RESEARCH ARTICLE SUMMARY

HIV TRANSMISSION

Selection bias at the heterosexual HIV-1 transmission bottleneck


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RATIONALE: If HIV-1 transmission involves selection for viruses with certain favorable characteristics, then such advantages should emerge as statistical biases when viewed across many viral loci in many transmitting partners. We therefore identified 137 Zambian heterosexual transmission pairs, for whom plasma samples were available for both the donor and recipient partner soon after transmission, and compared the viral sequences obtained from each partner to identify features that predicted whether the majority amino acid observed at any particular position in the donor was transmitted. We focused attention on two features: viral genetic characteristics that correlate with viral fitness and clinical factors that influence transmission. Statistical modeling indicates that the former will be favored for transmission, while the latter will nullify this relative advantage.

RESULTS: We observed a highly significant selection bias that favors the transmission of amino acids associated with increased fitness. These features included the frequency of the amino acid in the study cohort, the relative advantage of the amino acid with respect to the stability of the protein, and features related to immune escape and compensation. This selection bias was reduced in couples with high risk of transmission. In particular, significantly less selection bias was observed in men with genital inflammation and in women (regardless of inflammation status), compared to healthy men, suggesting a more permissive environment in the female than male genital tract. Consistent with this observation, viruses transmitted to women were characterized by lower predicted fitness than those in men. The presence of amino acids favored during transmission predicted which individual virus within a donor was transmitted to their partner, while chronically infected individuals with viral populations characterized by a predominance of these amino acids were more likely to transmit to their partners.

CONCLUSION: These data highlight the clear selection biases that benefit fitter viruses during transmission in the context of a stochastic process. That such biases exist, and are tempered by certain risk factors, suggests that transmission is frequently characterized by many abortive transmission events in which some target cells are nonproductively infected. Moreover, for efficient transmission, some changes that favored survival in the transmitting partner are frequently discarded, resulting in overall slower evolution of HIV-1 in the population. Paradoxically, by increasing the selection bias at the transmission bottleneck, reduction of susceptibility may increase the expected fitness of breakthrough viruses that establish infection and may therefore worsen the prognosis for the newly infected partner. Conversely, preventive or therapeutic approaches that weaken the virus may reduce overall transmission rates via a mechanism that is independent from the quantity of circulating virus, and may therefore provide long-term benefits to the recipient if transmission does occur.

Fitter viruses (red) are favored more in woman-to-man (bottom curve) than in man-to-woman (top curve) transmission. The probability that a majority donor amino acid variant is transmitted is a function of relative fitness, here estimated by the frequency of the variant in the Zambian population. Even residues common in the population are less likely to be transmitted to healthy men than to women, indicative of higher selection bias in woman-to-man transmission.

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Heterosexual HIV-1 transmission is characterized by a severe genetic bottleneck, in which infection is typically established by a single genetic variant selected from the large and diverse quasispecies typically present in the donor (1–7). The source of this bottleneck is likely mediated by multiple physical and immunologic factors that limit which virus particles can reach the genital tract, penetrate the mucosal barrier, productively infect target cells, and then traffic out of the mucosa for systemic dissemination—the sum total of which effectively blocks transmission in >99% of unprotected sexual exposures (8, 9).

One potential source of this bottleneck is the unique environment of the male and female genital tracts, which may feature different target cell populations from those the majority of viruses face in systemic infection. The envelope (Env) protein, expressed on the surface of virus particles, determines target cell specificity; thus, variations in target cell populations are likely to exert selection pressure on the virus. Indeed, selection pressure appears to favor viruses encoding Env proteins that use CCR5 as a co-receptor (10), that favor target cells more likely to be trafficked out of the gut (11), and that have higher Env concentrations (12). There is also evidence that Env proteins with lower levels of glycosylation (1–3, 13) and that are closer to ancestral sequences (14, 15) are similarly favored. Glycosylation serves as a steric shield from the humoral immune response (16), whereas the move toward an ancestral state may involve the reversion of immunological escape mutations. Because the naïve host lacks an HIV-specific adaptive immune response, these escape features are no longer necessary, and any fitness cost associated with them may become a hindrance in transmission.

If general fitness plays a role in transmission, then fitness preferences will manifest themselves in non-Env proteins as well, the possibility of which has been recently suggested by observations that transmitted founder viruses are relatively more resistant to interferon (12, 17), a phenotype that is unlikely to be dependent on Env. The ability of virus particles to grow and efficiently infect target cells has clear pathologic consequences, with in vitro measurements of viral replicative capacity correlating with viral loads (VLs) and CD4 decline in both acute and chronic infection (18–21). VL is also closely linked with the odds of transmission, raising the possibility that general in vivo viral fitness could play a significant role in the transmission process as well.

The severe nature of the transmission bottleneck suggests that the selection of the breakthrough virus is a stochastic process in which a virus with a modest growth advantage in the mucosal compartments would be more likely to succeed in establishing infection. When viewed across many linked transmission partners, such viral advantages should emerge as measurable statistical biases. We provide evidence here that the genetic bottleneck imposes a selection bias for transmission of amino acids that are consensus in the cohort and are predicted to confer increased in vivo fitness on viruses in the newly infected, immunologically naïve recipient. This bias is tempered by transmission risk factors, such as donor VL, genital inflammation, and recipient gender, and provides an overall transmission advantage for viral quasispecies that are dominated by viruses with high in vivo fitness.

**Results**

**Consensus residues are preferentially transmitted**

If some viruses with a general growth advantage are more likely to establish infection, then transmission of minority variants will be more frequent when the majority variant has lower fitness. To test this hypothesis, we collected plasma samples from 137 donors and their virologically linked seroconverting partners (recipients) a median of 46 days beyond the estimated date of infection (table SI) and compared the amino acid variants determined by Sanger sequencing at each position in the Gag, Pol, and Nef proteins. Restricting our analysis to the 228,362 residues in which a dominant (non-mixture) residue was observed at a given position in both partners, we observed a clear bias for transmission of cohort consensus (≥50%) residues, with 99.65% of donor variants that matched cohort consensus transmitted to the partner compared with 92.61% of variants that were defined as polymorphisms (216,589 of 217,348 versus 10,200 of 11,014; $P < 1 \times 10^{-16}$, Fisher’s exact test), indicating that donor minority variants are more likely to be transmitted to the partner compared with 92.61% of donor variants that matched cohort consensus transmitted to the partner compared with 92.61% of variants that were defined as polymorphisms (216,589 of 217,348 versus 10,200 of 11,014; $P < 1 \times 10^{-16}$, Fisher’s exact test), indicating that donor minority variants are more likely to be...
transmitted when the donor majority variant differs from the cohort consensus. An example of this bias was observed at Nef71, which is adjacent to the critical PxxP motif implicated in SH3 (Src homology 3) domain binding and major histocompatibility complex (MHC) class I down-regulation (22, 23). Among the 114 donors where the consensus arginine was observed as the dominant donor variant, arginine was transmitted to 112 recipients; in contrast, the dominant residue was transmitted from only 7 of 14 donors where polymorphic lysine or threonine was dominant ($P = 1 \times 10^{-6}$, Fisher’s exact test), suggesting a bias toward the transmission of consensus arginine at this site.

Within each couple, a median of 99.69% of donor sites matching cohort consensus were transmitted, compared to only 94.38% for polymorphic donor sites ($P<1 \times 10^{-16}$, signed-rank test) (Fig. 1A). Similar results were observed for sites in the donor where we observed a mixture of two amino acids in the population sequences but a single amino acid in the recipient; in these instances, consensus was still preferentially transmitted (median, 60%; $P = 1.9 \times 10^{-9}$, signed-rank test), suggesting that this result was not driven by perfectly conserved sites that may be similarly conserved in the host (Fig. 1B).

