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Reliable reconstruction of HIV-1 whole genome haplotypes reveals clonal interference and genetic hitchhiking among immune escape variants

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Abstract

Background: Following transmission, HIV-1 evolves into a diverse population, and next generation sequencing enables us to detect variants occurring at low frequencies. Studying viral evolution at the level of whole genomes was hitherto not possible because next generation sequencing delivers relatively short reads.

Results: We here provide a proof of principle that whole HIV-1 genomes can be reliably reconstructed from short reads, and use this to study the selection of immune escape mutations at the level of whole genome haplotypes. Using realistically simulated HIV-1 populations, we demonstrate that reconstruction of complete genome haplotypes is feasible with high fidelity. We do not reconstruct all genetically distinct genomes, but each reconstructed haplotype represents one or more of the quasispecies in the HIV-1 population. We then reconstruct 30 whole genome haplotypes from published short sequence reads sampled longitudinally from a single HIV-1 infected patient. We confirm the reliability of the reconstruction by validating our predicted haplotype genes with single genome amplification sequences, and by comparing haplotype frequencies with observed epitope escape frequencies.

Conclusions: Phylogenetic analysis shows that the HIV-1 population undergoes selection driven evolution, with successive replacement of the viral population by novel dominant strains. We demonstrate that immune escape mutants evolve in a dependent manner with various mutations hitchhiking along with others. As a consequence of this *clonal interference*, selection coefficients have to be estimated for complete haplotypes and not for individual immune escapes.

Keywords: Haplotype reconstruction, HIV-1 genome, Next generation sequencing, Clonal interference, Genetic hitchhiking

Background

Inside every human host, HIV-1 embarks upon an arms race to evade the host's immune responses [1,2]. A single (or a few) founder genetic lineage(s) establish the infection in the CD4+ cells localized in the mucosa at the port of entry [1,3-7]. HIV-1 then spreads to regional lymph nodes where it comes in contact with a large number of CD4+ T cells. Abundance of CD4+ T cells allows rampant viral replication resulting in millions of

viral particles per ml of plasma at peak viremia, occurring ~21–28 days post infection [3,8]. Interplay between HIV-1, target cell availability, and the host's immune system during the acute phase of infection leads to a gradual decline of the viral load, eventually leveling off to a "set point" [2,5,9]. The decrease in the viral load following peak viremia is attributed to CD4+ T cell depletion, the immune responses of HIV-1 specific B cells, and cytotoxic T cells (CTLs). A high mutation rate and a large population size allow HIV-1 to rapidly evolve immune escape mutants [10,11], and strong CTL selection pressures confer fitness advantage to viral lineages that harbor CTL escape mutations [3,12,13]. The efficacy and breadth of

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cognate CTL responses increases the within-host HIV-1 diversity after peak viremia, and plays a major role in HIV-1 evolution during the acute phase of infection [5,9,14-19].

Longitudinally collected samples have been used to study the evolution of escape mutations, which can be used to estimate the selection pressure imposed by the corresponding CTL response [11,20,21]. The rate of escape is typically estimated using a logistic curve to model the replacement of wildtype epitope by escape variant [22]. Several escape mutations of the same epitope may occur simultaneously in different viral lineages, but such multiple escape pathways for a single epitope are generally modeled as a single escape event [12,17,18,23,24]. Escape mutations typically result in a fitness cost to the virus, and along different escape pathways the virus may face a different selection pressure by the CTLs [1,20,21,23,25]. Therefore, it was recommended to consider different escape pathways within the same epitope as separate escape events [23].

Escape mutations acquired in different epitopes are typically considered independent events. But in reality multiple epitopes may escape either in the same lineage, or in different lineages of the same viral population [1,11]. Viral lineages containing multiple escape mutations are expected to cause a selective sweep, reducing viral diversity and driving other lineages to extinction. The concurrent presence of different escape mutations in different viral lineages may result in competition between them in their race towards fixation [23,26,27] via *clonal interference* [28,29]. Due to clonal interference, two beneficial mutations residing in two different lineages out-compete each other as well as the wild-type when one approaches fixation. Additional beneficial mutations can be sequentially acquired, ultimately resulting in escape mutations in multiple epitopes [17,21,23,27]. As a result of clonal interference, epitope escape mutations may also drive several non-epitope mutations, co-occurring in the same viral lineage, to fixation via *genetic hitchhiking* [26,30,31]. Clonal interference has recently been shown to play an important role in the evolution of influenza [26,27]. One would expect it to also influence the evolution of HIV-1, but as yet there is no evidence for this.

