Resistance to type 1 interferons is a major determinant of HIV-1 transmission fitness

Shilpa S. Iyerab,1, Frederic Bibollet-Rucheac,1, Scott Sherrill-Mixa,d,f, Gerald H. Learnb, Lindsey Plenderleithb, Andrew G. Smithb, Hannah J. Barbiana,b,1, Ronnie M. Russello, Marcos V. P. Gondimb, Catherine Y. Baharic, Christiana M. Shavoc, Yingying Li, Timothy Deckerc, Barton F. Haynesd,e,f, George M. Shawa,b, Paul M. Sharpb, Persephone Borrowf, and Beatrice H. Hahnb,2

*Department of Medicine, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA 19104; 1Department of Microbiology, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA 19104; 2Institute of Evolutionary Biology, and Centre for Immunity, Infection and Evolution, University of Edinburgh, Edinburgh EH9 3LF, United Kingdom; 3Human Vaccine Institute, Duke University Medical Center, Durham, NC 27710; 4Department of Medicine, Duke University Medical Center, Durham, NC 27710; 5Department of Immunology, Duke University Medical Center, Durham, NC 27710; and 6Nuffield Department of Clinical Medicine, University of Oxford, Oxford OX3 7FZ, United Kingdom

Contributed by Beatrice H. Hahn, December 8, 2016 (sent for review November 14, 2016; reviewed by Michael H. Malim and Greg J. Towers)

Sexual transmission of HIV-1 is an inefficient process, with only one or few variants of the donor quasispecies establishing the new infection. A critical, and as yet unresolved, question is the mucosal bottleneck select for viruses with enhanced transmission fitness. Here, we characterized 300 limiting dilution-derived virus isolates from the plasma, and in some instances genital secretions, of eight HIV-1 donor and recipient pairs. Although there were no differences in the amount of virion-associated envelope glycoprotein, recipient isolates were on average threefold more infectious (P = 0.0001), replicated to 1.4-fold higher titers (P = 0.0004), and were significantly more resistant to type I IFNs than the corresponding donor isolates. Remarkably, transmitted viruses exhibited 7.8-fold higher IFNα2 (P < 0.00001) and 39-fold higher IFNβ (P < 0.00001) half-maximal inhibitory concentrations (IC50) than did donor isolates, and their odds of replicating in CD4+ T cells at the highest IFNα2 and IFNβ doses were 35-fold (P < 0.00001) and 250-fold (P < 0.00001) greater, respectively. Interestingly, pretreatment of CD4+ T cells with IFNβ, but not IFNα2, selected donor plasma isolates that exhibited a transmitted virus-like phenotype, and such viruses were also detected in the donor genital tract. These data indicate that transmitted viruses are phenotypically distinct, and that increased IFN resistance represents their most distinguishing property. Thus, the mucosal bottleneck selects for viruses that are able to replicate and spread efficiently in the face of a potent innate immune response.

For obvious reasons, viruses cannot be collected from, or studied in, humans at the time of transmission. However, by sequencing plasma virion RNA (vRNA) in the first few weeks following transmission, it is possible to enumerate and infer the genome(s) of the virus(es) that established the infection (9, 12–14). In the absence of adaptive immune pressures, HIV-1 diversifies in a random fashion, with viral sequences exhibiting a Poisson distribution of mutations and a star-like phylogeny that coalesces to an inferred consensus sequence. This consensus sequence represents the genome of the virus that initiated the infection, termed the transmitted founder (TF) virus (9). Single genome amplification (SGA) of plasma vRNA, which precludes PCR artifacts such as Taq polymerase-mediated recombination (15–18), revealed that in the great majority (~80%) of sexual transmission cases, a single TF virus establishes the new infection (9, 12, 13, 19–21). The ability to infer and molecularly clone the genomes of TF viruses has permitted their biological characterization. Initial studies showed that TF viruses use the cell-surface molecules CD4 and CCR5 as their receptor and coreceptor, and replicate efficiently

Understanding the host and viral factors that influence HIV-1 transmission may aid the development of an effective AIDS vaccine. In 2015, ~2 million individuals were newly infected with HIV-1, the great majority of whom acquired the virus by sexual routes (1). Although a number of factors, such as high donor viral loads, genital inflammation, altered mucosal microbiota, and recipient gender, are known to increase the infection risk (2–4), virus transmission across intact mucosal surfaces is inherently inefficient, with only a small fraction (less than 1%) of unprotected sexual exposures leading to productive infection (5–8). This inefficiency is exemplified by a stringent population bottleneck, in which only one or a limited number of variants from the diverse quasispecies of the transmitting donor establish the new infection (9). Transmitted viruses are not usually the most abundant strains in the genital secretions of infected donors (10), and analyses of viral sequences from 137 matched donor and recipient pairs indicated that viruses with a more ancestral genotype are preferentially transmitted (11). These data suggested that mucosal transmission selects for variants with enhanced transmission fitness (11). However, the viral properties that contribute to this transmission fitness have not been defined.

Sexual transmission of HIV-1 is an inefficient process, with only one or few variants of the donor quasispecies establishing the new infection. A critical, and as yet unresolved, question is the mucosal bottleneck select for viruses with enhanced transmission fitness. Here, we characterized 300 limiting dilution-derived virus isolates from the plasma, and in some instances genital secretions, of eight HIV-1 donor and recipient pairs. Although there were no differences in the amount of virion-associated envelope glycoprotein, recipient isolates were on average threefold more infectious (P = 0.0001), replicated to 1.4-fold higher titers (P = 0.0004), and were significantly more resistant to type I IFNs than the corresponding donor isolates. Remarkably, transmitted viruses exhibited 7.8-fold higher IFNα2 (P < 0.00001) and 39-fold higher IFNβ (P < 0.00001) half-maximal inhibitory concentrations (IC50) than did donor isolates, and their odds of replicating in CD4+ T cells at the highest IFNα2 and IFNβ doses were 35-fold (P < 0.00001) and 250-fold (P < 0.00001) greater, respectively. Interestingly, pretreatment of CD4+ T cells with IFNβ, but not IFNα2, selected donor plasma isolates that exhibited a transmitted virus-like phenotype, and such viruses were also detected in the donor genital tract. These data indicate that transmitted viruses are phenotypically distinct, and that increased IFN resistance represents their most distinguishing property. Thus, the mucosal bottleneck selects for viruses that are able to replicate and spread efficiently in the face of a potent innate immune response.