**Selection bias can be modeled by a binomial process**

To further investigate the apparent bias against the transmission of non-consensus amino acids, we modeled transmission as a binomial mixture process, which assumes that each virus in the donor quasispecies is part of a subpopulation, and each virus within that subpopulation is equally and independently likely to establish infection (see Methods). Assuming a low probability of transmission, the odds that a donor amino acid $a$ at a given position is observed in the recipient founder virus $F$ is approximately the relative frequency of $a$ in the donor quasispecies multiplied by the relative selection advantage of $a$, given by

$$
\frac{\Pr(a \in F)}{\Pr(a \notin F)} \approx \frac{f_a}{1-f_a} \times \frac{p_a}{p_n}
$$

where $f_a$ is the frequency of viruses with $a$ in the donor quasispecies, and $p_a/p_n$ is the relative advantage for transmission that viruses with $a$ have over viruses without $a$ (Fig. 1B). $p_a$ is the a priori probability that a virus of type $a$ will establish infection, and similarly for $F$. We refer to this latter ratio as the “selection bias” and say that the transmission bottleneck is “unbiased” if the ratio is one (that is, if there is no selection advantage for or against $a$). The log of this approximation yields the following linear relationship:

$$
\log\odds(a \in F) = \log\odds(f_a) + \text{bias}_a
$$

in which the log-odds that the founder virus $F$ includes a virus with $a$ is approximately the log-odds of the frequency of $a$ in the donor quasispecies, shifted by the extent of selection bias [$\text{bias}_a = \log(\frac{p_a}{p_n})$] for or against $a$. We use “increased selection bias” to refer to an increase in the absolute value of the bias, and “decreased selection bias” to mean the bias moves toward zero. The bias can be modeled as a linear function of fixed or random effects, allowing the estimation of the effects of features of interest on overall selection bias.

**Deep sequencing confirms the statistical model for selection bias**

We confirmed the above relationship by deep sequencing of viruses from five linked transmission pairs, treating $a$ as an indicator that a virus matches the donor dominant variant in the virus quasispecies at a particular site, then treating each site as an independent set of observations, to obtain sensitive estimates of the frequency of each site ($f_a$) in each donor. We observed a linear relationship between the log-odds of $f_a$ and the observed log-odds of the transmission probability, with a clear bias against transmission of non-consensus polymorphisms (Fig. 2A), consistent with Eq. 1. For example, an amino acid observed in 85% of the donor viruses will be transmitted with 85% probability if it matches cohort consensus, compared to only a 65% probability if it does not.

**Odds of transmission are predicted by factors related to viral fitness**

Although the frequency of the amino acid in the cohort (cohort frequency) and that in the donor’s quasispecies (quasispecies frequency) were weakly correlated ($\text{Spearman} \rho = 0.18$, $P < 1 \times 10^{-6}$; fig. S1), each was a significant predictor in a multivariable logistic regression model of transmission ($P < 5 \times 10^{-6}$; Table 1), consistent with cohort frequency serving as a marker of selection bias. We therefore examined the relationship between the cohort frequency of each donor amino acid and the odds of its transmission in all 137 linked transmission couples, and observed a strong continuous relationship between cohort frequency and transmission probability, both at sites where a mixture of amino acids was observed in the donor (Fig. 2B) and at sites where a single residue was observed (Fig. 2C). The continuous nature of these transmission/frequency curves is striking and indicates that even small changes in the relative cohort frequency of an amino acid will have a measurable effect on the odds that that amino acid will be transmitted.

Although cohort frequency is not a direct measure of in vivo fitness, as it may also reflect founder effects or genetic drift, it likely correlates with population-wide in vivo fitness. We thus hypothesized that features that independently predict in vivo fitness would modulate the frequency/transmission curve. First, we found that the in silico predicted protein stability costs of amino acid substitutions modulated the transmission curve, such that amino acids with minimal impact on the protein structure were most likely to be transmitted (Fig. 2D). Our in silico measure of protein stability was, by construction, biased toward consensus residues, which were most likely to match the sequence of the protein used to define the crystal structure. Nevertheless, we found that, for any given cohort frequency, an amino acid that did not affect the structure was more likely to be transmitted than an amino acid with a large impact (in the case of polymorphisms), or than a residue that occurred at a site where many other residues were equally well suited for the structure. Similarly, we found that the number of putative compensatory mutations associated with a given amino acid residue (as estimated from statistical linkage (24)) was correlated with an increased probability of transmission, consistent with such mutations being fixed by the compensations, or of compensatory mutations reducing
Fig. 2. Viral fitness modulates selection bias in heterosexual HIV-1 transmission. The odds that the donor’s amino acid will be transmitted to the recipient is a function of the relative frequency of the amino acid in the quasispecies as well as the fitness of that amino acid, as estimated here by several independent metrics. Each plot shows the empirical transmission probability (odds on a log_{10} scale) of a variant as a function of one or more parameters. Empirical transmission probabilities (solid colored lines) are estimated counting the proportion transmitted within a continuous sliding window of width 1 log-odds with respect to the feature represented on the abscissa. All log-odds values are smoothed by adding a pseudo-count. Gray lines represent a quadratic fit to the sliding window averages; shaded areas represent 95% confidence intervals estimated using the percentile-t method on 1000 multilevel bootstraps. \( P \) values are taken from Table 1 (A) or Table 2 (D to F) and represent the \( P \) value from a multilevel logistic regression model in which all features are treated as continuous variables, as described in Methods. (A) The log-odds of transmission is linearly related to the relative in vivo frequency of the variant in the donor quasispecies, with a near 1-to-1 mapping for variants that match cohort consensus. In contrast, polymorphisms are uniformly less likely to be transmitted (\( N = 8314 \) observations over 5 couples). (B) Among \( N = 3115 \) donor sites containing two–amino acid mixtures from 137 couples, the probability of transmission is also strongly predicted by the relative cohort frequency of the amino acid. Transmission probability is with respect to a randomly chosen member of the mixture; the abscissa represents the relative frequency of that amino acid in the cohort compared to the other amino acid in the mixture. (C to F) Among \( N = 228,362 \) non-mixture donor sites from 137 couples, the odds of transmission is predicted by (C) the frequency of the amino acid in the cohort, (D) the relative impact of the variant on the stability of the protein structure (low impact implies high fitness), (E) the number of covarying sites statistically linked with the variant, and (F) whether the variant is consistent with immune escape from one of the donor’s HLA alleles (only polymorphic sites are shown). See Methods for feature definitions.
the viral dosage upon exposure, then the result-
fication by a constant factor
creases the ability of each virus to establish in-
some risk factors increase the odds that any
multilevel, multivariable logistic regression mod-
fixation after transmission could not be ruled
escaped viral sequences, though rapid escape and
This may suggest a selection advantage for pre-
reduce selection bias
aive advantages (that is, bias \( a = 0 \)).
Transmission risk factors reduce selection bias
The selection bias is defined above as the relative
abilities of viruses of type \( a \) to establish infection.
Some risk factors increase the odds that any
risk factor increases the ability of each virus to establish in-
fected by a constant factor \( c \), as opposed to
wants to achieve the frequency of exposure or
the viral dosage upon exposure, then the result-
where the donor viral population will not substan-
tially affect selection bias (see notes S1 and S2 and
Thus, these results suggest that the increased risk of
This is linked with higher donor VL is attributable, at
least in part, to increased in vivo viral fitness, consistent
with the observation that donor VL is correlated with
and higher early set-point VL in recipient partners
Contrast, the reduction in selection bias experienced by female recipients and male
with GUI suggests an overall reduced selection bias, which is more conducive to in-
to lower-fitness variants.

Variable reversion rates compensate for variations in selection bias
Transmission of immune escape amino acids characterized by low fitness often results in grad-
al reversion to high-fit (consensus) amino acids (35–39), suggesting that relative reversion
rates can serve as a marker for the transmission of
low-fitness variants. We hypothesized that, if
the selection bias acts against the transmission of
less-fit polymorphisms and such bias is reduced in
female recipients, then the founder viruses of
women will include a higher number of costly
variants, which will revert more quickly than the
variants transmitted to men. We therefore col-
llected longitudinal plasma samples for 81 of the
transmission pairs at an average interval of 3
months out to 24 months after infection. Con-
sistent with the selection bias analysis, consen-
sus residues were transmitted at a greater
proportion of polymorphic donor sites to male
recipients compared to female recipients (5.98% versus 4.22%); as hypothesized, the rate at which
transmitted polymorphisms reverted to consen-
sus was significantly faster among female
(0.24%/month) than male recipients (0.12%/month)
\((P = 0.016; \text{Fig. 4A and S4}), providing further
evidence that selection bias is less stringent in
male-to-female than in female-to-male transmis-
sion (Fig. 3D).