To study the role of clonal interference and genetic hitchhiking in HIV-1 evolution, one requires long, or ideally complete, genome sequences with high coverage from multiple time points. The availability of longitudinally sampled single genome sequences is rather limited, and next generation sequencing (NGS) is economically a more feasible alternative for single genome sequencing [1,7,32]. One large NGS dataset was generated by Henn et al. [11], when they performed NGS on complete HIV-1 genomes from 6 longitudinal samples in one patient (subject 9213). NGS allowed Henn et al. [11] to study

the dynamics of immune escape mutations in individual epitopes. However, due to the short sequencing read lengths of about ~ 400 bp, Henn et al. [11] were not able to distinguish whether multiple epitopes escape simultaneously in the same haplotype, or independently in distinct viral haplotypes. We here attempt to reconstruct whole genome haplotypes from this data. This is challenging due to two reasons [33,34]. First, a variety of sequencing and PCR errors are incorporated in sequencing reads, which require robust error correction techniques to identify the true mutations [33,35,36]. A second problem concerns the determination of contiguity of sequencing reads generated from a single viral genome haplotype. Differences in prevalence of the genome haplotypes, genetic distances between polymorphic sites, and the amount of sequencing errors, are important factors influencing the reliability of haplotype reconstruction. A number of algorithms have been developed for viral haplotype reconstruction [33,34,37,38], and among them PredictHaplo has been shown to perform at high precision in reconstructing gene haplotypes at a high sequence divergence [34,39].

We here demonstrate that reconstruction of whole genome HIV-1 haplotypes from NGS datasets is feasible using a read clean-up step before haplotype prediction with PredictHaplo. The reconstructed genome haplotypes provide important insights into the evolutionary dynamics of the viral quasispecies, as we show that the epitope escape dynamics are influenced by the strength and breadth of CTL selection, clonal interference, and genetic drift.

Results and discussion

Testing the feasibility of haplotype reconstruction using simulated datasets

Before reconstruction of HIV-1 whole genome haplotypes from next generation sequencing reads, we first tested the feasibility of the haplotype reconstruction pipeline using *in silico* datasets. We generated six *in silico* HIV-1 populations with nucleotide diversities (1, 2, 4 and 10%) [11,40,41] and frequency distributions (uniform or log-normal, see Methods). For each dataset, ART_454 software [42] was used to generate *in silico* next-generation sequencing reads. Each simulated population consisted of 9 master HIV-1 genomes that were 8800 bp long. For the 4 data sets with uniform frequency distributions (data sets U1, U2, U4 and U10, respectively), our pipeline reconstructed 9 haplotypes from the short reads artificially generated from the true 9 *in silico* genomes (see Methods). The reconstruction was accurate with a Hamming distance of just 1 to 9 nucleotides from the true genomes (Table 1).

The accuracy decreased when the population structure was changed to a log-normal frequency distribution (data sets L1, L2, L4 and L10, respectively). For instance,

Table 1 Haplotype reconstruction performance on simulated datasets

Dataset	Nucleotide diversity	No. of haplotypes reconstructed	Mean Hamming distance (MHD)	MHD of top 4 haplotypes	Range Hamming distance
U1	1	9	6	6	(6-6)
U2	2	9	4.33	2.25	(1-9)
U4	4	9	4.33	2	(2-9)
U10	10	9	4.66	2.25	(1-9)
L1	1	6	49.5	28.25	(11-93)
L2	2	5	46.2	23	(7-139)
L4	4	9	158.44	11	(1-427)
L10	10	7	283.85	65.25	(46-706)
LHV	1+	9	20.66	13.5	(13-37)

Simulated datasets were named according to the frequency distribution of their population and their nucleotide diversity. Populations with uniform and log-normal frequency distributions are labeled as U and L, respectively, followed by a number denoting the percentage nucleotide diversity between genomes. In all cases there were 9 “true” genomes.

for the L4 dataset with 4% nucleotide diversity, the 9 reconstructed haplotypes had a Hamming distance of 1 to 427 nucleotides from the corresponding true genomes (Table 1). Closer examination, however, revealed that the four most prevalent (>4%) haplotypes were accurately reconstructed, with Hamming distances of 1, 1, 19 and 23. Additionally, most of the errors were limited to either the start or end of the genomes, where the simulated coverage was low. Ignoring the initial and final 20 bases of the haplotype sequences, the four most prevalent haplotypes were identical to their closest true genomes.

Some regions of HIV-1 genome are known to have higher mutation rate than others [43]. To model the variability in mutation rate in HIV-1, we simulated a viral population (LHV) with genomes containing 3 hypervariable regions (see Methods). The 9 reconstructed genomes from the LHV data set had a Hamming distance of 13 to 37 nucleotides from the corresponding true genomes (Table 1), whereas the four most prevalent haplotypes were reconstructed at a Hamming distance of 13, 13, 13 and 15. For all data sets, the predicted frequencies of the reconstructed haplotypes matched those of the true genomes (results not shown).

