Phylogenetic approach to recover integration dates of latent HIV sequences within-host

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Given that HIV evolution and latent reservoir establishment occur continually within-host, and that latently infected cells can persist long-term, the HIV reservoir should comprise a genetically heterogeneous archive recapitulating within-host HIV evolution. However, this has yet to be conclusively demonstrated, in part due to the challenges of reconstructing within-host reservoir establishment dynamics over long timescales. We developed a phylogenetic framework to reconstruct the integration dates of individual latent HIV lineages. The framework first involves inference and rooting of a maximum-likelihood phylogeny relating plasma HIV RNA sequences serially sampled before the initiation of suppressive antiretroviral therapy, along with putative latent sequences sampled thereafter. A linear model relating root-to-tip distances of plasma HIV RNA sequences to their sampling dates is used to convert root-to-tip distances of putative latent lineages to their establishment (integration) dates. Reconstruction of the ages of putative latent sequences sampled from chronically HIV-infected individuals up to 10 y following initiation of suppressive therapy revealed a genetically heterogeneous reservoir that recapitulated HIV’s within-host evolutionary history. Reservoir sequences were interspersed throughout multiple within-host lineages, with the oldest dating to >20 y before sampling; historic genetic bottlenecks events were also recorded therein. Notably, plasma HIV RNA sequences isolated from a viremia blip in an individual receiving otherwise suppressive therapy were highly genetically diverse and spanned a 20-y age range, suggestive of spontaneous in vivo HIV reactivation from a large latently infected cell pool. Our framework for reservoir dating provides a potentially powerful addition to the HIV persistence research toolkit.

HIV, latency, reservoir, phylogenetics, evolution

HIV, like all retroviruses, integrates its genome into that of the infected host cell. Although actively infected cells typically die as a result of viral cytopathic effects or immune-mediated elimination, a minority [broadly estimated as one in every million resting CD4+ T cells (1, 2)] harbor integrated HIV DNA in a state of reduced transcriptional activity (or quiescence) over long periods (3–5). Termed latently infected cells or latent HIV reservoirs, these represent the major barrier to a cure as they can persist for years and can reactivate at any time to produce infectious virions (5–10). It is for this reason that combination antiretroviral therapies (cART), which do not act upon latent HIV reservoirs, need to be maintained for life.

Characterization of latent HIV sequences provides information about reservoir stability, distribution, and dynamics (11), which can in turn inform HIV elimination strategies. It is now established that a majority of sexually acquired HIV infections are initiated by a single transmitted/founder variant, from which descendant viral populations rapidly reaccumulate genetic diversity (12–20). Concomitantly, establishment of latent HIV reservoirs begins within hours or days following infection and continues as long as the virus is actively replicating within the host (21, 22). Given that latent HIV genomes can persist for decades, either in the original cell or clonal descendants thereof (23, 24), the reservoir should constitute a genetically heterogeneous archive of within-host HIV evolution, even after years of cART (25–27). However, while numerous studies have confirmed that the HIV reservoir is genetically diverse (11, 28–35), our knowledge of the within-host ancestor–descendant relationships of these sequences remains limited, because few studies (ref. 36 being a notable exception) have interpreted reservoir diversity in the context of HIV’s within-host evolutionary history. In particular, it remains unclear whether HIV sequences sampled from the reservoir during long-term suppressive cART truly recapitulate within-host HIV evolution or whether dynamic processes such as homeostatic proliferation (33), clonal expansion (11, 37–41), and/or preferential elimination of certain latently HIV-infected cells skew reservoir sequence composition over time. These knowledge gaps persist due to the challenges of studying within-host HIV dynamics over long timescales, combined with a lack of methods to infer latent HIV integration dates. While phylogenetic principles have been used to identify viral reservoirs (25), and techniques have been developed to detect latent lineages (42) and to quantify their turnover (43),

Significance

Studies characterizing within-host latent HIV sequence diversity have yielded insight into reservoir dynamics and persistence. Our understanding of these processes, however, can be further enhanced if reservoir diversity is interpreted in context of HIV’s within-host evolutionary history. Approaches to infer the original establishment (i.e., integration) dates of individual within-host latent HIV lineages would be particularly useful in this regard. We describe a phylogenetic framework to infer latent HIV ages from viral sequence information and apply it to latent HIV sequences sampled up to 10 y on suppressive therapy to yield insights into HIV reservoir dynamics. The ability to infer within-host latent HIV ages from sequence information has broad potential applications that may advance us toward an HIV cure.

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latent HIV deposition times have generally been estimated by assessing genetic similarity of these sequences to pre-cART plasma HIV RNA sequences (e.g., refs. 30 and 36). However, this approach is limited by the ability to previously sample similar sequences within-host. To our knowledge, no studies have directly leveraged information about within-host HIV evolutionary rates to infer latent HIV sequence ages.