For obvious reasons, viruses cannot be collected from, or studied in, humans at the time of transmission. However, by sequencing plasma virion RNA (vRNA) in the first few weeks following transmission, it is possible to enumerate and infer the genome(s) of the virus(es) that established the infection (9, 12–14). In the absence of adaptive immune pressures, HIV-1 diversifies in a random fashion, with viral sequences exhibiting a Poisson distribution of mutations and a star-like phylogeny that coalesces to an inferred consensus sequence. This consensus sequence represents the genome of the virus that initiated the infection, termed the transmitted founder (TF) virus (9). Single genome amplification (SGA) of plasma vRNA, which precludes PCR artifacts such as Taq polymerase-mediated recombination (15–18), revealed that in the great majority (~80%) of sexual transmission cases, a single TF virus establishes the new infection (9, 12, 13, 19–21). The ability to infer and molecularly clone the genomes of TF viruses has permitted their biological characterization. Initial studies showed that TF viruses use the cell-surface molecules CD4 and CCR5 as their receptor and coreceptor, and replicate efficiently

U
mucosal HIV-1 transmission | type I interferons | innate immunity | HIV-1 transmission pairs | HIV-1 transmission fitness

Significance

Effective prevention strategies are urgently needed to control the spread of HIV-1. A critical barrier to developing such strategies is the lack of understanding of the host antiviral defenses that control HIV-1 replication in the mucosa at the site of entry. Here, we characterized viruses from matched donor and recipient pairs to determine whether transmitted HIV-1 strains exhibit traits that increase their transmission fitness. Characterizing 300 limiting dilution-derived isolates, we identified several properties that enhance virus replication in the face of a vigorous innate immune response, of which resistance to type I IFNs is the most important. These results provide new insights into the HIV-1 transmission process and define possible new targets for AIDS prevention and therapy.


Reviewers: M.H.M., King's College London; and G.J.T., University College London. The authors declare no conflict of interest.

Freyly available online through the PNAS open access option.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession nos. KY111920–KY112054 and KY364480). Biological data and analysis code are archived on Zenodo, https://zenodo.org (doi: 10.5281/zenodo.216445).

1S.S.I., F.B.-R., and S.S.-M. contributed equally to this work.

2To whom correspondence should be addressed. Email: bhh@upenn.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1620144114/-/DCSupplemental.

www.pnas.org/cgi/doi/10.1073/pnas.1620144114

ES90–ES99 | PNAS | Published online January 9, 2017
in activated CD4^+ T cells but not macrophages (14, 22–25). Moreover, analysis of a comprehensive panel of infectious molecular clones (IMCs) showed that TF viruses packaged more envelope glycoprotein (Env), exhibited greater infectivity, bound to monocytes-derived dendritic cells and replicating to higher titers in CD4^+ T cells in the presence of the type 1 interferon IFNα2 than chronic control (CC) viruses (26). However, a potential confounder of these studies was the fact that TF and CC viruses were not derived from epidemiologically linked transmission pairs. To compare transmitted and nontransmitted viruses close to the time of transmission, two recent studies characterized the phenotype of viruses from matched donor and recipient pairs (27, 28). Examining various biological properties, including the sensitivity of donor and recipient viruses to IFNα2, both studies failed to identify viral traits that were indicative of enhanced transmission fitness (27, 28).

Innate immune responses, in particular type 1 IFNs, represent a potent first-line defense against many pathogens, including primate lentiviruses (29–33). Consistent with these data, treatment of rhesus macaques with pegylated IFNα2 increased the number of intrarectal challenges required to achieve systemic simian immunodeficiency virus (SIV) infection and decreased the number of transmitted founder viruses (34). Similarly, mucosal application of IFNα protected macaques from repeated intrarectal and intravaginal challenges with a simian–human immunodeficiency virus (SHIV) (35). Because type 1 IFNs are rapidly upregulated at mucosal sites of virus replication in SIV-infected macaques (36), and bioactive IFN levels are highly elevated during acute HIV-1 infection (37), we hypothesized that IFN-mediated antiviral activity contributes to the HIV-1 transmission bottleneck. To test this hypothesis, we generated a large panel of limiting dilution-derived isolates from the plasma and genital secretions of chronically infected donors and their matched recipients. Analyzing 300 such isolates, we identified a number of biological properties that are associated with increased transmission fitness, all of which serve to enhance HIV-1 replication and spread in the face of a vigorous innate immune response.

Results

Generation of Limiting Dilution HIV-1 Isolates from Sexual Transmission Pairs. Molecular cloning of HIV-1 genomes is labor intensive and thus limits the number of IMCs that can reasonably be characterized. Moreover, predicting which viral genomes are functional in chronically infected individuals is challenging, because immune escape mutations frequently incur fitness costs (38–45). Virus isolation represents an alternative to cloning, but currently cannot account for the biological variation of individual quasispecies members. Here, we used limiting dilution virus isolation to generate single virion-derived HIV-1 strains from eight sexual transmission pairs. These included four female-to-male (FTM) transmissions (subtype C) from southern Africa as well as from one male-to-female (MTF) and three men-who-have-sex-with-men (MSM) transmissions (subtype B) from the United States (SI Appendix, Table S1). In all but one case, the newly infected recipient was identified first as part of an acute infection cohort, whereas the transmitting partner was identified retrospectively. Phylogenetic analysis of SGA-derived virus sequences confirmed that all transmission pairs were epidemiologically linked (SI Appendix, Fig. S1) and showed that two recipients (CH378 and CH831) had acquired their infection from the same donor (CH742). Seven of the eight recipients were infected with a single TF virus, whereas the remaining subject (CH378) acquired at least two TF viruses (SI Appendix, Fig. S2). All subjects remained treatment naïve throughout the study.