When we included all selection bias features in a Cox proportional hazard model of reversion
from the early founder sequences through 24
months after infection, the relative hazard of all
but one feature (the structural impact of an
amino acid, which had no significant effect on
reversion) was consistent with what would be
predicted from selection bias: Selection features
consistent with increased viral fitness predicted
slower reversion, and selection features consistent
with increased susceptibility predicted faster
reversion (table S2). In addition, recipients who
were transmitted a higher number of polymor-
phisms at sites where their donor was poly-
orphic had lower early set-point VL (Spearman \( r = -0.34, P = 0.002; \text{Fig. 4B}), consistent with previous
reports that the in vitro gag fitness of early viral
isolates (19, 21), as well as the transmission of
HLA-B escape variants (40, 41), predicts early set-
point VL. This further corroborates the in vivo
fitness costs of polymorphisms that are actively
selected against during the transmission bottle-
neck and is consistent with our previous obser-
vation of a significantly lower VL in these women
early in infection (34).

Amino acid features predict odds of transmission of viral sequences and populations
The selection bias models described above result in a predicted log-odds that a given residue at a
given site will be transmitted. If we treat all sites
within an individual as independent (conditioned
on the protein), then the mean of the predicted
log-odds over a given viral sequence yields a
transmission index that estimates how likely
overall an individual sequence is to be transmitted.
Given the observed selection bias, we expect that
founder viruses will tend to have above-average
transmission indices relative to the donor quas-
Ispecies. Using limiting dilution single genome
amplification (SGA), we obtained a median of 19
(range, 4 to 27) gag sequences for each of 17
concerned the transmission indices of donor ampi-
cons to those of the linked founder
sequences. Overall, founder sequences had higher
than expected Gag transmission indices (Fig. 5,
A and B; \( P = 0.02 \)), though viruses with even
higher transmissibility indices were frequently
observed in the donor quasispecies, highlight-
ing the stochastic nature of transmission.

The observed variation in mean donor transmis-
sibility suggests that some quasispecies are,

Table 1. Donor quasispecies and cohort frequencies as additive predictors of transmission.

| Feature                     | \( \gamma \) estimate* | SE     | \( z \)      | \( \Pr(>|z|) \) | Likelihood ratio test† | \( \chi^2 \) (df) | \( \Pr(>\chi^2) \) |
|-----------------------------|------------------------|--------|------------|----------------|------------------------|-----------------|------------------|
| (Intercept)                 | 7.63                   | 0.557  | 13.70      | \(<1 \times 10^{-16}\) |                       |                 |                  |
| Donor quasispecies frequency‡| 0.56                   | 0.053  | 10.65      | \(<1 \times 10^{-16}\) |                       |                 |                  |
| Cohort frequency (cfreq)§   | 1.98                   | 0.541  | 3.66       | 2.5 \times 10^{-4} | 40.1 (2)              | 2.0 \times 10^{-9} |                  |
| cfreq²                      | 0.30                   | 0.121  | 2.45       | 0.014           |                       |                 |                  |

*Fixed effect parameters. Model was fit using logistic regression; Model fit was not improved by the addition of protein domain features or random effects. Compare to Fig. 2A. †Combined significance for sets of features was estimated using the likelihood ratio test. ‡The standardized smoothed log-odds of the frequency of the amino acid in the donor from deep sequencing. Smoothing factor is \( q = 1/50 \). §The standardized smoothed log-odds of the frequency of the amino acid in the cohort. Smoothing factor is \( q = 1/350 \).
on average, more transmissible than others and may therefore be more likely to establish infection. To test this, we obtained Gag, Pol, and Nef sequences from 181 risk-matched, chronically infected individuals who had transmitted to their partners. Overall, chronically infected partners who had transmitted exhibited higher median transmission indices than individuals who had not yet transmitted (P = 0.008; Fig. S5C), an effect that remained significant when controlling for the donor VL, recipient gender, and GUI risk factors (table S3).

### Discussion

The recognition that a single virus, or at most a handful of viruses, establishes infection led to great optimism that the defining characteristics of transmitted founder viruses would be readily identified, leading to a clear vaccine strategy. Although selection bias has been observed to act on the Env protein (1–3, 10, 12, 42), and may favor viruses that are relatively resistant to interferon-α (2), no deterministic features have yet been identified. Rather, the bottleneck appears to act at a stochastic level, favoring, though not exclusively, viruses with higher overall fitness in the context of the mucosal compartment. Here, we show that selection bias also acts on non-Env proteins and can be estimated by such generic features as the effect of a variant on protein stability, dependency of the variant on compensation, and the overall frequency of the variant in the cohort. That each of these features also predicts rates of reversion in the linked recipient further supports their role as markers of in vivo fitness. These observations confirm the hypothesis that minor fitness advantages play an important role in transmission beyond features that depend on the nature of target cells in the mucosa. Although the majority of these features likely correlate with fitness in many immunological compartments, the observation that variants linked to immune escape in the donor were less likely to be transmitted—and more likely to revert if they were transmitted—highlights the fact that in vivo fitness in chronic infection must account for immune pressures that may be absent at the site of transmission. As a result of selection bias, transmission often results in a step back in evolutionary time toward consensus, thereby slowing the rate of population-wide evolution, consistent with the low rate of population-wide evolution observed in the North American epidemic (45).

The observation that transmission risk factors reduce the selection bias further corroborates the role of fitness in transmission and provides important clues as to the mechanisms of increased risk. In the case of donor VL, whereas the quantity of virus during exposure likely plays a role in increased risk, it cannot explain the observed reduction in selection bias, suggesting that the underlying fitness typical of high VL is playing an important role as well (see note S1 and fig. S3 for simulation experiments and further discussion on this point). This in turn suggests that interventions that reduce VL without altering viral fitness (such as antiretroviral therapy in the absence of virologic escape) will have diminished effects on transmission compared to those that similarly reduce VL but additionally weaken the virus. Conversely, immunological escapes that confer a net advantage on the virus in the donor, and therefore result in higher VL, may nonetheless reduce the rate of transmission as a result of weakening the virus.

Selection bias was here measured by comparing genetic variants found in donor and recipient blood, and thus in principle could reflect selection occurring at any number of steps, including selection of viruses in the donor genital compartment or productive infection followed by reversion. Our previous SGA analysis of virus variants in the donor genital tract of Zambian transmission pairs argued against preferential selection in this compartment (44), and a deeper analysis of the reversion data also provides evidence that the selection bias observed above is