As evolution of a given HIV lineage effectively ceases upon integration (and only resumes once a new round of virus replication ensues), latent HIV sequences will display a characteristic discordance between their sampling date and their “true” (older) age based on their genetic divergence from the transmitted/founder virus. Inference of the latter date, along with a host-specific evolutionary rate, would thus allow a reservoir’s “true” age to be inferred from its sequence. Within-host ancestor-descendant relationships can be phylogenetically reconstructed using serially sampled HIV sequences (44, 45); moreover, by “fixing” tree tips at their respective sampling dates, phylogenies can be rescaled chronologically, allowing rates of evolution to be estimated directly (46). As the rate of evolution remains fairly consistent among actively replicating within-host HIV lineages, the divergence of descendant lineages from the transmitted/founder virus can be approximated, at least over the initial years of infection, by a “strict” molecular clock model (36, 47, 48).

Based on these principles, we developed a phylogenetic framework that recovers the integration date of latent HIV lineages by reconstructing within-host HIV evolutionary history from virus sequences sampled pre-cART.

**Results**

**Reconstructing Integration Dates Phylogenetically.** Phylogenetic inference of HIV evolutionary rates from time-stamped between-host HIV sequence datasets has been used to date key events in the pandemic’s history (49, 50), to reconstruct putative transmission histories (44, 51, 52), and to infer unknown sequence ages (53). We adapt these techniques to the within-host context with the goal of estimating latent HIV sequence ages (Fig. 1). We begin by inferring a maximum-likelihood phylogeny (54) from plasma HIV RNA sequences (“training data”) along with sequences whose ages are known (“censored data,” for example proviral DNA sequences subsequently sampled during suppressive cART) (Fig. 1A). We then identify the root location that optimizes the relationship between evolutionary distance and sampling time in the training data (Fig. 1B) and calibrate a strict molecular clock by fitting a linear model that relates their root-to-tip distances to their sampling dates, given by

$$D = \mu T + a,$$

where the linear model estimate $D$ denotes the genetic distance from the root, predictor variable $T$ denotes the sample collection date, $\mu$ denotes the evolutionary rate, and $a$ denotes the $y$-intercept (Fig. 1C). We then test for the presence of a molecular clock (i.e., evidence that sequence divergence increases over time) by comparing the model against the null model of zero slope (where $D$ is constant over time). Linear models need to fulfill two criteria to pass quality control: first, the Akaike information criterion (AIC) (55) needs to be at least 10 units lower than that of the null model, and second, the 95% CI of the model-estimated root date needs to contain or precede the first sampling date. For root estimates, the established (i.e., integration) date of each censored sequence $x$ is estimated by

$$x \sim \mu T^2 + \mu T + a / \mu,$$

where $T$ is the date of integration, $\mu$ is the evolutionary rate, and $a$ is the intercept (Fig. 1D). The uncertainty in $T$ is given by

$$T \sim \frac{\mu T^2 + \mu T + a}{\mu} / \nu,$$

where $\nu$ is the number of training sequences, and $D$ is the mean genetic distance and collection date of the training sequences, respectively (56).

**Framework Proof of Concept.** To test the concept, we simulated 1,000 phylogeny trees with 100 tips each under a strict molecular clock, censored the sampling dates for 50% of sequences at random, and inferred linear models from the remainder (representative dataset in Fig. 2A–C). All linear models yielded ΔAIC values above the predefined threshold (mean ΔAIC 187); however, 39/1,000 (4%) returned root date estimates whose 95% CIs were later than the first training time point and were thus rejected, yielding a 96% success rate. Linear models fit the training data very well: the overall scaled mean absolute error (MAE), calculated as the grand mean of the absolute error between predicted and true sampling dates of the training data, was 0.034 (i.e., 3.4%). That is, for datasets spanning a 1-y period, the linear model recovered training data sampling dates to within an average of 12 d. Dates of censored sequences were similarly recovered with minimal error: The overall scaled mean absolute difference (MAD, calculated as the grand mean of the absolute difference between predicted and true sampling dates of censored sequences, scaled by total training data timespan) was 0.037, with no evidence of distributional asymmetry using a nonparametric binomial test (57) (all 961 simulations returned $P$ values below the significance cutoff; see Materials and Methods) (Fig. 2D).