In reality, HIV-1 is known to have a complex population structure involving multiple co-occurring quasispecies [44]. To mimic this, we simulated a more complex population (LQ4_i) consisting of 9 master HIV-1 genomes

corresponding to 9 quasispecies, each surrounded by a varying number of closely related genomes (Table 2 and Figure 1A; see Methods). This was simulated a 100 times, and for each simulation a minimum of 5 and a maximum of 8 haplotypes were reconstructed (Table 2). The 4 most abundant haplotypes represented the 4 most prevalent quasispecies in the population, and these were more accurately reconstructed than the low frequency haplotypes. Most sequence errors were found in the initial and final 50 bp of the reconstructed haplotypes, exclusion of which resulted in a large reduction in the Hamming distance between the reconstructed haplotypes and the true genomes, especially for the prevalent haplotypes (Table 2).

We now present a detailed analysis of LQ4_1, as a representative example (the 9 other simulations were similar). The five most prevalent haplotypes (haplo_0, haplo_3, haplo_4, haplo_5 and haplo_6; see Figure 1) were reconstructed at a Hamming distance of 8 to 46 nucleotides from their closest sequences (Table 2), and were reconstructed as a “consensus genome” of their corresponding quasispecies (Additional file 1: Figure S1). However, the most abundant quasispecies containing 50 simulated genomes, was reconstructed as two haplotypes, i.e. haplo_0 and haplo_4, each “representing” 22 and 28 simulated genomes (Figure 1A). The remaining two haplotypes (haplo_1 and haplo_2) were reconstructed as consensus genomes of the five rare (frequencies <4%)

Table 2 Haplotype reconstruction of quasispecies datasets

Dataset	No. of haplotypes reconstructed			Mean Hamming distance (MHD)	MHD of top 4 haplotypes	MHD of top 4: pruned
	Mode	Min	Max			
LQ4_1 to LQ4_100	6	5	8	75.76	13.63	4.94

Haplotype reconstructions from populations having a simulated viral quasispecies structure. Nine quasispecies were present in each simulated population and the diversity between them was 4% (see Methods). The pruned haplotype genomes were created by excluding the initial and final 50 bp from the reconstructed haplotypes to avoid errors due to low read coverage. A 100 simulations were performed.

