously described express either low ($1.0 \times 10^4$ molecules/cell; R lines) or high ($4 \times 10^5$ molecules/cell; J lines) levels of CD4 with four different levels of CCR5 molecules expressed per cell (HI-R, 0; RC-30, 2.4 $\times 10^3$; RC-55, 2.1 $\times 10^3$; RC-49, 8.5 $\times 10^5$; HI-J, 0; JC-10, 2 $\times 10^3$; JC-6, 2.7 $\times 10^3$; and JC-53, 1.3 $\times 10^5$) (15). For each Env-pseudotyped virus, the percentage of JC-53 infectivity was calculated by dividing the luminescence of that cell line by the luminescence in JC-53 cells.

We initially evaluated the donor and recipient Envs for infectivity in the J-cell lines (Fig. 1), which express high CD4 levels approximating those of T lymphocytes. Expressed as a percentage of JC-53 activity, the virus with CXCR4-tropic NL4.3 Env infected JC-6, JC-10, and HI-J cells approximately equally (100.4%, 107.7%, and 93.32% of JC-53 activity, respectively). The previously described subtype B macrophage-tropic BA-L Env demonstrated a pronounced dependence on CCR5 levels, decreasing from 70.47% of JC-53 activity in JC-6 cells to 1.921% in JC-10 and 0.002% in HI-J cells. We also tested another subtype B R5-tropic Env, R14.11 (8), which displayed a much-reduced dependence on CCR5, maintaining almost 90% of JC-53 activity in JC-10 cells, which express 50-fold fewer CCR5 molecules. This activity decreased to 0.0001% in HI-J cells, which express no CCR5 molecules. In contrast, infection of these cell lines by the donor and recipient Envs as a whole mimicked the BA-L Env and declined from a median range of 18 to 137% in JC-6 cells to a range of 0.8 to 13.1% in JC-10 cells and 0.0001 to 0.007% in HI-J cells. Nonparametric Mann-Whitney $t$ test analyses of medians revealed that for all six pairs in JC-6, JC-10, and HI-J cells, there was no significant difference in infection between the donor and recipient Envs. Importantly, when medians from all donor Envs and all recipient Envs were grouped and compared by both Mann-Whitney and two-way analysis of variance (ANOVA), no significant difference in infection was observed between the donor and recipient groups in any of the J-cell lines (Fig. 1D to F).

We next evaluated infection of the R-cell lines that express various levels of CCR5 and a 10-fold lower level of CD4, which approximate levels on macrophages. Infection of these cells, relative to that of JC-53 cells, was drastically diminished for Envs from all six pairs and decreased further with decreasing levels of CCR5 (Fig. 2A to D). Medians ranged from 0.015 to 3.1% in RC-49 cells, 0.002 to 1.75% in RC-55 cells, 0.01 to 1.4% in RC-30 cells, and 0.0001 to 0.0038% in HI-R cells. For 5/6 transmission pairs, there was no statistically significant difference in infection between donor and recipient Envs in any of the R-cell lines. For one pair, transmission pair 53, recipient Envs more efficiently infected RC-49 cells ($P = 0.0159$), as well as RC-55 cells ($P = 0.0317$), RC-30 cells ($P = 0.0317$), and HI-R cells ($P = 0.0159$). Both BA-L and R14.11 Envs were able to infect R-cell lines better than a majority of transmission pair Envs, demonstrating 2.5% and 1.3% JC-53 activity, respectively, in RC-49 cells, which express the highest level of CCR5. They also both displayed similarly reduced infection efficiencies of 0.6% and <0.005% in RC-30 and HI-R cells, respectively. In contrast, NL4.3 Env exhibited infection efficiencies that varied inversely to CCR5 expression levels (5.4% in RC-49 cells, 7.2% in RC-55 cells, 19.5% in RC-30 cells, and 26.4% in HI-R cells), ruling out a general reduction in infectivity as CCR5 levels decreased. The 4-fold difference in infection efficiency between RC-49 and HI-R cells is consistent with a 6-fold difference in surface expression of CXCR4 between these two cell lines (our unpublished data). When donor Envs and recipient Envs from all transmission pairs were combined and analyzed by both Mann-Whitney and two-way ANOVA, there was again no significant difference in infection between medians of the two groups in any of the R-cell lines (Fig. 2E to H).