To test the framework’s performance on real HIV sequence datasets sampled at various depths, frequencies, and timespans, we applied it to 424 longitudinal partial HIV RNA env sequences.
Fig. 2. Framework proof of concept using simulated and published HIV sequences. (A) Representative rooted tree relating simulated longitudinal within-host plasma HIV RNA sequences with 50% of tips randomly assigned as training data (circles colored by sampling time point) or censored for molecular dating (open black circles). (B) Resulting linear model with ancestor traces overlaid. (C) Inferred dates of censored sequences; arrow indicates baseline sampling date. (D) Density plots of normalized error distributions (expressed as the absolute difference between predicted and true sampling dates of the censored sequences, scaled by the total dataset timespan, where 1 and 1 represent 100% and 100%, respectively) from 100 (of 961) successful simulations selected at random. (E–G) Same as A–C, but for within-host plasma HIV RNA sequences from LANL participant 13654 where 50% of tips were randomly assigned as training data. (H) Density plots of normalized error distributions for all six successful LANL RNA datasets. (I–K) Same as A–C, but for LANL participant 821 with HIV RNA and DNA sequences treated as training and censored, respectively. (L) Density plots of normalized differences between HIV DNA predicted and sampling dates for the six successful LANL RNA/DNA datasets.
The Los Alamos National Laboratory HIV sequence database (LANL) (58) from eight individuals with known infection dates (median 40 sequences per individual, collected at a median of 4.5 time points over a median 2.2 y) whose within-host sequence diversity was consistent with a molecular clock (SI Appendix, Table S1). After censoring 50% of sequences at random from each dataset, linear models could still be calibrated for six of them (representative data shown in Fig. 2 E–G; summary in SI Appendix, Table S1). In part because real datasets were on average smaller and spanned shorter time periods than simulated data, scaled prediction errors were higher. Nevertheless, the model predicted censored dates to within an average of 15% (SD 6.3%) of the total timespan of each dataset, where prediction errors reflected overall model fit to the training data (Pearson’s correlation between model MAE and MAD was $R = 0.98, P = 0.007$, with no significant distributional asymmetry (Fig. 2H and SI Appendix, Table S1). These observations demonstrate that the framework can be applied to sparse data (e.g., for individual 13333, a linear model was calibrated from only 19 training sequences sampled over 1.4 y).

We then applied the framework to seven published datasets (58) featuring plasma RNA and proviral DNA sequences sampled at various times pre- and post-treatment (median 142 sequences per individual collected at a median 11 time points over a median 5.8 y), although none featured HIV DNA sampled during long-term suppressive cART (SI Appendix, Table S1). Thus, while the majority of HIV DNA sequences should represent contemporary within-host strains, we hypothesized that a minority would represent latent genomes (59). To test this, plasma HIV RNA sequences were used as training data to reconstruct proviral DNA archival dates, with the expectation that datasets containing latent genomes would yield left-skewed date difference distributions due to latent genome archival dates preceding their sampling dates. Of the seven datasets, six (including three with only three training time points) yielded successful linear models. (Representative data are shown in Fig. 2 I–K and a summary is given in SI Appendix, Table S1.) Overall, HIV DNA dates were predicted to within an average of 16% (SD 5.9%) of the total training data timespan where MAD again correlated strongly with overall model MAE (Pearson’s $R = 0.98, P = 0.00048$; SI Appendix, Table S1). Moreover, significantly left-skewed date difference distributions were noted in two individuals, consistent with the presence of latent lineages (SI Appendix, Fig. S1 and Table S1). These results demonstrate that HIV sequence ages can be reconstructed from a wide range of datasets from individuals with various clinical histories, and that the framework can detect the expected presence of latent lineages.

**Dating the Latent HIV Reservoir.** As the framework is designed to date HIV sequences within (or released from) HIV reservoirs, we applied it to reconstruct integration dates of HIV sequences isolated from two individuals who had maintained prolonged plasma viral load (pVL) suppression on cART. Participant 1 was diagnosed with HIV in August 1996 and did not initiate a regimen that suppressed pVL to $<50$ copies HIV RNA per mL until August 2006 (Fig. 3A). Viremia rebounded in the fall of 2007 after an unsuccessful regimem change, but suppression was reacieved in April 2008 and sustained to the present day with the exception of one “blip” to 76 copies per mL in September 2015. Using single-genome amplification, we collected 102 HIV RNA nef sequences from 14 pre-cART plasma specimens spanning August 1996 to June 2006 as training data. We also collected 42 nef sequences sampled at four time points post-cART for molecular dating; these included proviral DNA sequences retrieved from whole blood collected in July 2011 and peripheral blood mononuclear cells (PBMC) in June 2016 (5 and 10 y post-cART, respectively) and HIV RNA sequences sampled during the September 2007 pVL rebound and the September 2015 viremia blip. These data included eight instances where identical HIV RNA sequences were isolated from the same or temporally adjacent plasma samples and eight instances where identical sequences were isolated from putative reservoirs (including two instances where the same sequence was isolated from the plasma viremia event in 2007 and PBMC sampled in 2016) (SI Appendix, Fig. S2A).