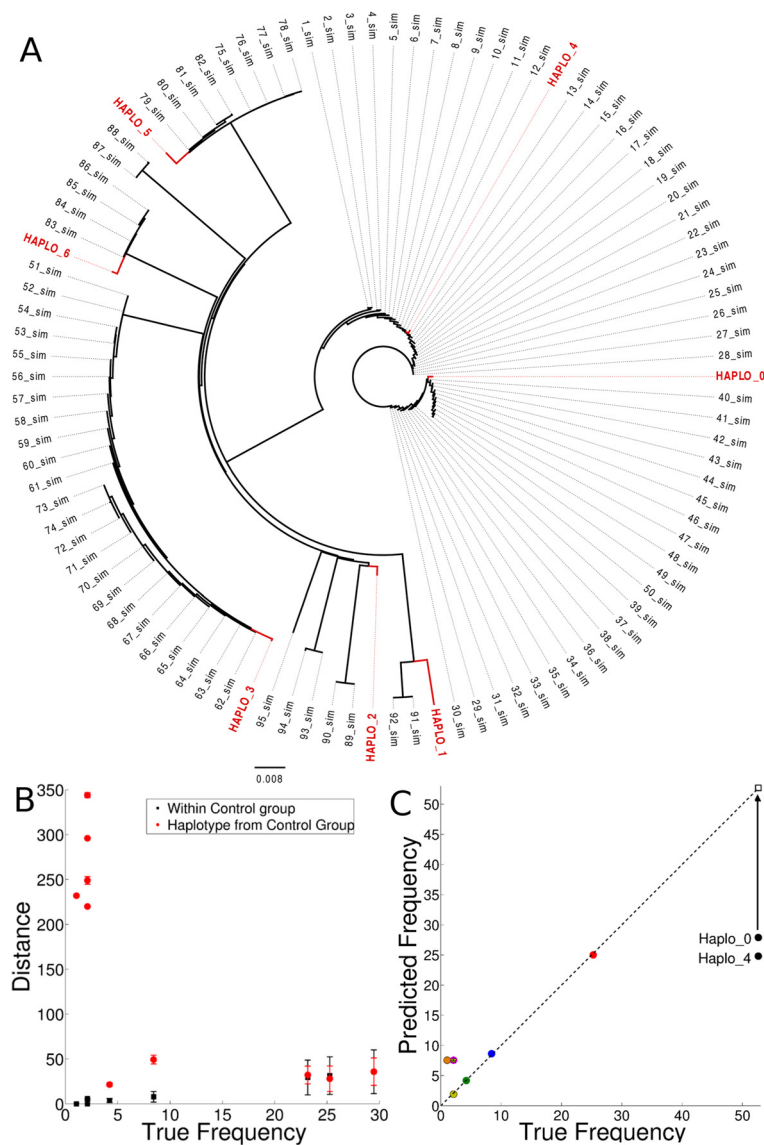


Figure 1 Accuracy of reconstructed haplotypes from simulated HIV-1 quasiespecies (LQ4_1). We simulated HIV-1 population (LQ4_1; Table 2) with 9 quasiespecies consisting of 50, 24, 8, 4, 2, 2, 2, 2 and 1, sequences respectively. The genome sequences within each quasiespecies contained at most 0.1% nucleotide diversity (see Methods). **(A)** Seven reconstructed whole genome haplotypes (red) were aligned with the 95 simulated genomes. Dark lines represent the branches of the phylogenetic tree (the dotted lines pointing outwards provide better visualization). **(B)** Average Hamming distance within each quasiespecies (black), and between the reconstructed haplotype and the quasiespecies at minimum Hamming distance (red). Because one haplotype can represent multiple quasiespecies, we used the phylogenetic tree to determine the genomes represented by a haplotype. Markers represent the mean Hamming distance and error bars represent 1 standard deviation. **(C)** The haplotype frequencies (vertical axis) predicted from haplotype reconstruction pipeline are plotted against the true quasiespecies frequencies (horizontal axis). The dominant quasiespecies was reconstructed as two haplotypes (Haplo_0 and Haplo_4) and summation of their predicted frequencies is indicated by an open square. The haplotype Haplo_2 (predicted frequency of 7.55%) represents four low prevalence quasiespecies: one quasiespecies with true frequency of 1.05% (orange circle) and three with true frequencies of 2.1% (see the overlapping points next to orange circle). The points lying on the dashed diagonal line represent perfect haplotype frequency predictions.

quasiespecies (Figure 1A). Variation in the Hamming distance between the reconstructed haplotype and genomes of the corresponding quasiespecies was comparable to the intrinsic variation within the quasiespecies (Figure 1B). The dominant quasiespecies were accurately reconstructed, but

the less frequent quasiespecies were reconstructed at a much higher Hamming distance (Figure 1B). Since haplotypes are reconstructed as a consensus of each quasiespecies (Additional file 1: Figure S1), variations between the reconstructed haplotypes and their true genome can

at least partly be attributed to the viral quasispecies population structure. The predicted prevalence of the dominant haplotypes was comparable to the true frequencies of the corresponding quasispecies (Figure 1C). Summarizing, these simulations suggest that the haplotype reconstruction pipeline works with high fidelity for complete HIV-1 genome haplotype reconstruction using 454 sequencing reads with realistic nucleotide diversities of 1 to 10%, and genotype frequencies larger than 4%.

We have used PredictHaplo to reconstruct genome haplotypes. Several other algorithms are available for haplotype reconstruction [34,37-39]. To test the performance of these algorithms we also reconstructed haplotypes using two other algorithms, QuRe and ShoRAH. For the *in silico* HIV-1 data sets (given in Tables 1 and 2), each containing 9 true genomes, ShoRAH reconstructed ≥ 187 haplotypes with a ≥ 117.80 mean Hamming distance for each data set (Table 3). For the same *in silico* HIV-1 data sets (Tables 1 and 2), QuRe reconstructed ≥ 21 haplotypes with a ≥ 211.18 mean Hamming distance for each data set (Table 3). Corroborating a previous study by Schirmer et al. [34], we see that ShoRAH and QuRe overestimate the number of haplotypes and have lower precision to predict “true” haplotypes (Table 4). PredictHaplo reconstructs haplotypes more reliably than both ShoRAH and QuRe even if we only consider the four most prevalent haplotypes (Tables 1, 2, and 3). Due to the large number of spurious haplotypes predicted by ShoRAH and QuRe, the predicted haplotype frequencies do not match with the true frequencies (results not shown), whereas the haplotype frequencies predicted by PredictHaplo were accurate (as shown in Figure 1C). Note that previous studies have shown that both QuRe and PredictHaplo reconstruct shorter haplotypes (length ~ 1400 bp) at high precision [39]. Since the aim of our study was not to benchmark algorithms but to study the evolutionary dynamics of immune escape mutations. We continue with PredictHaplo as it produced the lowest number of

Table 4 Precision of haplotype reconstruction algorithms

Algorithm	U2	U4	U10	L2	L4	L10	LQ4_1
PredictHaplo [53]	1	1	1	0.60	0.44	0	0.57
ShoRAH [38]	0.10	0.18	0.39	0	0	0	0
QuRe [37]	0	0	0.01	0	0	NA	0

We calculated the precision by taking the ratio between the number of accurately reconstructed haplotypes over the total number of reconstructed haplotypes. An haplotype was considered to be accurately reconstructed if the Hamming distance to the “true” genome was ≤ 25 . Recall values are not listed as most of them are 0 for ShoRAH and QuRe.

spurious whole genome haplotypes, and accurately predicted the haplotype frequencies. The other algorithms may prove more apt to address different questions.