To generate limiting dilution-derived viral isolates, plasma as well as cell-free fractions of cervicovaginal lavage (CVL) and semen (SEM) samples were end-point diluted and used to infect activated normal donor CD4^+ T cells in 24-well plates. According to a Poisson distribution, a dilution that yields positive cultures in no more than 30% of wells should contain a single infectious unit more than 80% of the time. Cultures were maintained for 20 d, tested for p24 antigen production, and virus positive wells were expanded further in normal donor CD4^+ T cells for an additional 10 d. The resulting viral stocks were used for all subsequent genetic and biological analyses.

To ensure that the limiting dilution isolates were indeed single virion derived, we sequenced all stocks before biological characterization. Briefly, 5’ and 3’ half genome were PCR amplified, MiSeq sequenced, and the resulting reads assembled to generate an isolate-specific consensus sequence. Viral reads were then mapped to this consensus sequence and the extent of genetic diversity was examined for each position along the genome. Isolates that exhibited more than 15% diversity at any one position in the alignment were considered to contain more than one variant and thus removed from further analysis. To control for the emergence of phenotypically distinct variants in the culture, we also generated limiting dilution isolates from all acutely infected subjects even though TF IMCs were available for two recipients (14, 26). Using plasma samples collected closest to the time of transmission, we generated 95 donor and 61 recipient isolates (SI Appendix, Table S1). Virus isolation from CVL and SEM samples was more challenging, because of lower viral loads, frequent bacterial and yeast contaminations, and the fact that many genital secretions were inherently cytotoxic for CD4^+ T cells (46). Nonetheless, we were able to generate limiting dilution isolates from the CVL or semen samples of three transmitting donors (SI Appendix, Table S1).

Limiting Dilution HIV-1 Isolates Are Representative of the Donor Quasispecies. To determine whether the limiting dilution isolates were representative of the viral quasispecies present in both donors and recipients, we compared all isolate-derived sequences to SGA-derived vRNA sequences amplified directly from the blood of the same individual. In phylogenetic trees of 3’ half genome sequences, isolate and plasma vRNA sequences were completely interspersed (Fig. 1A and SI Appendix, Fig. S3). To assess whether isolate and plasma viral sequences from chronically infected donors were segregated, we calculated their genealogical sorting index (gsi) (47). Two donor samples yielded gsi values that were higher than expected from random segregation (SI Appendix, Table S2A). In one case (CH212), available isolates represented only two of three diverse viral lineages present in this donor’s quasispecies, indicating limited sampling (SI Appendix, Fig. S3F). In the other case (CH728), two pairs of near identical isolate sequences indicated repeat culture of the same virus (SI Appendix, Fig. S3C). Collapsing one of these to a single sequence rendered the gsi value nonsignificant. For all other subjects, there was no evidence of compartmentalization of plasma and genital secretion isolates from donors CH492 and CH742 (SI Appendix, Fig. S3 and Table S2B).

Increased Env Content Is Not a Characteristic Feature of Transmitted Viruses. Comparing viruses from unrelated subjects, we previously reported that TF IMCs package on average 1.9-fold more Env than viruses circulating in the plasma of chronically infected individuals (26). To examine the Env content of matched donor and recipient isolates, we generated viral stocks in normal donor CD4^+ T cells, depleted these of microvesicles, purified virions using antibody-coated magnetic beads, and quantified Env by ELISA per unit of RT activity. We found that plasma isolates varied widely in the amounts of Env that they packaged, but failed to identify consistent differences between donor and recipient isolates. Recipient isolates packaged either significantly
Transmitted Viruses Exhibit Increased Particle Infectivity. We previously reported that TF viruses were more infectious than the corresponding plasma isolates, but this was not the case for the other two donors (Fig. 1B and SI Appendix, Fig. S4A). When data from all pairs were combined, no significant differences in Env content were observed between donor and recipient isolates, plasma and genital secretion isolates, and subtype B and C recipient isolates (Fig. 1C). These data indicate that mucosal transmission does not select for viruses with an increased Env content.

Transmitted Viruses Replicate to Higher Titers. The replicative capacity of viruses can influence their reproductive ratio (R₀) and thus their ability to expand an initial infection (51). Comparing IMCs from unrelated subjects, we previously failed to detect differences in the growth potential of TF and chronic HIV-1 strains (26), and similar results were reported for donor and recipient viruses from transmission pairs (27, 28). Here, we compared the replicative capacity of limiting dilution-derived isolates in normal donor CD4⁺ T cells. Using equal numbers of particles for viral input (1 ng of RT activity), we monitored the growth kinetics of a subset of isolates (n = 25) for 9 d by measuring p24 antigen in culture supernatants every 48 h. We then determined the area under the curve (AUC) and compared it with p24 values measured at individual time points. This analysis revealed a strong correlation between the AUC and p24 production at day 7 (r = 0.99, P < 0.0001). We thus used the latter as a measure of viral replicative capacity for all remaining isolates.

Transmitting donor isolates varied widely in their replicative capacity, and this was also true for some recipient isolates. However,
recipient isolates replicated on average between 1.2- and 1.7-fold more efficiently than viruses isolated from the corresponding donors (Fig. 1F and SI Appendix, Fig. S4C). These differences were significant for seven transmission pairs, with a trend observed for the eighth pair (Fig. 1F and SI Appendix, Fig. S4C). In contrast, genital secretion isolates did not exhibit an increased replicative capacity. Combining results from all transmission pairs, we found that on average recipient isolates grew to 1.4-fold higher titers than their corresponding donor isolates ($P = 0.004$), whereas no significant differences were observed between plasma and genital secretion isolates, or between recipient isolates representing subtype B and C infections (Fig. 1G). These data indicate that mucosal transmission selects for viruses with enhanced replicative capacity.