### Table 2. Multilevel multivariable logistic regression model of transmission selection bias.

| Feature | \( \gamma \) estimate* | SE | \( z \) | \( \Pr(>|z|) \) | Likelihood ratio test† | \( \chi^2 \) (df) | \( \Pr(>\chi^2) \) |
|---------|-------------------------|----|------|-------------|------------------------|-----------------|------------------|
| (Intercept) | 6.43 | 0.558 | 11.53 | <1 × 10\(^{-16} \) | | | |
| Cohort frequency (cfreq)‡ | 1.70 | 0.119 | 14.24 | <1 × 10\(^{-16} \) | | | |
| cfreq\(^2\) | 0.24 | 0.019 | 12.28 | <1 × 10\(^{-16} \) | | | |
| No. of covarying sites | 0.04 | 0.012 | 3.35 | 8.2 × 10\(^{-4} \) | | | |
| Susceptible to | −0.60 | 0.142 | −4.18 | 2.9 × 10\(^{-5} \) | | | |
| recipient HLA | | | | | | | |
| Donor Esc polymorphism : Gag§¶ | 0.00 | 0.253 | 0.00 | 0.998 | | 13.3 (3) | 0.004 |
| Donor Esc polymorphism : Pol | −0.69 | 0.197 | −3.49 | 4.9 × 10\(^{-4} \) | | 22.2 (3) | 5.9 × 10\(^{-5} \) |
| Donor Esc polymorphism : Nef | 0.48 | 0.326 | 1.48 | 0.140 | | | |
| Risk index* | 0.15 | 0.084 | 1.74 | 0.081 | | | |
| Risk index : cfreq | 0.14 | 0.067 | 2.15 | 0.032 | | | |
| Risk index : cfreq\(^2\) | 0.06 | 0.015 | 3.65 | 2.6 × 10\(^{-4} \) | | | |
| Estimated time since infection | −0.16 | 0.132 | −1.18 | 0.236 | | | |
| p17§‡ | 0.22 | 0.228 | 0.97 | 0.333 | | | |
| p17 : cfreq | 0.19 | 0.103 | 1.83 | 0.067 | | | |
| p24 | 1.72 | 0.285 | 6.03 | 1.7 × 10\(^{-5} \) | | | |
| p24 : cfreq | 0.64 | 0.116 | 5.47 | 4.6 × 10\(^{-8} \) | | | |
| p15 | 0.65 | 0.241 | 2.71 | 0.007 | | | |
| p15 : cfreq | 0.28 | 0.106 | 2.66 | 0.008 | | | |
| Protease | 0.62 | 0.307 | 2.03 | 0.042 | | | |
| Protease : cfreq | 0.15 | 0.135 | 1.09 | 0.278 | | | |
| RT | 0.62 | 0.208 | 2.98 | 0.003 | | | |
| RT : cfreq | 0.15 | 0.095 | 1.60 | 0.109 | | | |
| Integrase | 0.50 | 0.225 | 2.23 | 0.026 | | | |
| Integrase : cfreq | 0.19 | 0.105 | 1.78 | 0.076 | | | |
| Nef | 0.97 | 0.236 | 4.12 | 3.8 × 10\(^{-5} \) | | | |
| Nef : cfreq | 0.41 | 0.310 | 1.34 | 0.181 | | | |
| Nef CD4/MHC domains | 0.50 | 0.104 | 4.80 | 1.6 × 10\(^{-6} \) | | | |
| Nef CD4/MHC domains : cfreq | 0.52 | 0.133 | 3.88 | 1.0 × 10\(^{-4} \) | | | |
| Structural frequency (sfreq)‡‡ | 0.33 | 0.144 | 2.29 | 0.022 | | 24.2 (3) | 2.2 × 10\(^{-5} \) |
| sfreq : cfreq | 0.49 | 0.129 | 3.80 | 1.5 × 10\(^{-4} \) | | | |
| sfreq : cfreq\(^2\) | 0.13 | 0.029 | 4.45 | 8.6 × 10\(^{-6} \) | | | |

*Fixed effect parameters. Model was fit using multilevel logistic regression. Model fit was not improved by the addition of quadratic interaction effects between cohort frequency and protein domains or couple ID. See Methods for feature definitions. Compare to Figs. 2 and 3. †Likelihood ratio test performed between the full model and a model excluding the grouped set of features. ‡Cohort frequency was standardized (zero mean, unit variance). §Donor CTL escape features were scaled by 1 − cfreq to reflect the probability that de novo escape occurred in the donor. ¶Colon (:) signifies a multiplicative interaction. **Standardized (zero mean, unit variance) donor VL plus one if the recipient is female or a male with GUI. ††Protein domain features are treated as covariates. It is not clear whether significance implies a different relationship between cohort frequency and odds of transmission, or simply reflects variations in mean donor quasispecies diversity. §§Defined as the expected frequency of an amino acid in the cohort based on the impact of that amino acid on the protein structure (see Methods; frequency was standardized). Structural features were evaluated separately from the rest of the model because crystal structures are available for only a subset of sites. Model estimates reflect model fit using all parameters. Likelihood ratio test is against a null model including only the main parameters, but fit on sites with structural information. §§Random effects were applied to each couple. The intercept and the slope of cohort frequency were allowed to vary as a bivariate Gaussian. Maximum likelihood SDs are reported. The maximum likelihood covariance term is presented as a correlation.
not an artifact of rapid reversion in the narrow time frame of acute infection. First, nonparametric estimates of reversion rates place the rate of reversion at 0.12% per month for males (0.24% for females), a rate that holds constant after 3 months of infection. In contrast, the inferred rate of reversion peaks during the transmission window with a maximum that is an order of magnitude higher than the constant rate observed.

Fig. 3. Transmission risk factors reduce selection bias in heterosexual HIV-1 transmission. The empirical log-odds of transmission is plotted as a function of the frequency of each variant in the cohort, as defined in Fig. 2. Donor VL near the time of transmission, sex of the recipient, and presentation of GUI in male recipient partners each affect the selection bias. (A to C) Individuals are segregated by donor VL levels used in previous studies of transmission risk (27, 28). High donor VL reduces transmission selection bias in (B) female-to-male, but not in (C) male-to-female, transmission. (D to F) Male recipients appear to have increased selection bias compared to female recipients, an effect that is mitigated by the presence of GUI (D and E) or high donor VL (F). Quadratic polynomial fit (solid gray lines) and 95% confidence intervals (shaded area) were estimated as in Fig. 2. P values are estimated from a nonparametric, block-bootstrap method that tests the null hypothesis that the normalized areas under two curves are identical (see Methods for details).
during the remainder of the sample period (Fig. S4). It is unlikely that reversion rates would change so suddenly, and phylogenetic analyses of single genome amplified viral sequences from these early time points in this cohort (3, 6, 44) strongly argue against the founder virus undergoing significant rapid reversion in the first few days after transmission that must be enabled by rapid selective sweeps. Moreover, the observation in this study that features that reduced selection bias predict faster reversion is not consistent with the proposition that selection bias is an artifact of rapid reversion, as it would require reversion rates to be faster in men in the first month of infection and faster in women in the following years. These data thus argue that selection bias occurs primarily at the site of transmission, and suggest that sexual exposure frequently results in nonproductive infection of target cells until viruses with higher fitness gain a foothold for successful dissemination.

The observation that sequence features alone can predict the odds of transmission for a particular virus population highlights the importance of transmission selection bias and provides a clear mechanism for risk factors that reduce selection bias by increasing virulence or susceptibility. In addition, transmission of even subtly weaker viruses, either by increased susceptibility that allows transmission of less fit viruses from the donor quasispecies or because all variants in the donor quasispecies have lower fitness, may result in a clinical advantage for recipients (40, 41). Although the advantage of such subtle effects may be short-lived because of increased reversion that typically restores viral fitness, previous reports indicate that the replicative fitness costs of early viral sequences result in a sustained clinical advantage for the linked recipient (19, 21). Paradoxically, by increasing the selection bias at the transmission bottleneck, reduction of susceptibility would increase the expected fitness of breakthrough viruses that manage to establish infection and may therefore worsen the prognosis for the newly infected partner. Conversely, preventative or therapeutic approaches that even marginally weaken the virus may reduce overall transmission rates via a mechanism that is independent from the quantity of circulating virus and may provide long-term benefits even upon successful transmission.

Methods

Study subjects

All participants in the Zambia Emory HIV Research Project (ZEHRP) discordant couples cohort in Lusaka, Zambia, were enrolled in human subjects protocols approved by both the University of Zambia Research Ethics Committee and the Emory University Institutional Review Board. Before enrollment, individuals received counseling and signed a written informed consent form agreeing to participate. The subjects selected from the cohort were initially HIV-1 serodiscordant partners in cohabiting heterosexual couples with subsequent intracouple epidemiologically linked HIV-1 transmission (45–47). Epidemiological linkage was defined by phylogenetic analyses of HIV-1 gag/pol sequences from both partners (48). Viral isolates from each partner in the transmission pair were closely related, with median and maximum nucleotide substitution rates of 1.5 and 4.0%, respectively. In contrast, the median nucleotide substitution rate for unlinked HIV-1 C viruses from the Zambian cohort and elsewhere was 8.8± (49). The algorithm used to determine the estimated date of infection was previously described by Haaland et al. (5). All patients in this cohort were antiretroviral therapy naive. Zambian linked recipients were identified with a median (interquartile range) estimated time since infection (ETI) of 46 (42 to 60.5) days, at which time plasma samples were obtained from both the transmitting source partner (donor) and the linked seroconverting partner (recipient). All of the transmission pairs included in this study are infected with subtype C HIV-1.

A control group of 181 not yet transmitting (NYT) HIV-positive partners were selected from discordant couples enrolled for a minimum of 1 year, and matched as a group with the transmitting couples for a risk factor score derived from data on recent (i) sexual activity with the primary partner, (ii) sperm count in vaginal wash of female partner, (iii) pregnancy history, and (iv) GUI disease. These factors were used to create a risk profile for every transmission pair, and then NYT partners in each of four successive risk strata were selected in the same proportion to the representation of donor in each of the four strata, so that the two sets of HIV-positive partners were frequency-matched by their risk profile. Summary statistics for donor and NYT individuals are available in table S1.