Experimental validation of the reconstructed haplotypes

The same methodology is now applied to the NGS data from a patient described in Henn et al. [11]. In brief, this patient presented during acute infection with a viral load of 9.3×10^6 copies/ml, and henceforth this is referred to as day 0. Six longitudinal serum samples were collected at day 0, 3, 59, 165, 476 and 1543 post presentation by Henn et al. [11]. The peak viral load was observed at day 3. The first four samples represent the acute phase dynamics, while the last two samples fall in the chronic phase of infection. Deep sequencing (Roche 454 Genome Sequencer FLX Titanium) was performed on each sample using four overlapping PCR amplicons spanning the complete protein coding HIV-1 genome. The average fold sequence coverage for the samples from day 0, 3, 59, 165, 476 and 1543 was 667.7, 724.4, 750.5, 299.7, 227.6 and 540.7, respectively. CTL epitope escapes restricted by subject’s HLA alleles (A01, A24, B38, B44 and Cw04) were studied using a local read analysis, and confirmed using IFN-gamma ELISPOT assays. The most dominant CTL responses at day 59 were directed

Table 3 Comparison of haplotype reconstruction algorithms

Dataset	ShoRAH			QuRe		
	No. of haplotypes reconstructed	MHD	MHD: top 4	No. of haplotypes reconstructed	MHD	MHD: top 4
U2	187	117.80	0	104	228.66	168.50
U4	201	201.49	0	146	416.38	230.50
U10	200	315.515	1	86	607.10	83.75
L2	210	217.47	186	21	260.24	245.25
L4	205	577.97	336.25	48	565.29	510.75
L10	344	1182.13	886.25	NA	NA	NA
LQ4_1	200	220.485	60.75	27	211.18	103

Simulated viral populations given in Tables 1 and 2 were also analyzed by two alternative haplotype reconstruction algorithms, i.e., ShoRAH and QuRe [33,37,38]. All datasets contained 9 “true” genomes. NA indicates that QuRe was terminated after two weeks of CPU run time (3.1 GHz 16 core Intel Xeon processor, 128GB RAM).

against the Nef-RW8 and Vif-W19 epitopes. To validate the sequencing reads, Single Genome Amplification (SGA) was performed for *vif* gene sequences from day 59. The public availability of this data allowed us to test the validity of our whole genome haplotype reconstruction pipeline.

Validation of reconstructed haplotypes by SGA

As the first validation step, the 95 SGA *vif* sequences from day 59 [11] were aligned with the 6 *vif* gene sequences extracted from the whole genome haplotypes reconstructed using the NGS data from day 59. Phylogenetic analysis showed that five reconstructed *vif* sequences were identical to 48 out of 95 SGA *vif* sequences (Figure 2A). Of the remaining SGA sequences, 11 were found in clades that were represented by at least one reconstructed haplotype (e.g., B7, B42, B57, B81 and B130 were represented by d59_1). 16 low frequency SGA variants, depicted as singletons in Figure 2A, were represented by one haplotype (d59_2). Thus, 75 of the 95 observed SGA sequences were represented by 6 reconstructed genome haplotypes. The remaining 20 SGA

sequences were prevalent at low frequencies, and were not reconstructed as a unique haplotype. Even though we reconstruct whole genome haplotypes, we apparently detect haplotypes with mutations in just a few bases within a region of 600 bp (spanning at least 2 reads), confirming that global haplotypes accurately capture local features.

Temporal variations in haplotype prevalence frequencies

Next to reconstructing the sequences, we should also be able to predict the correct genotype frequencies of the haplotypes. Due to transmission bottlenecks, HIV-1 infection is typically established by a single (or a few) genetic lineage(s) [1,4-6,23], and the HIV-1 population does not exhibit much diversity until the peak viremia [11]. One should therefore expect one (or a few) dominant HIV-1 haplotype(s) before and around the time viremia peaks. Around the peak viremia, immune selection leads to diversification, i.e. a decrease in the frequency of the dominant quasiespecies, and an increase in frequency of other quasiespecies in the population [1,23]. Reconstructing the haplotypes from every sample, we found that the