Transmitted Viruses Are Uniformly Resistant to Type I Interferons. We previously reported that TF viruses are more resistant to IFNα2 than viruses from chronically infected individuals (26, 52). However, two subsequent studies of linked transmission pairs failed to confirm this phenotype, with one study finding no differences in IFNα2 resistance between transmitted and nontransmitted viruses (27), and the other reporting transmitted viruses being more IFNα2 sensitive (28). To resolve these differences, we tested the IFN sensitivity of the limiting dilution-derived isolates, but with some experimental modifications. First, instead of testing only IFNα2, we measured the antiviral effect of a second potent inhibitor of HIV-1, IFNβ (35, 52). Second, rather than examining the effect of only a single IFN inhibitory dose (26–28, 53), we determined the half-maximal inhibitory concentration (IC$_{50}$) of both IFNα2 and IFNβ for every single isolate. This was done by treating normal donor CD4$^+$ T cells with increasing quantities of IFN, infecting them with equal amounts of virus, and culturing the cells for 7 d while replenishing IFN-containing medium. Virus replication was then measured for each IFN concentration as the amount of p24 produced at day 7 and plotted as the percentage of viral growth in the absence of IFN, which was set to 100% (Fig. 2A and B). As an independent measure of IFN resistance, we also measured viral replication at the highest IFN dose and expressed this residual replication capacity (Vres) as the percentage of viral growth in the absence of IFN (Fig. 2A and B and SI Appendix, Fig. S5 B and D).

For each transmission pair, plasma isolates from donors exhibited a wide range of sensitivities to both IFNα2 and IFNβ, whereas recipient isolates were much less variable as well as uniformly more resistant to both IFNα2 and IFNβ (Fig. 2 C and D).

![Image](Image)
(picograms per milliliter) (** P<0.001; **** P<0.0001), recipient isolates were either suppressed below the limits of p24 detection or reached Vres values of 0.01 (r=0.33, P<0.004), indicating only a partial overlap in the effects of the two IFN subtypes on the activation state, survival, and IFN stimulated gene (ISG) expression levels of CD4+ T cells.

Combining data from all transmission pairs, we found that recipient isolates were on average significantly more resistant to both IFNα2 and IFNβ than the corresponding donor isolates, exhibiting 7.8-fold (P<0.00001) and 39-fold (P<0.00001) higher IC50 values, respectively (Fig. 2 D and F). Moreover, recipient isolates had 35-fold (P<0.00001) and 250-fold (P<0.00001) greater odds of replicating at the highest IFNα2 (Fig. S6B) and IFNβ (Fig. S6D) doses than the great majority of donor viruses, respectively. These differences were not dependent on the viral subtype (Fig. 2 D and F). Thus, resistance to type 1 IFNs is a characteristic feature of transmitted viruses.

**Selection with IFNβ, but Not IFNα2, Yields Donor Isolates with a Transmitted Virus-Like Phenotype.** To search for IFN-resistant viruses in donor plasma, we treated CD4+ T cells with high doses of IFNα2 (4.0 pg/mL) or IFNβ (0.44 ng/mL) 24 h before virus isolation. The rationale was to maximally up-regulate antiviral ISGs in these target cells (without causing cell toxicity), thereby weakly for donor genital secretion isolates (IFNα2: r=0.34, P<0.05; IFNβ: r=0.40, P<0.01), indicating that IC50 and Vres provide different measures of the antiviral effect of IFNs in these compartments. Similarly, IC50 values for IFNα2 and IFNβ correlated only weakly (r=0.33, P<0.045), indicating only a partial overlap in the effects of the two IFN subtypes on the activation state, survival, and IFN stimulated gene (ISG) expression levels of CD4+ T cells.

**Fig. 3.** Biological characterization of IFNα2- and IFNβ-selected donor and recipient isolates. (A, C, E, G, and I) IFNα2 IC50 (picograms per milliliter) (A), IFNβ IC50 (picograms per milliliter) (C), viral Env content (mass ratio of gp120 and RT) (E), particle infectivity (RLU per picogram of RT) (G), and replicative capacity in CD4+ T cells (nanograms of p24 antigen per milliliter) (I) values are shown for limiting dilution-devoid viral isolates from one representative matched donor (D-CH492) and recipient (R-CH427) pair. In each panel, untreated (dark green), IFNα2-selected (light green), and IFNβ-selected (yellow) isolates from the donor (D-492) are compared with untreated (red), IFNα2-selected (dark pink), and IFNβ-selected (light pink) isolates from the corresponding recipient (R-CH427). Boxes show the interquartile range, a black bar within each box indicates the geometric mean, and whiskers span the range of the data, respectively. Asterisks indicate significant differences (determined by unpaired t test) between groups (*P<0.05; **P<0.01; ***P<0.001; ****P<0.0001). Because IFN selection did not alter the phenotype of recipient isolates, only statistical comparisons of donor isolates to untreated recipient isolates are shown. (B, D, F, H, and J) Hierarchical Bayesian regression models were used to test the population-wide fold change of IFNα2 IC50 (B), IFNβ IC50 (D), Env content (F), particle infectivity (H), and replicative capacity in CD4+ T cells (J) across all transmission pairs between untreated and IFNα2-selected donor isolates (blue), untreated and IFNβ-selected donor isolates (green), untreated and IFNα2-selected recipient isolates (gray), and untreated and IFNβ-selected recipient isolates (yellow). The fold change between untreated donor and recipient plasma isolates (red), as in Figs. 1 and 2, is also shown for comparison. A dashed vertical line marks a fold change of 1, indicating no effect. The estimated posterior probability distribution for each parameter is shown along with a table summarizing the expected fold change and the probability that the effect is <1 [or where indicated by an asterisk (*) the probability that the effect is >1].
simulating, at least in part, conditions during the earliest stages of HIV-1 infection. As a control, the same approach was used to isolate viruses from recipient plasma. As expected, the number of viral isolates recovered from pretreated CD4+ T cells was lower than from untreated CD4+ T cells, especially when IFN-β was used for selection (SI Appendix, Table S1). Thus, whereas IFN-α2 pretreatment yielded plasma isolates for all donors and recipients, only three donors and two recipients yielded IFN-β preselected plasma isolates. This was as expected because the selection dose of IFN-β was six orders of magnitude higher than the average IFN-β IC50 value of all isolates (IFN-α2 doses higher than 5.5 pg/mL caused cell toxicity). Phylogenetic analyses of full-length genome sequences revealed no evidence of compartmentalization of selected and unselected isolates (SI Appendix, Fig. S7 and Table S2C).