Parameter definitions and methods

Amplification and sequencing of gag, pol, and nef genes

Viral RNA was extracted from 140 μl of plasma samples using the Viral RNA Extraction Kit.
Gag-pol population sequences were generated using nested gene-specific primers. Combined reverse transcription polymerase chain reaction (RT-PCR) and first-round synthesis was performed using SuperScript III Platinum One-Step RT-PCR (Invitrogen) and 5 µl of viral RNA template. RT-PCR and first-round primers include GOF (forward) 5'-ATTTGACTAGGAGGCTAGAA-3' and VfOR (RT-PCR and reverse) 5'-CTCTAGGAGACTCTGACCC-3'. Second-round PCR was performed using Expand High Fidelity Enzyme (Roche) and 1 µl of the first-round PCR product. Nested second-round primers include GIF (forward) 5'-TTTGACTAGGAGGCAGGAA-3' and VfBR (reverse) 5'-CTCTATACTGGATGTTTACTCGAAC-3'. Nef sequences were generated in a similar fashion, using an additional set of nested gene-specific primers. RT-PCR and first-round primers include VfI (forward) 5'-GGGTTTATTACAGGGACAGCAGG-3' and OMF9 (RT-PCR and reverse) 5'-GCACCTCAAGGCGAAGCTTTATTGAGGCTTA-3'. Second-round primers include VfI2 (forward) 5'-GCAAATCTCTGTGACAAGGTTGAAGGG-3' and OMF9 (reverse). An average of 800 RNA templates was added to the One-Step RT-PCR. Three positive amplicons per individual were pooled, representing on average 2400 input genomes, and purified with the PCR Purification Kit (QIAGEN). Purified products were sequenced by the University of Alabama at Birmingham DNA Sequencing Core. Sequence chromatograms were analyzed using Sequencher 5.0 (Gene Codes Corp.), and degenerate bases were denoted using the International Union of Pure and Applied Chemistry (IUPAC) codes when minor peaks exceeded 25% of the total peak height in both forward and reverse reads. Codons containing degenerate bases were defined as “mixtures,” whereas those with no evidence of degenerate bases or with minor peaks comprising less than 25% of the total were defined as “dominant variants.”

Sequences were codon-aligned to the HXB2 reference sequence using HIVAlign (www.hiv.lanl.gov/content/sequence/VIRALIGN/viralign.html), followed by hand-editing. For all analyses, we considered sites where a dominant variant was identified, and we excluded sites where a gap or a stop codon was present. In cases where transmission was considered, a site was excluded if a mixture, gap, or stop codon was observed in either the donor or the recipient. For transmission indices, exclusion criteria were based on the sequence in question alone—no information was taken from the individual’s partner. For a given couple, a residue was defined to have been transmitted if the same amino acid was observed in both donor and recipient. For Fig. 2B (transmission from mixtures), we limited the analysis to mixtures consisting of two amino acids in the donor, then randomly selected one of the residues to test if it transmitted. For Fig. 1B (proportion of mixtures that transmitted consens), we limited the analysis to donor mixtures consisting of two amino acids, one of which matched cohort consensus, then measured the per-couple proportion of these sites in which the consensus residue was transmitted.

Protein domains were used as covariates in the modeling. Protein domains were defined as follows: Gag was split into p17, p24, and p15; Pol was split into protease (Pr), reverse transcriptase (RT), integrase, and the Gag-Pol transframe (GagPolTF) region. The CD4 and MHC down-regulation domains of Nef, here defined as HXB2 positions 2, 17–26, 57–58, 62–65, 69–81, 154–155, 164–165, and 174–175 (49), were treated as a separate Nef domain.

454 sequencing was performed on five donors to estimate the quasispecies frequency of donor variants. For each donor, we amplified from an average of 13,000 RNA templates and obtained two overlapping PCR amplicons spanning the entire protein-coding region of the HIV-1 genome. Pooled amplicons were acoustically sheared to produce fragments between 300 and 800 bases in length. Batched, barcoded samples were amplified by emulsion PCR and sequenced on a 454 Junior (Roche) as described previously (50). To achieve sufficient detection of minor variants, we required a targeted coverage per site of 250-fold. The raw sequence output (“reads”) were assembled by Vicuna (51) and V-FAT (Broad Institute, www.broadinstitute.org/scientific-community/science/projects/viral-genomics/v-fat) to form a single genome that represents the majority base at each nucleotide position (the consensus assembly). The reads were then corrected for systematic 454 errors, such as homopolymer indels and carry forward/incomplete extensions (CAFIEs), and aligned to the consensus assembly using previously developed software RC454 and V-Phaser.

Fig. 5. Sequence-derived transmission index predicts transmission. The transmission index of a sequence was calculated as the mean of the expected log-odds of transmission for each site in the sequence, as estimated by a logistic regression model that included a second-order polynomial of cohort frequency, the number of covarying sites, and offsets and cohort-frequency interactions for each protein domain. Transmission indices were computed out-of-sample using leave-one-out cross-validation. (A and B) The transmission index for individual donor Gag amplicons compared to the transmission index for the linked founder Gag sequence. Models trained on Gag alone. (A) Transmission index for each couple. Black bar represents the median transmission index of donor sequences. Four sequences with transmission index < 0.67 were excluded as outliers. (B) Transmission index of each founder virus, median-centered against the transmission indices for linked donor sequences (P value from two-tailed Wilcoxon signed-rank test). (C) The overall transmission index of Gag, Pol, and Nef is significantly different between donor and potential source partners in risk-matched discordant couples (P value from two-tailed Mann-Whitney rank-sum test).
(30), as well as with custom programs written in Perl. These alignments were hand-refined to match the corresponding population sequences. Codon positions in which the 454-derived dominant variant differed from the Sanger sequencing-derived amino acid were excluded. The quasispecies frequency of a codon was taken to be the fraction of reads spanning a codon position that contained the codon. Amino acid frequencies were defined to be the sum of the frequencies of codons encoding the same amino acid. Sites with a read depth of less than 10 were excluded.

Limiting dilution SGA sequencing of gag was performed on 11 donor-recipient pairs, as previously described (39), but using gag-specific primers. Briefly, full-length gag was amplified from peripheral blood mononuclear cell DNA by nested PCR using the following primers: outerfor1 5′-AAGTAAGACAGAGGAGATCTCTCGAC-3′, gagR2b 5′-GCCAAGAGTGATTGAGGG-3′, innerfor1 5′-TTTCAAGAGGAGGCTGAAGAA-3′, and innerenv1 5′-GATCATGTCCTGTGGTAAGAGGAC-3′. In addition, six pairs of gag SGA sequences were taken from near-full-length genome sequences amplified by similar methods to those described previously (44). Sequences were aligned to the cohort Sanger sequence alignments.

Cohort frequencies were defined with respect to sequences taken from 375 (gag), 327 (pol), and 350 ( nef) chronically infected individuals in the Zambian cohort. The donor and NTY individuals studied here were the subset for whom gag, pol, and nef sequences were available. The cohort frequency was taken to be the proportion of individuals with a given amino acid, excluding all individuals with gaps, stop codons, or missing data at the site in question. Amino acid mixtures containing k amino acids contributed 1/k observations to each amino acid in the mixture. Cohort consensus was defined to be residues observed in a majority (≥50%) of sequences, whereas polymorphism was defined as all non-majority (<50%) residues. All residues at highly polymorphic sites in which no residue was observed in at least half the population were thus defined as polymorphisms.

Smoothed log-odds ratios were used to transform cohort and quasispecies frequencies as input variables for the logistic regression models, as well as for visualization of cohort, in vivo, and transmission frequencies. The smoothed log-odds account for finite sampling by including a prior that pushes log-odds ratios toward 0. For a probability p with smoothing factor q, p ∈ [0, 1], the smoothed log odd ratio is defined as \( \log(p + q) - \log(p + q) \), which is equivalent to adding a pseudo-count of qN if the probability is a proportion derived from N observations. Here, we use q = 1/350 for cohort frequencies and q = 1/50 for quasispecies frequencies. For visualization, empirical transmission frequencies are smoothed by the same factor as the variable to which it is being compared.