As identical HIV sequences are not phylogenetically informative, a rooted maximum-likelihood phylogeny was inferred from unique HIV RNA nef (training data) and 34 unique putative reservoir sequences (censored data). The characteristic genetic bottlenecks that typify within-host HIV evolution (15, 16, 20) are clearly visible in the “ladder-like” tree shape (48, 60) (Fig. 3B) and in the temporally ordered plasma HIV RNA nef amino acid alignments (SI Appendix, Fig. S2B). Training data root-to-tip patristic distances also correlated strongly with sampling times (Pearson $R = 0.92, P < 9.8 \times 10^{-35}$), indicative of a molecular clock. Consistent with the reservoir as a genetically diverse archive of within-host HIV evolution [where it would be expected that reservoir sequences would be dispersed throughout a phylogeny of viruses sampled over time from an individual (25)], censored sequences were embedded within multiple within-host lineages and exhibited overall diversity comparable to that of pre-cART plasma HIV RNA sequences sampled over a decade (mean patriscic distances of 0.12 vs. 0.095 expected substitutions per base, respectively). The linear model fit the training data well, particularly in the early years (overall $\Delta$AIC = 172; MAE = 1.1 y), yielding an estimated nef mutation rate of $3.9 \times 10^{-5}$ substitutions per base per d and an estimated root date of August 1995 (Fig. 3C). Evolutionary rate and root date reconstructions were confirmed using established Bayesian methods (grand mean rate $4.1 \times 10^{-5}$; 95% highest posterior density (HPD) $3.1 \times 10^{-5}–5.2 \times 10^{-5}$) substitutions per base per d; grand mean root date December 1995 (95% HPD July 1995–May 1996).

Further supporting the reservoir as a within-host HIV evolution archive, unique proviral DNA sequences sampled in 2016, after nearly 10 y of suppressive cART, dated to between
1997 and 2007, making them an average of 14 y old (Fig. 3D). Moreover, consistent with multiple ancestral HIV lineages contributing to viremia recrudescence upon therapy discontinuation, four of the five HIV RNA sequences isolated from the 2007 viremia rebound dated to distinct pre-cART time points (one to January 2002 and the others to early 2006). The single HIV RNA sequence isolated from the 2015 plasma viremia blip dated to August 2006, around the time of cART initiation, suggesting the blip originated from spontaneous reactivation of a latent HIV lineage seeded just before therapy. While the linear model returned 2009 integration dates for two sequences, in both cases their 95% CIs overlapped with the last period of uncontrolled viremia (SI Appendix, Fig. S3). Overall these results suggest that HIV evolution was not ongoing during suppressive cART in this participant, at least not in blood.

Participant 2 was diagnosed with HIV in April 1995 (Fig. 4A). In July 2000 he initiated an incompletely suppressive dual ART regimen that was maintained until August 2006, when he initiated cART. Plasma viremia was suppressed to <50 copies per mL by December 2006 and maintained until May 2011, after which intermittent viremia blips occurred, the highest of which was in March 2013 (1,063 copies per mL). We collected 47 plasma HIV RNA nef sequences from four pre-ART time points between February 1997 to December 1999 and 100 plasma nef sequences from two time points between April 2001 and August 2006 while on dual ART, for use as training data. An additional 30 nef sequences sampled up to 10 y post-cART, including plasma sequences from the March 2013 viremia blip and PBMC-derived HIV DNA sequences sampled in August 2016, were isolated for molecular dating. We noted 16 cases where identical HIV sequences were isolated from the same or different time points, including one case where a sequence isolated from the 2013 plasma viremia blip exactly matched one isolated from plasma HIV RNA in 2005 (SI Appendix, Fig. S4A).

The phylogeny inferred from 119 unique plasma HIV RNA and 18 putative reservoir sequences revealed two initial viral lineages that underwent a severe genetic bottleneck following dual ART (Fig. 4B and SI Appendix, Fig. S4B). Again, training data root-to-tip distances correlated strongly with sampling times (e.g., Pearson $R = 0.61$, $P = 3.4 \times 10^{-7}$ for pre-ART period) and putative reservoir sequences were dispersed throughout all lineages. These included one ancestral subclade branching close to the root that included five unique sequences isolated from both reservoir samplings, whose most recent common ancestor (MRCA) gave rise to the clade that disappeared from circulation after dual ART. Two linear models were trained, one for the ART-naive period (ΔAIC = 16; MAE = 0.85 y) which yielded an estimated mutation rate of 1.1 \times 10^{-3} substitutions per base per d and an estimated root date of February 1993 and the second for the dual-therapy period (ΔAIC = 14; MAE = 3.0 y; estimated mutation rate 3.0 \times 10^{-4} substitutions per base per d) (Fig. 4C).

Evolutionary rate and root date estimates were again consistent with those estimated using Bayesian methods (grand mean rate 8.8 \times 10^{-4} [95% HPD 6.8 \times 10^{-4}–1.1 \times 10^{-3}] substitutions per base per d; grand mean root date February 1994 [95% HPD January 1993–March 1994]).