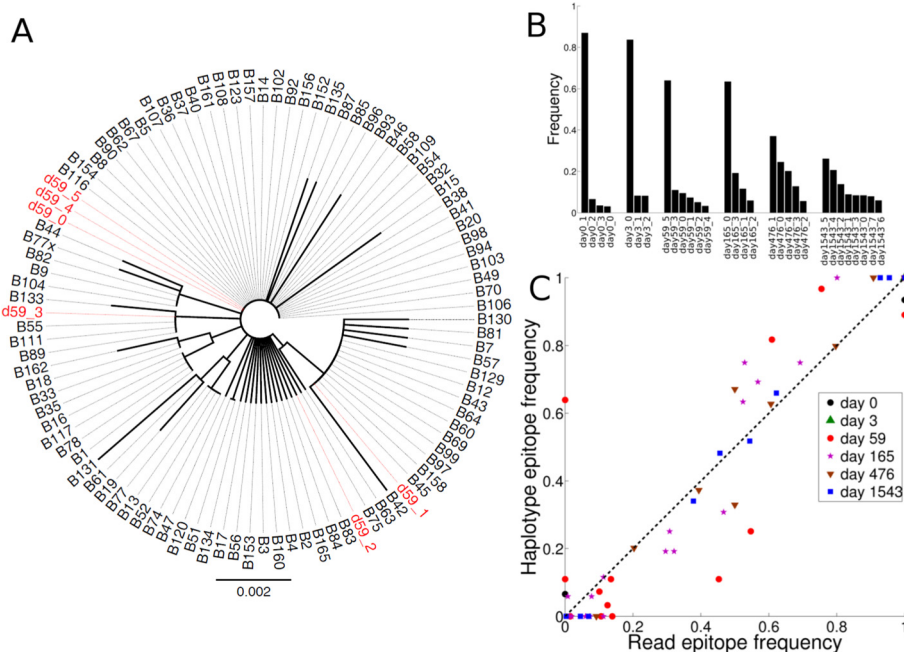


Figure 2 Validation of reconstructed haplotypes from subject 9213. (A) Neighbor-joining phylogenetic tree generated from the global alignment of the *vif* genes extracted from the 6 haplotypes reconstructed from the day 59 data (red) and the SGA *vif* gene sequences (black) obtained from [11]. **(B)** The predicted haplotype frequencies for subject 9213 plotted for each sample (day 0, day 3, day 59, day 165, day 476, and day 1543). Note that the variation in read coverage (667.7, 724.4, 750.5, 299.7, 227.6 and 540.7, respectively) does not seem to influence the number of predicted haplotypes. **(C)** Comparison of the epitope frequencies estimated from reconstructed haplotypes and those from the local analysis of sequencing reads described in [11]. Markers represent different days. Variants of each epitope are represented as different observations. Several epitope variants with 100% read frequency and a 100% predicted frequency in reconstructed haplotypes appear as overlapping points in the upper right corner on the diagonal. The overlapping points were not considered for the correlation analysis. The Spearman's rank correlation coefficient between predicted epitope frequencies extracted from complete genome haplotypes and the local read frequencies was $\rho = 0.85$ (p -value $< 10^{-15}$).

number of reconstructed haplotypes varied from 3 to 8 (Figure 2B). Although the viral population at day 0 (i.e., before peak viremia) consisted of 4 haplotypes, only one haplotype (d0_1) was present at a high frequency (~87%), confirming that this infection was established by a single founder virus lineage. At day 3 (around peak viremia), the same dominant haplotype (d3_0, identical to d0_1) was present at a similar high frequency (~83.6%). Diversification led to a decrease in the frequency of dominant haplotypes at day 59 and 165. Several haplotypes with comparable frequencies were reconstructed for day 476 and 1543, indicating diversification and expansion of multiple lineages during the early chronic phase of infection (Figure 2B). The reconstructed haplotypes thus suggest a realistic scenario where the infection in subject 9213 was started by a single haplotype, followed by an increase in viral diversity resulting in the establishment of multiple co-dominant viral quasispecies.

Validation of whole genome haplotypes using predicted epitope frequencies

As a third validation, we compared the frequencies of the CTL epitopes, obtained by summation of the predicted frequencies of those haplotypes containing the CTL epitope, with the observed read frequencies described in Henn et al. [11]. The average nucleotide diversity between whole genome haplotypes from the same sample was 128 nucleotides. As the epitope escape mutations comprise only a small fraction of all the polymorphic sites used to infer the whole genome haplotypes, this comparison is a fairly independent validation. Since it is trivial that the epitopes containing no variation in a given sample will always exhibit 100% read and haplotype frequencies, they were not considered for the correlation analysis. The predicted frequencies of the epitopes were accurate and tightly correlated with the observed frequencies (Spearman's $\rho = 0.85$, $p\text{-value} < 10^{-15}$) (Figure 2C). Some of the low frequency epitope variants were not predicted by haplotype reconstruction, and they could either be sequencing errors, or true variants not containing sufficient contiguous reads to allow their reconstruction. The Nef RW8-T5M, the Gag A01-R6K and the Gag A01-E3A escape mutations, which were not detected by the local read analysis of Henn et al. [11], were predicted at $\geq 6.5\%$ frequency on the day 0, and 59 haplotypes (red, and black circles on vertical axis, Figure 2C). Henn et al. [11] performed several additional clean-up steps (read phasing, read profiling, and removal of reads partially spanning an epitope), which apparently discarded the reads containing these Nef and Gag variants. Because the Nef RW8-T5M and the Gag A01-R6K mutations were present at subsequent time points [11], we think their detection by the pipeline is correct. To summarize, we can correctly predict the epitope frequencies by reconstructing whole genome haplotypes.