**Virologic and clinical parameters**

HIV plasma VL was determined at the Emory Center for AIDS Research Virology Core Labo

ratory using the Amplicor HIV-1 Monitor Test (version 1.5b, Roche). Early set-point VL in recipients was defined as the earliest stable nadir VL value measured between 3 and 9 months after infection and which did not show a significant increase in value within a 3- to 4-month window, as previously described (39). Donor VL was defined as the VL determined at or near the time of seroconversion in the previously HIV-negative partner.

GUI in the linked recipient was defined as at least one instance where genital inflammation or ulceration was noted on a physical examination or was treated between enrollment and seroconversion or during the 12-month period before seroconversion for individuals enrolled for more than 12 months, as previously described (59). Recipient GUI data were missing for three men and three women.

**Physics-based estimation of structural impact of point mutations**

One possible mechanism by which a mutation can reduce overall fitness is by altering the stability of the viral protein. We therefore used an in silico estimate of protein stability to estimate the impact of a point mutation on the viral structure, then used these estimates to define the energy-based expected frequency of each amino acid. In particular, we first estimated the thermodynamic stability changes caused by each of the 20 amino acids at each site using the FoldX software package (http://foldx.erg.es), as previously described (33). Briefly, the structures of p17 [Protein Data Bank (PDB) code: 2GOL] (54), p24 hexamer (PDB: 3HeE) (55), protease dimer (PDB: 3DLO) (56), RT (PDB: 1DLO) (57), integrase (PDB: 1B1S) (58), and Nef (PDB: 1EFN) (59) were mutated to the clade C consensus sequence (as defined by the consensus of our combined southern African cohort), and the FoldX optimization procedure and probability-based rotamer libraries were used to remove steric clashes and other estimation errors and to reconstruct missing side-chain atoms (60). Then the absolute changes \( \text{Δ}G \) in the Gibbs free energy were estimated using the FoldX software for each of the 20 amino acids. Next, we converted these predicted changes in free energy into a probability distribution for each site. The structurally based expected frequency \( E_{ij}(f_i) \) (structural frequency) of amino acid j and site i was defined using a normalized negative exponential Boltzmann distribution

\[
E_{ij}(f_i) = \exp(-|\text{Δ}G_{ij}|) \sum_{j=1}^{20} \exp(-|\text{Δ}G_{ij}|)
\]

This measure thus captures the relative impact on the structure of the 20 amino acids at a given site: If all amino acids result in roughly the same protein stability, then the expected frequency of each amino acid will be 1/20. We use the absolute value of the change in Gibbs free energy on the assumption that the structural stability of the viral protein is optimized in vivo. Thus, by construction, all consensus residues will have \( |\text{Δ}G_{ij}| = 0 \). Nevertheless, the expected frequency of consensus residues at different sites will vary on the basis of the predicted impact of the other residues at each site.

The smoothed log-odds of \( E_{ij}(f_i) \), with smoothing factor \( q = 1/350 \) to match cohort frequency smoothing, was then used as a feature in the models. Estimated values for structural frequency (\( E_{ij}(f_i) \)) are available in table S4.

**HLA-HIV associations and covariation**

For HLA class I genotyping, genomic DNA was extracted from whole blood or buffy coats (QIAamp blood kit, Qiagen). HLA class I genotyping relied on a combination of PCR-based techniques, involving sequence-specific primers (Invitrogen) and sequence-specific oligonucleotide probes (Innomogenetics), as described previously (61). Ambiguities were resolved by direct sequencing of three exons in each gene, using kits (Abbott Molecular Inc.) designed for capillary electrophoresis and the ABI 3130x DNA Analyzer (Applied Biosystems).

Correlations between HIV amino acids and HLA types were estimated using a phylogenetic dependence network, as previously described (24). Briefly, a maximum likelihood phylogeny was estimated for each protein using Phyml (version 3.0; (62), using the general time reversible substitution model, a γ distribution over substitution rates, and inferred nucleotide and constant site probabilities. A phylogenetically corrected logistic regression model (63), conditioned on the PhyML-inferrred phylogenies, was then used to assess the significance of an HLA allele in determining the amino acid for a given site. Forward selection was used to identify HLA alleles that correlate with a given amino acid at a given site. The model was run twice: once including other sites as covariates (covariation), and once without. To increase power, all variables were treated as binary, and all residues were tested against all HLAs (at “4-digit” subtype and “2-digit” type levels). All associations significant at \( q < 0.2 \) (corresponding to a false discovery rate of 20%) in either run are available in table S4.

An amino acid at a given site in a given individual was defined to be consistent with escape in that individual if (i) the individual expresses an HLA with an association at that site, and (ii) either the residue is positively correlated (referred to as “Adapted” in the literature) with the HLA, or any other residue is negatively correlated (referred to as “NonAdapted” in the literature) with the HLA. “Donor escape” is thus a binary variable that indicates whether the residue is consistent with escape from the donor. In multivariable models, we weight the donor escape binary variable by the probability that escape was selected in the donor, which is estimated as one minus the frequency of the residue in the cohort. A residue is susceptible to an individual if (i) the individual expresses an HLA with an association at that site, and (ii) the residue is positively correlated (referred to as “Adapted” in the literature) with that HLA. “Recipient susceptible” is thus a binary variable that indicates whether an amino acid is putatively susceptible to an HLA expressed by the recipient. We limited analyses to associations identified at \( q < 0.01 \), indicating that 99% of the associations are
expected to be nonspurious. These generally represent the strongest associations and are characterized by higher escape frequencies in individuals expressing the HLA and lower background frequencies in individuals not expressing the HLA.

Covariation among HIV sites was determined using the phylogenetically corrected logistic regression. However, rather than building a dependency network using forward selection, we liberally kept all pairwise associations significant at $q < 0.01$. Thus, a mutation at site $a$ that initiates a chain reaction of compensation at site $b$ followed by site $c$ will be picked up as two separate covarying sites in the pairwise analysis. Thus, this pairwise analysis, while identifying indirect associations, will characterize the breadth of downstream compensation events expected to result from a given point mutation. We defined the number of covarying sites of a particular amino acid at a particular site to be the number of unique positions that were significantly associated (positively or negatively) with that amino acid. The numbers of covarying sites for each amino acid are available in Table S4.

HLA and covariation associations were trained on a multicohort data set of 2066 chronically C-infected, antiretroviral-naïve individuals with HIV sequence and high-resolution HLA type information. These cohorts have been previously described, but were here merged together for the first time to yield greater statistical power. Briefly, in addition to the Zambian individuals described above ($n = 360$), the cohort consists of individuals from Durban, South Africa ($n = 968$) (64, 65), Bloemfontein, South Africa ($n = 260$) (66), Kimberley, South Africa ($n = 26$) (67), Gabarone, Botswana ($n = 386$) (68), and southern African subjects attending outpatient HIV clinics in the Thames Valley area of the United Kingdom ($n = 66$), originally from Botswana, Malawi, South Africa, and Zimbabwe (67). From these individuals, population sequences were available for Gag-p17/p24 ($n = 1897$), Gag-p15 ($n = 1135$), Pol-Pr ($n = 1335$), Pol-RT ($n = 1364$), Pol-Int ($n = 698$), and Nef ($n = 1366$). High-resolution HLA types were missing or ambiguous for at least one allele in 239 of 2066 (11.5%) non-Zambian individuals. For these, a probability distribution over haplotypes was estimated using a machine learning approach that infers haplotype frequencies, as previously described (68) and extensively validated for this purpose (70). The inferred HLA completion probability distributions were used as a prior for the phylogenetic-logistic regression analysis, as previously described (70).

**Statistical modeling of transmission selection bias**

**Infection as a binomial process**

The observation that the majority (>99%) of sexual encounters among heterosexual partners do not result in transmission, coupled with the observation that the majority of transmissions are established by a single founder virus, implies that this process is stochastic. Here, we assume that the number of transmitted founder viruses is distributed binomially, parameterized by the number of viruses in the donor genital compartment and the probability that each virus will establish infection. We then build on this model to estimate the probability that the founder virus population contains a particular genotype, and use this to model selection bias.

Suppose the average sexual encounter is characterized by $n$ viruses present in the infected partner’s genital compartment, and that the a priori probability that any particular virus will establish infection is $p$. If the probability that a given virus establishes productive infection is independent of the state of other viruses, then the total number $T$ of viruses establishing infection is a binomially distributed random variable. Of particular interest is the probability that at least one virus establishes infection, providing the rate $r$ of transmission, given by

$$r = \Pr (T > 0; n, p) = 1 - (1 - p)^n = np$$

where the approximation follows from a Taylor series expansion because the rate is small: observed rates of transmission have been reported in the range 0.01 to 0.001 (9).