Reservoir sequences were dated using the linear model inferred from the lineages in which they resided, yielding integration dates that preceded collection dates by up to 21 y (Fig. 4D). Two observations are notable. First, the five sequences in the most ancestral subclade, which included sequences sampled from the 2013 viremia blip as well as from proviral DNA in 2016, dated to between December 1994 and May 1996, before the participant’s first plasma sampling. Detection of archival sequences in viremia blips is consistent with the reservoir harboring reactivation-competent viral lineages that date back to transmission. Second, sequences from the 2013 viremia blip were genetically diverse and dated to between 1994 and 2010, suggesting that spontaneous in vivo reactivation of numerous ancestral HIV lineages led to this viremia episode.

**Framework Is Robust to Minimal Training Data.** Training linear models from ~100 HIV sequences sampled over a decade is not feasible in most cases, so we tested model robustness to sampling depth and frequency. We began by evaluating the framework’s ability to retrieve known plasma HIV RNA dates using progressively sparser training data (“RNA censoring validation”). To do this, we randomly subsampled training time points from participant 1’s data to generate 1,096 new training datasets (featuring all 91 combinations comprising two training time points, 100 datasets each comprising between 3 and 11 training time points, all 91 combinations comprising 12 training time points and all 14 combinations comprising 13 training time points), where the remainder were censored. As all datasets contained the same sequences, a single phylogeny was inferred where the root location (and thus the resulting linear model) differed for each dataset, depending on which sequences were designated “training” vs. “censored.” Overall MAE distributions (here, expressed as the difference between known and model-predicted sampling dates for all sequences) as well as their respective concordance coefficients (61) are shown in Fig. 5A–C. To provide context on maximal model performance, the full (14 time points) dataset yielded an overall MAE of 1.3 y (Fig. 5B) and a concordance coefficient of 0.90 (Fig. 5C).

These results demonstrated that the framework could be reliably applied to as few as three training data time points, and sometimes even two. For example, 86% of linear models trained on three time points passed calibration (Fig. 5A) and yielded a median MAE only 14% more than the full dataset (1.4 vs. 1.3 y, respectively; Fig. 5B) and a median concordance coefficient only 4.4% lower than the full dataset (0.86 vs. 0.90 respectively; Fig. 5C).
5C). Remarkably, linear models trained on only two time points (and an average of only 13 training sequences) were still successful in 31% of cases. In general, however, models trained on more time points, where the first was as early as possible and where sampling spanned the widest temporal range, were more likely be successful (Fig. 5 D–F). For example, all models where the first training time point was before 1998 (i.e., within ~3 y of the estimated root date) and the last was in 2000 or later were successful (Fig. 5F). Our observation that failed models returned a 2.1-fold higher grand mean MAE than passing ones further corroborates our model assessment criteria (Materials and Methods and Fig. 5 D and E).

The framework’s ability to reconstruct sequences of unknown age from sparse training data (“RNA subsampling validation”) was similarly robust (SI Appendix, Fig. S5). The above-described 1,096 temporally subsampled training datasets were used to reconstruct integration dates of putative reservoir sequences: here, however, unused plasma HIV RNA sequences were removed such that phylogenies inferred from fewer training data would be progressively sparser. Again, reservoir establishment dates were reliably retrieved. For example, 84% of linear models trained on only three time points passed calibration (SI Appendix, Fig. S5A) and yielded reconstructed sequence ages with a median MAD of only 1.5 y and a median concordance coefficient of 0.87 from those estimated using the full training dataset (SI Appendix, Fig. S5 B and C). For context, 1.5 y represents 7.5% of the total (20-y) follow-up period for this participant. Overall, results indicate that our framework is robust to limited sampling.

**Framework Is Robust to Rooting Strategy.** Rooting phylogenies is inherently difficult. To validate our primary rooting strategy (RTT; see Materials and Methods), we outgroup-rooted all datasets on the HIV-1 subtype B reference strain HXB2 (OGR). Outgroup rooting the eight LANL plasma HIV RNA datasets identified two that did not subsequently exhibit clock-like signal; thus, for the remaining six we censored 50% of the sequences at random and used the remainder as training data. While only three of the resulting linear models passed, these yielded dates highly concordant with RTT-recovered ones (0.98 overall; Fig. 6D). Outgroup rooting the seven LANL HIV RNA/DNA datasets sets, which were generally larger than the HIV RNA datasets, yielded five successful linear models (compared with six of seven by RTT), which returned dates highly concordant with RTT-estimated ones (0.97 overall; Fig. 6B). For participants 1 and 2, OGR yielded slightly earlier MRCA dates than RTT (April 1995 and April 1990, respectively; SI Appendix, Fig. S6). This is
likely because, while RTT is somewhat biased toward placing the root on a branch relating the earliest samples (which is unlikely to be long), OGR root placement has a more uniform probability distribution along the tree (as the outgroup is relatively genetically distant). Folding on a longer branch will thus place the root further back in time relative to the earliest samples on its tips. Nevertheless, OGR-estimated latent HIV dates were highly concordant with RTT-estimated ones (0.96 overall; Fig. 6C).