IC50 determinations confirmed that donor isolates from IFN-β pretreated cells were indeed more IFN resistant than those derived from untreated CD4+ T cells, whereas no changes were observed for recipient isolates (Fig. 3). For example, IFN-α2- and IFN-β-selected plasma isolates from donor CH492 had mean IFN-α2 and IFN-β IC50 values that were 7.6-fold and 31-fold higher than those of untreated plasma isolates (Fig. 3 A and C). However, resistance to one IFN subtype did not always predict resistance to the other. For donor CH492, IFN-α2 preselected isolates that were also highly resistant to IFN-α2 (Fig. 3A), whereas IFN-α2 preselected isolates were also resistant to IFN-α2 selected recipient isolates (Fig. 3B). In contrast, IFN-α2–selected isolates were 7-fold less resistant to IFN-β than IFN-β-selected isolates, and these in turn were 2-fold less resistant than untreated recipient isolates (Fig. 3D). Similar results were obtained for Vex, which showed that IFN-β2 selection did not generally yield IFN-β-resistant isolates, and that IFN-β-selected isolates were less resistant to IFN-β than untreated recipient isolates (SI Appendix, Fig. S6). Interestingly, IFN selection did not increase the IC50 or Vex values of recipient isolates, suggesting that transmitted viruses are already maximally resistant to both of these IFN subtypes (Fig. 3 B and D and SI Appendix, Fig. S6 B and D).

Having generated IFN-α2 or IFN-β preselected isolates, we next examined their biological properties. For donor CH492, IFN-α2 and IFN-β pretreatment resulted in isolates that packaged 2.0- and 3.3-fold more Env than untreated isolates, respectively (Fig. 3E). However, no significant differences in Env content were detected between treated and untreated isolates when data from all subjects were combined (Fig. 3F). However, pretreatment with IFN-α2 and IFN-β resulted in donor isolates that exhibited increased infectivity. This was observed for donor CH492 (Fig. 3G) as well as donor isolates combined (Fig. 3H). IFN-α2 and IFN-β pretreatment yielded plasma isolates that were on average 2- and 2.2-fold more infectious, respectively, than isolates obtained without selection, although neither pretreated group was as infectious as the recipient isolates. Interestingly, IFN-β pretreatment had no effect on the infectivity of recipient isolates (Fig. 3H).

Reasoning that IFN pretreatment may favor the outgrowth of viruses that replicated to higher titers, we compared the replicative capacity of IFN-selected and -unselected donor and recipient isolates. Indeed, pretreatment of CD4+ target cells with IFN-β resulted in donor isolates that replicated more efficiently than untreated viruses, both for CH492 (1.3-fold; Fig. 3J) and all donor isolates combined (1.3-fold; Fig. 3J). However, this was not observed when CD4+ T cells were pretreated with IFN-α2. Surprisingly, IFN-α2–selected isolates replicated significantly less well, both for donor CH492 (1.7-fold; Fig. 3J) and all donor isolates combined (2.1-fold; Fig. 3J). For each of the seven donors, IFN-α2 treatment selected isolates whose replicative capacity was much reduced compared with untreated isolates despite higher infectivity and in some cases greater amounts of packaged Env (e.g., CH492). These data indicate that IFN-α2 and IFN-β selection can have opposing effects on some viral properties, and that in contrast to previous suggestions (27), IFN resistance is not simply a consequence of a higher replicative fitness. As expected, IFN-α2 and IFN-β selection did not increase the growth potential of recipient isolates (Fig. 3I). Taken together, these results indicate that both IFN-α2– and IFN-β-resistant viruses are present, albeit at low levels, in the plasma of chronically infected individuals, and that in vitro treatment of CD4+ T cells with IFN-β, but not IFN-α2, selects isolates that approach the phenotype of transmitted viruses (Fig. 3 and SI Appendix, Fig. S6).

Transmitted Viruses Are More Efficiently Released from Infected Cells. We previously reported that CD4+ T cells infected with TF viruses released larger quantities of cell-free virions than cultures infected with CC viruses (54). However, because only two TF and two CC IMCs were studied, we examined this property in a much larger number (n = 127) of matched donor and recipient isolates. To quantify particle release from infected CD4+ T cells, we measured the amounts of cell-free and cell-associated p24 antigen 7 d postinfection, and used these values to calculate the percentage of p24 that was released into the supernatant. Consistent with our previous observations (54), we found that donor isolates produced on average much less cell-free virus than recipient isolates (Fig. 4), although the total