**A model for selection bias**

The question of selection bias can now be phrased as the question of whether $p$ depends on the type of viruses. Suppose the population of viruses in the infected donors is grouped into two types: type $a$ and type $\overline{a}$. Such binarization can be defined arbitrarily, but in this study, we categorize the viruses on the basis of whether they contain the dominant amino acid variant at a particular site ($a$ or not $a$). Extending the above formulation, we can write $n = n_a + n_{\overline{a}}$ and $p = f_a p_a + (1 - f_a) p_{\overline{a}}$, where $n_a/f_a$ is the number of viruses of type $a$, written in terms of the frequency $f_a$ of $a$ in the quasispecies; $n_{\overline{a}} = (1 - f_a)n$ is the number of viruses of type $\overline{a}$; and $p_a, p_{\overline{a}}$ are the a priori probabilities that a virus of type $a$ or $\overline{a}$ will establish infection, respectively. Then the total number of transmitted viruses becomes $T = T_a + T_{\overline{a}}$, for binomially distributed random variables $T_a$ and $T_{\overline{a}}$, which represent the total number of transmitted viroms of type $a$ and $\overline{a}$, respectively.

In the context of transmission selection bias, a natural quantity of interest is the odds that a virus of type $a$ is in the population of viruses that establish infection, conditional on infection being established, which is approximately

$$\frac{\Pr (T_a > 0 | T > 0)}{\Pr (T_{\overline{a}} > 0 | T > 0)} \approx \frac{\Pr (T_a > 0 | T > 0)}{\Pr (T_{\overline{a}} > 0 | T > 0)} \approx \frac{n_a p_a}{n_{\overline{a}} p_{\overline{a}}} = \frac{f_a p_a}{(1 - f_a) p_{\overline{a}}}$$

where the first step follows because $a$ and $\overline{a}$ are mutually exclusive and complete and our assumptions of independence and low rates of infection imply that the probability of transmitting both $a$ and $\overline{a}$ is negligible (see note S2).

The log of Eq. 3 fits nicely into the logistic regression framework:

$$\ln \left( \frac{\Pr (T_a > 0 | T > 0)}{\Pr (T_{\overline{a}} > 0 | T > 0)} \right) = \ln \left( \frac{f_a}{1 - f_a} \right) + \ln \left( \frac{p_a}{p_{\overline{a}}} \right)$$

$$= \beta_f + x \beta_p$$

where the offset term $\beta_f$ estimates $\ln(f_a/(1 - f_a))$, $x = (x_1, \ldots, x_9)$ is a row vector of features and $\beta = (\beta_{f}, \beta_p)$ is a column vector of weights. From Eq. 4 we see that the ratio $p_a/p_{\overline{a}}$ has the effect of biasing the probability that $a$ is in the founder virus, which is otherwise determined by the frequency of $a$ in the donor quasispecies. Thus, we define the bias with respect to $a$ to be

$$\text{bias}_a = \ln \left( \frac{p_a}{p_{\overline{a}}} \right)$$

and say transmission is “unbiased” if bias$_a = 0$. By fitting ($\beta_f, \beta_p$) to the observed data consisting of all dominant variants observed in all individuals, we can estimate the effects of our L features on selection bias and test the null hypothesis that a given feature has no effect on selection bias (that is, $\beta = 0$).

A key observation of this formulation is that the log-odds that the majority variant is transmitted is equal to the log-odds of the frequency of that variant in the quasispecies, plus or minus some bias term. This is validated in Fig. 2A, where the log-odds transmission probability is equal (within the limits of estimation) to the log-odds quasispecies frequencies for consensus residues, but is shifted down for polymorphisms. This formulation further predicts that donor quasispecies frequency will be the primary determinant of transmission except in the most extreme cases of selection bias, consistent with previous reports (7).

**The effect of transmission risk factors on selection bias**

Transmission risk factors may increase the risk of transmission in one of two ways: (i) by increasing the number of viruses $n$ that have an opportunity to establish infection, for example by increasing VI or increasing the number of sexual exposures, or (ii) by increasing the probability $p$ that any one virus can establish infection, by either increasing the transmission fitness of all virus particles or increasing the susceptibility of the uninfected partner.

A key observation from Eq. 4 is that $n$ is completely absent, indicating that the number of exposures or quantity of virus present at the time of exposure will not alter the selection bias. In contrast, suppose all individual viruses are equally more likely to establish infection (for example, due to increased susceptibility in the uninfected partner), by a quantity $c$. Then, the selection bias with respect to $a$ becomes

$$\text{bias}_a = \ln \left( \frac{p_a + c}{p_{\overline{a}} + c} \right)$$

which converges toward 0 as $c$ becomes relatively large. Thus, individuals with high risk
factors will experience a reduced selection bias, as observed in Fig. 3. Our observation that VL reduces selection bias in female-to-male transmission indicates that VL increases transmission risk in this population, at least in part, by serving as a marker of increased transmission fitness, and not simply because of exposure to a higher quantity of virus particles.

The absence of neither and the simple form of Eq. 6 are the result of our assumptions of independence of transmission among virions and of a low overall rate of transmission. These assumptions warrant further exploration and are discussed in supplementary notes.

**Multilevel logistic regression (generalized linear mixed models)**

Parameter estimation and hypothesis testing under logistic regression assumes independence of observations: in this case, each site in each individual is treated independently. However, as observed in fig. S2, the relationship between cohort frequency and transmission probability differs among proteins (indicating non-independence among sites within the same protein), and as suggested by Eq. 6, all sites within a couple may experience higher or lower transmission probabilities as a result of couple-specific risk factors (indicating non-independence among sites within the same couple). In this section, we describe our specification of a multilevel, multivariable logistic regression model (also known as a generalized linear mixed model) to account for these non-independences (26). A multilevel logistic regression is similar to a standard logistic regression, but with random variables embedded in the definition of some of the coefficients. In the current context, an instantiation of a random variable is indexed by the transmission couple, allowing the coefficients to be constant among observations drawn from the same individual, but to vary randomly between individuals.

For $N$ total observations over $J$ proteins in $M$ couples, let $p_{ijk}$ be the probability that the dominant variant observed at position $i (i = 1, \ldots, N)$ in protein $j (j = 1, \ldots, J)$ in couple $k (k = 1, \ldots, M)$ is transmitted. Let $X = \{x_{ijk}\}$ be an $N \times L$ data matrix of $L$ features. The features of the model are given in Table 2 and described in detail in the “Parameter definitions and methods” section. Here, we call attention to two features of particular interest: let the column vectors $X_r$ and $X_e$ be the cohort frequencies and risk indices (the aggregate of sex, VL, and GUI, defined above) for the observations. Because we call special attention to these features, we define a new $N \times (L - 2)$ predictor matrix $W$, such that $X = [W X_r X_e].$

Then, the multilevel logistic regression is defined by level one:

$$\log\text{odds}(p_{ijk} | X) = \beta_{0ijk}^{(0)} + \beta_{ijk}^{(1)} x_{iw} + \beta_{ijk}^{(2)} x_{e} ^2$$