Biological results

Phylogenetic analysis of reconstructed haplotypes

A hallmark of HIV-1 evolution during the acute phase of infection is the selection by the immune system [1,45]. The successive replacement of viral haplotypes in the phylogenetic trees in Figure 3 reveals that HIV-1 in subject 9213 also underwent a selection driven evolution. The *nef*, and *vif* genes, which are targeted by dominant CTL responses [11], and the *env* gene targeted by 3 subdominant CTL responses [11], exhibit a strong temporal phylogenetic signal of successive replacement of dominant haplotypes (Figure 3A-C). Strong temporal selection causes all day 1543 haplotypes to form a monophyletic clade diverging from the clade containing the dominant haplotype at day 476 (d476_1) (Figure 3A-C). Using codon selection analysis we found that 11 codons in *nef*, 4 codons in *vif* and 50 codons in *env* were under positive selection (see Methods). The *gag*, and *pol* genes targeted by subdominant CTL responses exhibit intermediate phylogenetic signals for temporal selection (Figure 3D and E). The trees of the remaining genes (*rev*, *tat*, *vpr*, and *vpu*) did not reveal immune selection driven evolution (Figure 3F-I). Codon selection analysis showed that no codon was under positive selection in *tat*, *vpr*, and *vpu*; whereas 1 codon in *rev* was under positive selection (see Methods). The complete genome phylogeny captured the temporal dynamics exhibited by the genes under identified CTL selection, and showed similar successive replacement of dominant haplotypes (Figure 4). Thus, phylogenetic analysis indicates temporal replacement of the dominant quasispecies, and that this signal is most evident in the genes targeted by the CTLs.

Clonal interference and epitope escape

Studying the 7 identified CTL epitopes in the whole genome haplotypes reveals that the virus explores multiple escape pathways in most epitopes. Different combinations of escape mutations were found in different haplotypes from the same population (Figure 5). The presence of different combinations of escape mutations in each haplotype should confer different fitness advantages to them resulting in clonal interference between the viral haplotypes. This illustrates that the selective advantage of escape mutations has to be considered in combination. The population at day 59 contained 6 escape mutations distributed over 4 different epitopes, 2 in Nef A24-RW8, 2 in Vif B38-WI9, 1 in Gag A01-GY9 and 1 in Env A01-RY9 (Figure 5). Interestingly, haplotype d59_3 containing 2 of the CTL escape mutations disappeared before day 165, while d59_5, which was the dominant haplotype at day 59 with only one escape mutation survived, and additionally evolved an anchor position escape mutation in the Vif B38-WI9 epitope by day 165. Interestingly, haplotype d165_3 contained 4 epitope escapes and was not