Fig. 4. Particle release capacity of matched donor and recipient isolates. (A) Donor and recipient isolates were tested for their ability to be released from infected CD4+ T cells. The percent of viral release was determined as the ratio of cell-free p24 divided by the total amount (cell associated plus cell free) of p24 7 d postinfection. Only a subset of isolates (n = 132) was tested. Values are color coded by transmission pair. (B) A hierarchical Bayesian regression model was used to estimate the population-wide fold change in the odds of release (the probability of release divided by the probability of retention) of p24 between untreated and IFN-α2–selected donor plasma isolates (blue), untreated and IFN-β–selected donor plasma isolates (green), untreated donor plasma and genital secretion isolates (purple), untreated donor and recipient plasma isolates (red), untreated and IFN-α2–selected recipient isolates (gray), and untreated and IFN-β–selected recipient isolates (yellow). A dashed vertical line marks a fold change of 1, indicating no effect. The estimated posterior probability distribution for each parameter is shown along with a table summarizing the expected fold change and the probability that the effect is <1 (analogous to a one-sided P value).
amount of p24 in these cultures was comparable. Plasma and genital secretion isolates from chronically infected donors released on average 31% and 38% of their total p24, respectively, whereas recipient isolates released 65%. In addition, IFN-selected isolates released more p24 than unselected donor isolates, although this effect was less pronounced for IFNo2 (42%) than for IFNβ (64%). Combining results from all isolates, the odds of p24 antigen being released from CD4+ T-cell cultures infected with IFNo2 and IFNβ-selected donor isolates were 1.6-fold and 3.8-fold higher, respectively, than from cultures infected with untreated donor isolates, and the odds of release were even higher (4.2-fold) for untreated recipient isolates (Fig. 4B). In contrast, no differences were observed for donor genital secretion isolates as well as for IFN-treated and untreated recipient isolates (Fig. 4B). Thus, mucosal transmission selects for viruses with a significantly enhanced particle release capacity, suggesting that the production of cell-free viruses is important in the transmission process.

Discussion
An effective AIDS vaccine will need to prevent acquisition of HIV-1 at mucosal surfaces (5). In this context, it is critical to know whether transmitted viruses possess unique biological properties that predispose them to establish new infections more efficiently. This is a controversial topic, because some studies have reported TF-specific traits (22, 24, 26, 52, 55), whereas others have failed to confirm these results (27, 28, 53, 58, 59). Some of these discrepancies are likely due to the fact that most previous analyses did not compare HIV-1 strains from transmission pairs. Using a more rigorous approach, two recent studies characterized viruses from matched donors and recipients, but failed to identify viral properties that were indicative of enhanced transmission fitness (27, 28). These findings led to the prevailing view that HIV-1 transmission is a stochastic process in which any reasonably fit virus has the potential of crossing the mucosa.

Both transmission pair studies characterized only very few donor and recipient viruses, using either infectious molecular clones (27) or peripheral blood mononuclear cell (PBMC)-derived bulk cultures (28). Reasoning that this approach had likely led to erroneous conclusions, we used limiting dilution isolation to generate a much larger number of donor and recipient viruses for phenotypic comparisons. We also used plasma rather than PBMCs for virus isolation to preclude the characterization of archived HIV-1 strains, generated genital secretion isolates for a subset of donors, and examined viral properties, such as virion release and resistance to IFNβ, which have not been previously characterized. Finally, we rendered plasma from treated donors and recipients, but failed to identify viral properties that were indicative of enhanced transmission fitness (27, 28). These findings led to the prevailing view that HIV-1 transmission is a stochastic process in which any reasonably fit virus has the potential of crossing the mucosa.

Nonetheless, to visualize the biological properties examined for all virus isolates (particle Env content, infectivity, replicative capacity, IFN IC50, and Vres values) in combination, we conducted a principal component analysis (Fig. 5 A and B). This approach revealed two major groups, one that contained all plasma and genital secretion isolates from chronically infected donors, and another that included all plasma isolates from acutely infected recipients (Fig. 5A). The fact that there was no overlap between these groups indicates that transmitted viruses are phenotypically distinct. This conclusion was confirmed when IFN-treated isolates were plotted on the same principal components (Fig. 5B). Whereas most IFNNo2-selected donor isolates grouped within the untreated donor cluster, most IFNβ-selected donor isolates overlapped the cluster of recipient viruses (Fig. 5B).

To quantify these relationships, we calculated the distance between each virus and its pair-matched recipient average of the first two principal components (Fig. 5C). As expected, untreated and IFN-treated recipient isolates were the closest to the recipient average, exhibiting only minimal variation. In contrast, untreated donor plasma and genital secretion isolates as well as IFNNo2-selected donor isolates were most distant from the average position of their respective recipient isolates and exhibited a wide distribution of distances. Interestingly, IFNβ-selected donor isolates were much closer to their recipient isolate average, consistent with IFNβ selection yielding a transmitted virus-like phenotype. We also examined the accuracy with which an isolate could be predicted to be derived from either a donor or a recipient on the basis of the seven biological properties examined (Fig. 5D). This analysis showed that IFN IC50 and Vres values predicted donor and recipient isolates with near 100% accuracy, indicating that resistance to type 1 IFNs is the most distinguishing characteristic of transmitted viruses.

If IFN resistance represents such a discerning feature, why did previous transmission pair studies miss this property? As shown in Fig. 2, chronic viruses exhibit a wide range of IFN IC50 values, indicating that random selection of just two such viruses per transmission pair by Deymier et al. (27) and Desrosiers et al. (28) could not account for real donor/recipient differences. Moreover, measuring viral release in response to a single IFN dose (26–28, 53) is likely less accurate than a formal IC50 determination. It should also be noted that the resistance of HIV-1 to IFNs is not constant during the course of infection. IFN resistance declines rapidly within the first 6 mo (52, 57), but then increases again when subjects progress toward AIDS (52, 60, 61). Thus, depending on when during the course of infection a virus is transmitted to another person, donor viruses may be more or less IFN resistant. For example, viruses from donors who transmit during acute HIV-1 infection or immediately following treatment interruption as described by Oberle et al. (28) would be expected to exhibit much higher levels of IFN resistance than viruses from subjects who transmit during asymptomatic chronic infection. In addition, PBMC cultures may reactivate latent viruses, which would be expected to exhibit IFN resistance levels consistent with their entry into the latent pool.