This model assumes a quadratic effect between cohort frequency and odds of transmission. The quadratic polynomial was chosen by first testing a model that included only protein effects and cohort frequency as a cubic on cohort frequency, then observing that the linear and quadratic effects, but not the cubic effects, were significant.) Each of the $\beta$ terms are composite, mixed-effect terms, defined by level two:

$$\beta_{ijk}^{(0)} = \gamma_{00} + W_i \Gamma + \gamma_{0x} x_r + \gamma_{0e} e_e + \epsilon_{0k}$$

$$\beta_{ijk}^{(1)} = \gamma_{10} + \gamma_{1x} x_r + \gamma_{1e} e_e + \epsilon_{1k}$$

$$\beta_{ijk}^{(2)} = \gamma_{20} + \gamma_{2x} x_r + \gamma_{2e} e_e + \epsilon_{2k}$$

where $\Gamma$ is a column vector of fixed effects, the $\gamma$ terms are fixed effects, with separate $\gamma_{0x}$ and $\gamma_{0e}$ for each protein domain, and the $\epsilon$ terms are random effects, which are normally distributed as

$$\epsilon_{jk} \sim N(0, \Sigma_0; \sigma_0, \sigma_1, \sigma_0)$$

with an independent sample taken for each couple $k$. Thus, the offset $\beta_{ijk}^{(0)}$ (Eq. 8) is determined by a grand mean ($\gamma_{00}$), a linear combination of all predictors and their coefficients ($X \Gamma$), and protein- and couple-specific offsets; the linear term $\beta_{ijk}^{(1)}$ (Eq. 9) is determined by a grand mean ($\gamma_{10}$), the risk ($\gamma_{1x}$), and protein- and couple-specific slopes; and the quadratic term $\beta_{ijk}^{(2)}$ (Eq. 10) is determined by a grand mean ($\gamma_{20}$) and the risk ($\gamma_{2x}$). Proteins are treated as fixed effects. Because the couples in our study represent a random draw from an assumed population, the couple-specific offset and slope terms ($\epsilon_{0k}, \epsilon_{1k}, \epsilon_{2k}$) are treated as random effects, with the effect for any given couple drawn from a bivariate normal distribution in which the offset and slope are allowed to be correlated. The final model learns the variance-covariance matrix $\Sigma_0$, which specifies the level of variability in slope and offset among couples, then integrates out ($\epsilon_{0k}, \epsilon_{1k}, \epsilon_{2k}$). The multilevel logistic regression was carried out using the glimr routine of the lme4 package (77) in R v3.0.2 (72). Protein- and couple-specific linear effects significantly improved the fit of the model ($P < 1 \times 10^{-8}$ by likelihood ratio test); quadratic effects did not improve the fit ($P > 0.4$ for both protein- and couple-specific terms). The final fit model is shown in Table 2.

**Transmission index**

Given a set of features $x = x_1, x_2, \ldots$ and a trained model ($\beta_0, \beta$, $\beta_{ijk}$), we can compute the expected log-odds of transmission over all sites in the sequence. Sites containing mixtures, stop codons, or gaps are ignored. To avoid overfitting, models for the transmission index were estimated using leave-one-out cross-validation, such that the transmission index for each donor was computed using a model inferred from data that did not include that couple. The NYT transmissibility scores were taken from randomly selected models learned from the leave-one-out donor-recipient training runs to ensure that any differences in observed variance were not due to differential model variance. Transmission indices were computed using a subset of features: log-odds cohort frequency, and its square; the number of statistically linked co varying sites; and corrections for subproteins (PI, P2, P5, GagPolT, protease, RT, integrase, and the Nef functional domain, as described above). SGA sequences were only available for Gag; we therefore used models trained on Gag alone for Fig. 5, A and B. For computational efficiency, random effects were not used in model inference for transmission indices.

**Statistical analyses**

**Empirical transmission probability curves**

To visualize the probability of transmission as a function of the frequency of a variant in the cohort or donor quasispecies (Figs. 2 and 3 and fig. S2), we used a sliding window approach in which we measure the observed proportion of sites that were transmitted within a given window. For example, for a given donor quasispecies frequency $f$, the empirical transmission probability corresponding to $f$ is the proportion of all variants $i$ such that $\log\text{odds}(f) = \log\text{odds}(f_{i}) < w$ for some window size $w$. The reported values are the empirical transmission frequency and the mean cohort frequency, over all observations within the window. We use $w = 1$, and only include values of $f$ with at least 20 points in the window.

To estimate 95% confidence intervals for an empirical transmission probability curve, we use a block bootstrap approach using the percentile-t method (73). Briefly, for each of $B = 1000$ bootstrap replicates, we sample with replacement the couples, then the sites within each sampled couple. We then estimate the empirical transmission probability for each cohort frequency value observed in the complete data set. The percentile-t 95% confidence interval is then estimated independently for each cohort frequency value.

In Fig. 3, we report $P$ values for comparing two experimental transmission probability curves (for example, comparing male-to-female versus female-to-male transmission). The statistic we used was the difference in the mean empirical transmission probability calculated over all cohort frequencies with at least 20 observations within the sliding window. Mean empirical transmission probabilities were calculated using the trapezoid method over the cohort and transmission frequencies output by the sliding window method. We then compared the observed difference in means to a normal distribution with mean 0 and SD $\hat{\sigma}$ to test the null hypothesis that the observed difference between the means of the two curves was zero. $\hat{\sigma}$ was estimated as the SD of the difference in means observed over the $B = 1000$ bootstrap replicates used to construct the confidence intervals.

**Reversion analysis**

The rates of reversion from polymorphism to consensus, for sites in which a polymorphism was present in both the donor and recipient,
were estimated as a function of fitness and susceptibility features. The date of reversion or loss to follow-up was determined relative to the date of the first available recipient sample. Thus, reversion rates are conditional on a polymorphism being present in both the donor sample and the first recipient sample. (Because we tracked reversion to consensus, the polymorphism in the recipient may differ from that in the donor.) The date of reversion was defined as the midpoint between the last non-mixture polymorphism and the first non-mixture consensus, minus the date of the first recipient sample. Mixtures were counted as missing data. Hazard ratios (HRs) were estimated using a Cox proportional hazard model. To account for assumed non-independence among sites sampled from the individuals, a multilevel bootstrap was performed (level 1 = sites, level 2 - individuals), with 1000 replications. Reported HR values are those from the full model. P values are computed from the SEs of the bootstrap HR values, assuming a standard normal distribution. Only sites with available structures were used in the Cox proportional hazard model (table S2); all sites were used for fig. 4. Statistical modeling was carried out using MATLAB (MATLAB and Statistics Toolbox Release 2012b, The MathWorks Inc.).

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ACKNOWLEDGMENTS

We thank all the volunteers in Zambia who participated in this study and all the staff at the Zambia Emory HIV Research Project in Lusaka who made this study possible. We would like to thank J. Allen and M. Hurlston for technical assistance and sample management; P. Farmer, Ph.D., for database design and management; and I. Brili, K. Wall, X. Li, C. Lippert, N. Fusi, J. Listgarten, and Z. Brumme for helpful discussions. This study was funded by R01 AI64066 and R37 AI52351 (E.H.) and the International AIDS Vaccine Initiative (to S.A.), and made possible in part by the support of the American people through the U.S. Agency for International Development (USAID). The contents are the responsibility of the study authors and do not necessarily reflect the views of USAID or the U.S. government.

This work was also supported, in part, by the Virology Core at the Emory Center for AIDS Research (grant P30 AI050409); the Yorkes National Primate Research Center base grant (P51RR00165) through the National Center for Research Resources PS1RR165 and by the Office of Research Infrastructure Programs/OI PS1001132; NIAID grants PO1-AI07445 (T.M.A.) and U01 AI66654 (R.S.); and Multidisciplinary AIDS Training Grant NIH T32-AI07387 (R.B.). J.P., D.T.C., and M.S. were supported in part by Action Cycling Fellowships. T.N. was supported by the International AIDS Vaccine Initiative, the South African Department of Science and Technology and the National Research Foundation through the South Africa Research Chairs Initiative, by an International Early Career Scholar award from the Howard Hughes Medical Institute, and by the Victor Daitz Foundation. E.H. is a Georgia Eminent Scientist award from the Howard Hughes Medical Institute, and by the Victor Daitz Foundation. E.H. is a Georgia Eminent Scholar. All viral sequences not previously published have been submitted to GenBank accession numbers JNO10476-JNO14465, JQ219842, and KM048382-KM050767. The data reported in this paper are tabulated in the supplementary online material.

SUPPLEMENTARY MATERIALS

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Tables S1 to S4

28 March 2014; accepted 13 June 2014

10.1126/science.1254031
HIV needs to be fit to transmit

Although you might not think it, it's hard to catch HIV. Less than 1% of unprotected sexual exposures result in infection. What then leads to transmission? Carlson et al. determined the amino acid sequence of viruses infecting 137 Zambian heterosexual couples in which one partner infected the other (see the Perspective by Joseph and Swanstrom). The authors then used statistical modeling and found that transmitted viruses are typically the most evolutionarily fit. That is, compared to other viral variants in the infected person, the transmitted virus most closely matches the most common viral sequence found in the Zambian population.

Science, this issue 10.1126/science.1254031; see also p. 136