OGR models were also generally robust (SI Appendix, Figs. S7 and S8). In the RNA censoring validations, OGR models required slightly more training time points (e.g., whereas 86% of RTT models passed validation, only 73% of OGR ones did so; SI Appendix, Fig. S7 A and E) and returned higher error (e.g., when trained on the full dataset, MAE was 1.9 y for OGR compared with 1.3 y for RTT) (SI Appendix, Fig. S7 B and F), yielding median concordance coefficients of 0.79 vs. 0.90, respectively (SI Appendix, Fig. S7 C and G). OGR models also yielded lower ΔAIC values than RTT ones (SI Appendix, Fig. S7 D and H).

Nevertheless, successful OGR models yielded comparable distributions regardless of training dataset size. For example, the median MAE of OGR models trained on only three time points was only 2.7% higher (1.95 vs. 1.89 y, respectively), and the median concordance coefficient only 4.5% lower, than that of the full dataset (0.76 vs. 0.79) (SI Appendix, Fig. S7 F and G). In the RNA subsampling validations, OGR performed nearly as well as RTT (SI Appendix, Fig. S8). For example, compared with dates recovered using the full training dataset, the median MAD of latent HIV ages recovered using only three training time points was 1.8 y for OGR and 1.5 y for RTT (SI Appendix, Fig. S8 B and F), yielding median concordance coefficients of 0.85 and 0.87, respectively (SI Appendix, Fig. S8 C and G). These results indicate that the framework is robust to rooting strategy.

Additional Validations. Since relatively few phylogenetic studies have focused on nef, we wished to confirm that evolution in this region is representative of the rest of the HIV genome. To do this we analyzed a published dataset of 16 nearly full genome HIV sequences sampled from an ART-naive individual over ∼3.5 y (62). We confirmed that, for each within-host HIV sequence, the number of differences from the baseline consensus within nef correlated strongly with that in the remainder of the HIV genome (Pearson $R = 0.96$, $P = 2.0 \times 10^{-8}$). Moreover, a partition-homogeneity test applied to nef vs. env sequences yielded $P = 0.44$, indicating that we cannot reject the null hypothesis that these regions evolved under the same evolutionary process. These observations support within-host variation in nef as being representative of the HIV genome.

Finally it is possible that, even during untreated HIV infection, a minority of plasma HIV RNA sequences may have been recently released from the reservoir. Such sequences would display reduced relative evolutionary rates and could reduce model fit. However, tests for such sequences within participant 1’s training data (Materials and Methods) identified only one such outlier, and removing it from the analysis yielded latent HIV ages that were 99% concordant with original estimates (SI Appendix, Fig. S9). These observations indicate that such sequences are relatively uncommon during untreated HIV infection and that their presence does not substantially affect framework performance.

**Discussion**

The ability to infer latent HIV integration dates would greatly enhance our understanding of reservoir longevity and dynamics. We developed a simple phylogenetic framework to do this and began by demonstrating its ability to recover known sampling dates on a variety of within-host datasets with acceptable error rates (e.g., in participant 1’s HIV RNA training data spanning a 10-y period, the model recovered known sampling dates to within an average of 1.3 y of their actual values; Fig. S5B). Subsequent application of the framework to date HIV sequences recovered after up to ~10 y on cART supported several long-held notions. Results confirm proviral DNA in PBMC as a genetically heterogeneous archive recapitulating within-host HIV evolutionary events dating back to transmission (Figs. 3 and 4) (36). Isolation of 18-y-old HIV sequences from a viremia blip on otherwise suppressive cART (Fig. 4D) supports the idea that such events can be explained by antigen-driven reactivation of latently infected cells (63) and extends a report of a long-lived (>17 y) defective HIV sequence (64) by demonstrating that replication-competent viral lineages can similarly persist long-term before spontaneously reactivating. Isolation of HIV RNA sequences ranging from 3 to 18 y old from this same viremia blip extends a report that in vivo treatment with latency reversal agents (65) can activate diverse latent HIV lineages by further suggesting that natural stimuli can do the same. This observation also supports a model where latently HIV-infected cells reactivate relatively frequently [although specific rates remain a matter of debate (66, 67)]. Our detection of plasma RNA sequences during participant 1’s treatment interruption that exactly matched HIV DNA sequences isolated from their reservoir also corroborates a previous report (68). Of note, the fact that we failed to identify a single HIV sequence whose date estimate’s 95% CI unambiguously fell within the period of complete viremia suppression also supports the notion that residual HIV replication on cART, at least as measured in blood, is absent or negligible in this individual (26, 31, 36, 69–71).