None of the previous transmission pair studies analyzed viral resistance to IFNβ, which produced the most pronounced donor/recipient differences. In contrast, the 39-fold higher IFNβ IC50 values of recipient isolates (Fig. 2F) is likely more biologically relevant, because many donor viruses were already more than 50% inhibited at the lowest IFNβ dose (Fig. 2F). This finding explains why the donorrecipient differences for IFNβ Vres values are so much higher than the corresponding IC50 values and why these differences are not observed for IFNNo2 (Fig. 2F and SI Appendix, Fig. 5D). Whereas both IC50 and Vres values provide an indicator of IFN resistance, they seem to describe only partially overlapping biological effects. For example, the strong correlation of both IFNNo2 and IFNβ IC50 and Vres values for donor plasma isolates likely indicates restriction by an IFN dose-driven increase in ISG activity. In contrast, the lack of a similarly strong correlation for donor genital secretion isolates suggests that some of these viruses are restricted by ISGs whose inhibitory activity is not IFN dose dependent. In addition, Vres may be a more relevant indicator of IFN resistance during the acute phase of infection when IFN levels are particularly high in the mucosa, whereas IC50 may be a more appropriate measure of systemic immune activation during later stages of infection. Future studies will need to determine the full range of IFNNo2 and IFNβ IC50 and Vres values in HIV-1-infected subjects over time.

Not all viral properties studied contributed, or contributed equally, to HIV-1 transmission fitness. For example, virion- associated Env content, although previously identified as a characteristic feature of TF viruses (26), did not differentiate donor and recipient isolates (Figs. 1C and 5D). Nonetheless, in half of
Fig. 5. Phenotypic properties distinguishing donor and recipient isolates. (A) Principal component analysis was used to visualize properties that were determined for all viral isolates (Env content, particle infectivity, replicative capacity, IFNα2 IC50, IFNβ IC50, IFNα2 Vres, and IFNβ Vres) in combination. The positions of untreated donor plasma (green), donor genital secretion (purple), and recipient plasma (red) isolates are shown on the first two components. Length and direction of arrows show how each variable contributes to the two axes. The minimum spanning ellipses that contain all data points for each group are shown in corresponding colors. (B) To visualize the effect of IFN selection, IFNα2-selected (green), and IFNβ-selected (yellow) donor isolates are plotted together with IFNα2-selected (dark pink) and IFNβ-selected (light pink) recipient isolates on the same principal components as in A. Minimum spanning ellipses encompassing the untreated donor plasma isolates (green), donor genital secretion isolates (purple), and untreated recipient plasma isolates (red) as shown in A were retained. (C) To quantify the groupings apparent in A and B, we calculated the distance of the first two principal components for each isolate to the average position of the corresponding untreated recipient isolates for that transmission pair. Isolates are color coded by transmission pairs and grouped in A and B. (D) The accuracy with which the seven viral properties predicted whether an isolate came from a donor or recipient was measured using receiver operating characteristic curves. Each line indicates the trade-off between true and false positive rates as a threshold is moved through the range of the data. Shading indicates the 95% confidence interval of the true positive rate. The dashed line indicates the expected performance of a predictor with no relationship to donor–recipient status. A line that reaches a true positive rate of 100% with a 0% false positive rate indicates that there is perfect separation between donor and recipient isolates.

The need to overcome innate defenses is also exemplified by the fact that recipient and IFNβ-selected donor isolates produced much higher levels of cell-free virus than the corresponding untreated donor isolates (Fig. 4). Type 1 IFNs induce tetherin, which prevents the release of virus particles from the plasma membrane of infected cells. HIV-1 counteracts tetherin using its Vpu protein, which binds tetherin and prevents its expression on the cell surface (62–64). However, TF Vpu proteins do not seem to counteract tetherin more effectively than the Vpu protein of chronic viruses (65). Moreover, TF-infected CD4+ T cells were shown to produce more cell-free virions even in the absence of Vpu (54). Although we have not mapped the genomic region(s) responsible for the significantly enhanced virion release capacity of recipient isolates, it is unlikely that Vpu alone is responsible. In fact, several isolates that differed significantly in their particle release function encoded identical vpu genes (SI Appendix, Fig. S7). Thus, it seems clear that other as-yet-unknown factors must be responsible for the increased particle release function of recipient (and IFNβ-selected donor) isolates, which may be critical to enhance virus spread in the mucosa during the first rounds of replication when extracellular IFN levels are particularly high.

In summary, we have identified resistance to type 1 IFNs, in particular IFNβ, as a key determinant of HIV-1 transmission fitness. This observation is consistent with previous studies showing that innate responses in the mucosa immediately following infection are inducing a potent antiviral state through the up-regulation of ISGs, many of which have anti-HIV-1 activity (63, 64, 66–73). All IFN subtype signals through the same heterodimeric receptor (30), but differences in receptor binding and/or downstream signal transduction pathways are thought to be responsible for IFN subtype-specific biological effects (74–77). IFNβ has been reported to bind the IFN receptor (IFNAR) with the highest affinity (76) and ligates the IFNAR1 chain in an IFNAR2-independent manner, resulting in the expression of a distinct set of genes (78). Either of these properties could explain its greater potency in placing selection on the transmitted virus pool. Nonetheless, IFNβ selection did not fully recapitulate the biological properties of recipient isolates despite the extremely high dose that was used to treat the target cells (Figs. 3D and 5B and C and SI Appendix, Fig. S6D). These results indicate that additional factors, possibly including IFNα2 and/or other IFN subtypes, shape the transmitted founder phenotype. Because there are a total of 13 IFNα subtypes as well as other type 1 IFNs such as IFNα, some of which inhibit HIV-1 even more potently in vitro and in animal models (79–81), it will be critical to evaluate to what extent they contribute alone, or in combination, to the HIV-1 transmission bottleneck.
Materials and Methods

Study Subjects. Transmission pairs were identified in the context of the Center for HIV/AIDS Vaccine Immunology (CHAVI-001) acute infection cohort (82). Plasma samples were obtained from eight acutely infected subjects (recipient- and donor-transmission partners (donors), with epidemiological linkage confirmed by viral sequence analysis (SI Appendix, Fig. S1). For a subset of donors, cervicovaginal lavage and semen samples were also available. Relevant epidemiological information is listed in SI Appendix, Table S1. Written informed consent was obtained from each subject and the study was approved by the institutional review boards of the University of Pennsylvania and Duke University.