Results from this study demonstrate that in this subtype C-infected cohort of heterosexual couples, in which recent transmission from one partner to the other has occurred, donor and recipient Envs display a strict dependence on CCR5 levels as well as a requirement for high levels of CD4 receptor. Thus, although the majority of subtype C Envs has been shown to use CCR5 and rarely switch to CXCR4 tropism (2, 14, 18), this is the first study to evaluate the ability of primary subtype C Envs from chronically infected individuals and their acutely infected partners to utilize different levels of receptor and coreceptor. The requirement for high CCR5 levels by subtype C Envs is consistent with reports that infectivity by macrophage-tropic subtype B primary isolates are highly dependent on this coreceptor (20). Previous studies of the cell lines used in this report have also demonstrated a strict dependence on CCR5 expression by subtype B isolates from acutely infected individuals (15). However, in a study of several subtype B primary isolates, including BA-L, high levels of either CCR5 or CD4 could compensate for low levels of the other, and maximal infectivity was observed in JC-10 cells that express only trace levels of CCR5 (15). This was not observed for the subtype C Envs or BA-L in the current analysis. The differences in infection of JC-10 cells observed for BA-L in the current study and the prior report likely reflect the presence of multiple genetic variants in the primary HIV BA-L isolate used in the latter, versus the Env clone used in this study. Indeed, when previously described cloned Envs (BBB, NNB, NBB) were tested in a replication-competent green fluorescent protein (GFP) reporter system for entry into the cell lines used in this study, the infectivities were consistent with the prior report (data not shown) (19). Using V1-V5 chimeric replication-competent subtype A viruses, Etemad et al. (5) observed reduced replication in JC-10 cells for those chimeras containing the acute virus fragment relative to that for those containing fragments from chronically infected partners, although both replicated on these cells less efficiently than on JC-53 cells. Thus, it is possible that viruses from subtype A transmission pairs differ in their CCR5 requirements from those we have observed for subtype C Envs.

Results from the high CD4 J-cell lines revealed a fairly uniform infectivity among the Envs when both CD4 and CCR5 are abundant. As receptor and coreceptor levels are decreased, the infectivity profile becomes significantly more heterogeneous, likely reflecting subtle differences in the ability of individual Envs to interact with the target cell. This is consistent with a recent report showing a similarly heterogeneous infection profile in macrophages with these same Envs (9). While the infectivity profile of recipient Env clones would be predicted to be relatively homogeneous due to the small number of amino acid differences between these clones, there was an occasional outlier detected among the recipients within a given pair. Sequencing analyses of two sets of recipient sequences with clear outliers in entry phenotypes revealed nonsynony-
FIG. 2. Low-CD4-expressing cells. Panels present results from cells expressing decreasing levels of CCR5 (RC-49 > RC-55 > RC-30 > HI-R). Experiments are plotted as described in the legend for Fig. 1. A star indicates statistical significant difference ($P < 0.05$) between the medians of the donor and recipient Env's.
mous nucleotide changes unique to the outlier clone in each case that could be responsible for reduced infectivity; in pair 205, amino acid changes were present in the C1 and V2 loop region, and in pair 53, changes were observed in the bridging sheet of gp120 and the MFPR region of gp41 (data not shown).

In our initial studies of subtype C transmission pairs, where we observed that Envs with more-compact variable loops were more frequently found in recipient partners, we suggested that this could reflect differential usage of receptors and/or coreceptors on key target cells early in infection. The current findings using epithelial cells expressing different levels of these molecules, presented here, are not consistent with this hypothesis and argue that the virus emerging from the genetic bottleneck is not selected for based on a unique pattern of CD4/CCR5 receptor utilization on target cells of the genital mucosa. Instead it appears that these viruses, like those replicating in the chronically infected partner, are effectively adapted to growth in CCR5-expressing, mucosal CD4+ T cells.

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