While ours is not the first study to infer reservoir temporal origins via analysis of pre-cART HIV sequence variation, previous
studies have generally inferred phylogenies solely on the basis of sequence homology (e.g., ref. 30) and have “dated” latent lineages to the period when similar sequences circulated in plasma [a recent elegant study by Brodin et al. (36) significantly advanced the field by inferring host-specific pre-cART evolutionary rates from longitudinal HIV sequences, yet latent reservoir establishment dates were still inferred via genetic similarity to pre-cART sequences]. While reasonable and appropriate, such approaches are limited by their reliance on retrospective sampling of plasma HIV RNA sequences similar to those retrieved from the reservoir, which is not always possible, particularly for individuals who are not diagnosed until later in infection. Recovering latent HIV sequence ages via calculation of a host-specific evolutionary rate overcomes this limitation, allowing the inference of HIV reservoir ages dating back to transmission (72) even if sampling does not extend that far back (e.g., Fig. 4). Our method is also fast. While unknown sequence dates can be recovered using Bayesian approaches (53), these require an additional parameter for each unknown date, significantly increasing computational requirements such that only a small number of sequences can be dated at a time. Our framework can recover all unknown sequence dates simultaneously using a desktop computer and yielded root date and rate estimates concordant with Bayesian estimates. Our framework is also robust to rooting strategy. RTT, which places the node before the root position in the relationships between the training data root-to-tip distances and sampling dates, will inherently yield better-fitting linear models than OGR. Nevertheless, OGR performed nearly as well as RTT at reconstructing unknown HIV sequence ages (Fig. 6), even on sparse training data (SI Appendix, Figs. S7 and S8). Data requirements are also relatively modest. We dated latent HIV lineages in an individual who did not achieve pVL suppression on cART until a decade after infection, using as few as two training time points (although more were preferable). Framework application to minimal training data should generally be feasible for individuals who initiate cART early, a practice that is consistent with our previous work on reconstrcuting HIV infection dates from sequence data (44, 45, 72).

Some limitations merit mention. The framework assumes that within-host HIV evolution can be adequately modeled using a strict molecular clock, which may not be ideal over prolonged timeframes (14). While we addressed variable within-host evolutionary rates in participant 2 by inferring a linear model for each treatment era, adaptation of the framework to work under alternative clock models is warranted. Our framework also assumes that latent lineages remain unchanged until sampling (that is, infection is not followed by a latent reversion to a lower evolutionary rate). As rates of within-host HIV evolution are genotype-specific, training and censored sequences must be derived from the same HIV region. While our previous work suggests that the framework should be adaptable to any HIV gene (72), only nef and env were validated here. To ensure that HIV sequences represented true within-host lineages we used single-genome amplification; however, only a limited portion of the HIV genome was analyzed and integration sites were not mapped. As such, we cannot confirm whether the studied HIV DNA sequences reside within intact proviruses, nor can we positively state that identical sequences derive from latently infected, clonally expanded cell populations. Moreover, as proviral DNA was amplified from PBMC, cellular origin of HIV lineages remains unknown. Finally, as latent HIV ages were only estimated for two participants who initiated cART late, we cannot generalize too broadly. For example, while our observation that HIV sequences sampled during long-term cART recapitulated pretherapy HIV RNA diversity was consistent with Brodin et al. (36), these sequences did not predominantly date to the period shortly before cART as reported in their study. Whether this is attributable to modest participant numbers, methodological differences [e.g., Brodin et al. (36) used next-generation sequencing and inferred reservoir ages via genetic similarity], or substantial interindividual diversity in latent HIV reservoir dynamics and composition merits further investigation.

Despite these limitations, our framework can potentially address key knowledge gaps. For example, framework application to recrudescence plasma HIV RNA sequences following spontaneous or therapeutic latency reversal could advance our understanding of the relationships between reservoir age, lineage origin, and reactivation potential. The framework could be expanded to infer reservoir tissue or cellular origin by adapting phylodragny methods (73–75) to a within-host context. It could be used to investigate the relationship between reservoir age and HIV genomic integrity (provided the viral region used for dating is intact) or genomic integration site (e.g., by using the LTR for model calibration) and should be scalable to next-generation sequence data (44). In conclusion, our framework could allow dating of viral sequences from several time points with regard to the genetic composition and longevity of the latent reservoir and represents a versatile and potentially powerful addition to the HIV persistence research toolkit.