Generation of Limiting Dilution HIV-1 Isolates. Plasma samples as well as cell-depleted genital secretions were end-point diluted and used to infect 1 × 10^6 positively selected, activated CD4^+ T cells (pooled from multiple donors) in 24-well plates. Cultures were maintained for 20 d, tested for p24 antigen (26), and virus positive wells were expanded in normal donor CD4^+ T cells for an additional 10 d. The resulting (one time expanded) viral stocks were used for all genetic and biological analyses.

Isolate Sequencing. Viral RNA was extracted from isolate stocks, reverse transcribed, and the resulting cDNA was used to amplify overlapping 5' and 3' genome halves in separate PCR reactions (14, 26). Amplicons were sequenced using an illumina NGS platform, and paired-end reads were assembled to generate a sample-specific reference sequence. Viral reads were mapped against the reference and the extent of genetic diversity was examined for each position along the alignment. Isolates that exhibited more than 15% diversity at any one position were judged to contain more than one variant and removed from further analysis.

Phylogenetic Analyses. Nucleotide sequences were aligned using CLUSTALW v2 (83), with ambiguous regions removed. Maximum likelihood trees with bootstrap support (1,000 replicates) were constructed using PhyML v3.1 (84) with evolutionary models selected using jModelTest v2.1.4.0 (85) or for larger datasets, RAxML using a GTR+gamma model (86). The gis was used to calculate the degree of phylogenetic association of sequences (47).

Env Content and Particle Infectivity. Particle-associated Env content was measured as described (26), with minor modifications (SI Appendix). Virion infectivity was measured using the TZM-bl reporter cell line as described (26), except for adding the fusion inhibitor T1249 (0.01 mg/mL) to prevent multiple rounds of infection.

Replicative Capacity and IFN Resistance. To determine the IFNα2 and IFNβ concentrations required to inhibit virus replication by 50% (IC_{50}), activated CD4^+ T cells were left untreated or cultured in the presence of increasing concentrations of IFNα2 (0.000067 pg/mL-5.5 pg/mL) or IFNβ (0.000067 pg/mL-0.44 pg/mL) for 24 h, infected with equal amounts of virus (1 ng of RT), and cultured for 7 d, while replenishing IFN-containing medium every 48 h. Virus replication was measured as the amount of p24 produced at day 7 and plotted for each IFN concentration as the percentage of viral growth in the absence of IFN. Residual viral replication (Vres) was measured at the highest IFNα2 and IFNβ concentrations and expressed as the percentage of replication in the absence of IFN. Replication in the untreated controls was used to determine the replicative capacity of viral isolates. Experiments were conducted in pooled CD4^+ T cells from multiple normal subjects (SI Appendix provides details).

Isolation of IFN-Resistant Viruses. To generate IFN-resistant viruses, pooled CD4^+ T cells were treated with high (but nontoxic) doses of IFNα2 (4.0 pg/mL) or IFNβ (44 pg/mL) for 24 h before the addition of end-point diluted donor and recipient plasma. Limiting dilution isolates were generated and maintained as described above.

Virion Release. Pooled CD4^+ T cells were infected with equal amounts of virus and maintained for 7 d. To quantify particle release, both cell-free and cell-associated amounts of p24 antigen levels were measured, and the percentage of p24 that was released into the supernatant was calculated.

Statistical Analyses. For intrapair comparisons, P values were derived using Welch’s unequal variance t test to compare log-transformed values. Population-wide fold changes of viral properties were estimated using hierarchical Bayesian regression models (87), which accounted for (i) nested measurements within a transmission pair, (ii) multiple transmissions from a single donor, (iii) heteroscedasticity between groups, and (iv) censored data where measurements were less than a given value. These models assumed that observations were normally distributed, with mean and variance drawn from population-level distributions (SI Appendix, SI Materials and Methods). Posterior probability distributions were estimated using Markov chain Monte Carlo sampling, implemented in Stan v2.12.0 (88) and R v3.3.1 (89).

The models were used to estimate the fold change of log-transformed functional data (or logit-transformed proportional data) between donor and recipient plasma isolates as well as donor plasma and genital secretion isolates, along with effects of HIV-1 subtype and IFNα2 or IFNβ selection. Fold change for viral properties was based on the estimated posterior mean, and probability values were calculated from the estimated cumulative posterior probability for a fold change of <1 (or >1 in the case of a posterior mean of <1) for the population-level parameters. Principal component and receiver operating characteristic analysis (90) were performed using R v3.3.1.

ACKNOWLEDGMENTS. We thank Shanki Sethi and Michelle Krysztofak for artwork and manuscript preparation and the University of Pennsylvania’s Center for AIDS Research Human Immunology, Flow Cytometry, and Viral and Molecular Core Facilities for reagents and protocols. This work was supported by the National Institutes of Health (Grants R01 AI114266, R01 AI111789, and P30 AI45008), the Center for HIV/AIDS Vaccine Immunology and Immunogen Discovery (Grant UM1 AI 1100645), and the BEAT-HIV Delta Collaboratory (Grant UM1 AI 126620). S.S., S.S.-M., H.J.B., and R.M.R. were supported by Training Grants T32 AI079782 and AI005440; P.B. is a Jenner Institute Investigator.