Materials and Methods

Simulations. Birth–death models have been used to study speciation (76) and epidemics (77) and to model the proliferation of virus lineages within-host (78). We generated 1,000 maximum likelihood phylogenies with 100 tips each under a birth–death model using serial-sampled simulations in which pairs per d, the death rate = E8965 10−5 per d, and the sampling function in the R package Treesim (80). The model was parameterized by fitting it to data from McCloskey et al. (72). We used BEAST v2.3.2 (81) to estimate the posterior distribution defined by the serial model with a birth-death skyline tree prior (77), a strict molecular clock, and an HKY85 model of nucleotide substitution (82). The lineage birth rate was estimated as 5.12 × 10−4 per d, the death rate was 5.01 × 10−5 per d, and the sampling probability as 5.24 × 10−4. We used NELSI (83) to introduce temporal variation in individual branch lengths by assigning rates of evolution to these trees by drawing values from a Gaussian distribution with a mean μ = 1.96 × 10−4 substitutions per generation and SD = 1.42 × 10−5. To model uncertainty in phylogenetic reconstruction we simulated sequence evolution along each tree with INDELible v1.03 (84) under an HKY85 substitution model where parameters were drawn from distributions derived from published within-host HIV datasets (stationary distributions 0.42, 0.15, 0.15, and 0.28 for Α, Г, C, and T, respectively; transition bias parameter 8.5) (72).

Published Datasets. Two datasets from the LANL were assembled for model evaluation (SI Appendix, Table S1) (58). The first comprised 424 longitudinal plasma HIV RNA partial env sequences from seven individuals from ref. 72 and one from ref. 62, selected based on three predefined criteria: individuals were treatment-naive and had two or more clonal or single-genome sequences per time point where the baseline sample was within 186 d of infection or seroconversion and where at least one subsequent time point was ≥6 mo later. We also required that the overall within-host phylogeny display molecular clock signal (discussed below). The second dataset comprised 361 plasma HIV RNA-derived and 545 PBMC-derived partial env sequences that were first or second sampled after a significant latency time (determined by a lack of viral RNA in plasma, to whom at least three RNA time points were available, where the earliest HIV RNA time point was at or before the earliest HIV DNA time point (SI Appendix, Table S1). Infection date did not need to be known and the dataset comprised both treatment-naive (14, 85) and treated (86) persons.

Participant Recruitment and Sampling. The framework was applied to two participants recruited from the British Columbia Centre for Excellence in HIV/AIDS for whom longitudinal archived plasma (and in one case, a single archived whole-blood specimen), dating back to 1996, were available. Participants provided blood in summer 2016 from which PBMC were isolated by standard density gradient separation (Histopaque-1077; Sigma), counted (TC20 automated cell counter; Bio-Rad), and cryopreserved at −150 °C in 90% FBS:10% DMSO. This study was approved by the Providence Health Care/University of British Columbia and Simon Fraser University research ethics boards (harmonization protocol H15-03077) and both participants gave written informed consent.

Single-Genome HIV Sequencing. HIV RNA was extracted from 0.5 μL plasma using the NucliSENS EasyMag system (BioMerieux) and genomic DNA was extracted from 200 μL whole blood or 5 μL PBMC using the Invitrogen genomic DNA isolation kit. Nef was selected for framework application due to its high within-host diversity and richness of phylogenetic signal. Single-genome amplification was performed by limiting dilution using high-fidelity enzymes and HIV sequence-specific primers optimized for amplification of multiple HIV-1 group M subtypes. For HIV RNA extracts, cDNA was
Verification of Root Date Estimates Using Bayesian Methods. Linear model--
derived evolutionary rate and root date estimates were validated using BEAST v1.8.4 (95). Posterior distributions for the root date were estimated using a SRD06 substitution model (96), a lognormal uncorrelated relaxed molecular clock (97), and a coalescent Gaussian Markov random field Bayesian skyrider tree prior (98) in four parallel Markov chain Monte Carlo chains with 10^6 iterations each. After discarding the first 10% as burn-in, chains were combined with LogCombiner v2.4.2 (81) and analyzed in Tracer v1.6 to ensure convergence and effective sample size values >200 for all parameters.

Comparison of nef to the Rest of the Genome. We analyzed a published
longitudinal within-host near-whole genome HIV sequence dataset from an
untreated individual (GenBank accession no. DQ853436-65) (62), removed
14 sequences that exhibited evidence for recombination (88), and aligned
the remaining 16 to HXB2 using MAFFT v7.313 (99). We used Pearson's
correlation to evaluate the relationship between the number of nucleotide
differences between each sequence and the consensus of the earliest time
point within nef vs. the rest of the HIV genome (ignoring gaps). We also
used the partition-homogeneity test implemented in PAUP* v4.0a (100, 101)
with 1,000 replicates to compare evolution in nef vs. env.

Statistical Analyses. Statistical calculations were performed using R v3.3.3. Cs
were calculated using the inverse.predict function of the cemical package in
R. Distributions asymmetry in reconstructed dates was tested using a two-
tailed nonparametric binomial test followed by a Bonferroni correction
(57). Plots were generated using ggplot2 and ggtree (102) packages in R.
Divergence times for ancestor traces were estimated using the estimate
dates function of the ape package in R (103, 104).

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