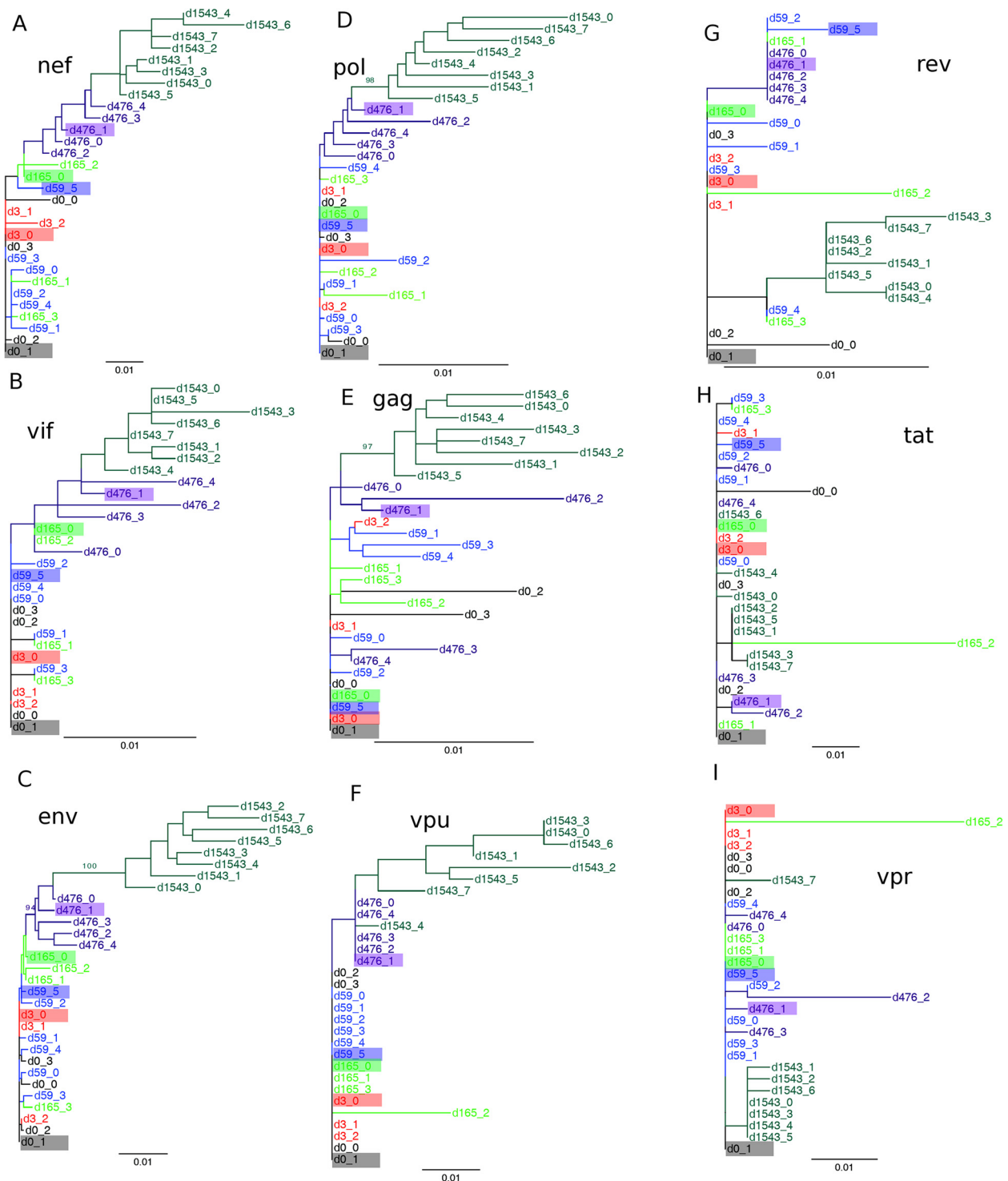
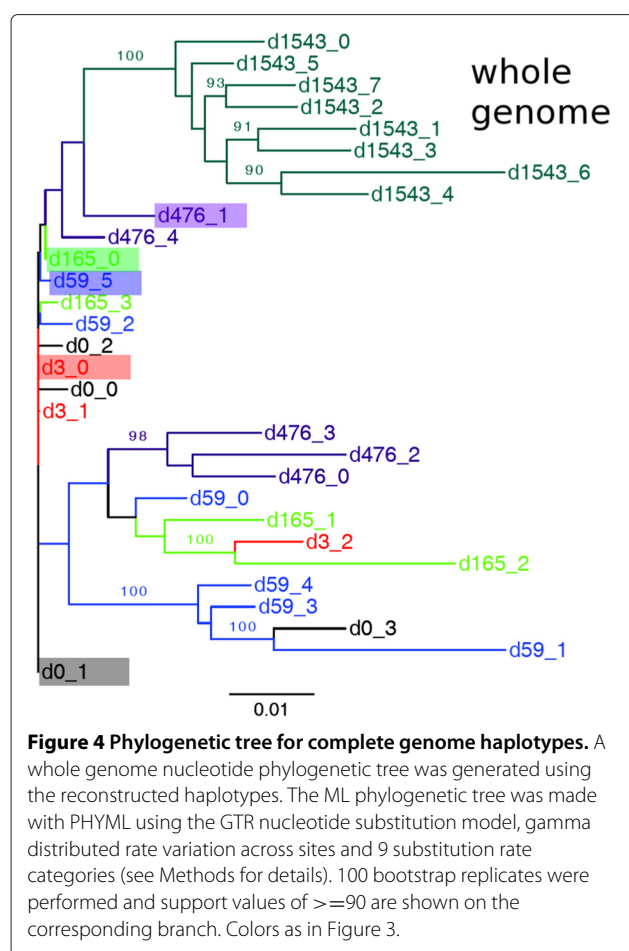


Figure 3 Phylogenetic trees of the reconstructed HIV-1 genes extracted from genome haplotypes. The phylogenetic trees generated from the nucleotide sequences of the (A) *nef*, (B) *vif*, (C) *env*, (D) *pol*, and (E) *gag* genes, that are targeted by one or more CTLs [11] exhibit a strong to intermediate signal for temporal selection. The *vpu* (F), *rev* (G), *tat* (H), and *vpr* (I) genes exhibit weak or no signal for temporal selection in their phylogenetic trees. The maximum likelihood (ML) phylogenetic trees were reconstructed with PHYML using the GTR nucleotide substitution model, gamma distributed rate variation across sites and 6 substitution rate categories (see Methods for details). 100 bootstrap replicates were performed and support values of ≥ 90 are shown on the corresponding branch. Colors denote different temporal haplotypes: day 0 (grey), day 3 (red), day 59 (blue), day 165 (green), day 476 (purple) and day 1543 (teal). The dominant haplotype for each sample is highlighted by a colored box.



selected over time, whereas haplotype d165_0 containing two escape mutations was selected (Figure 5).

An epitope typically contains two anchor positions that are essential for binding the MHC molecule. An epitope with an anchor position escape therefore results in the evasion from all potential CTL responses, as the epitope is no longer presented by the MHC molecule. This should confer a higher fitness compared to non-anchor position escapes which can be targeted by other CTL populations. This could explain why haplotype d165_0 with an anchor position escape (I9V) for the dominant Vif epitope was selected over haplotype d165_3 with the non-anchor S8A escape (Figure 5). Additionally, the strength of CTL selection may influence the fate of other epitope escape mutations. The fixation of haplotype d165_0, containing the escape mutations Vif B38-I9V and Nef A24-T5M from two dominant CTL responses, could therefore explain why the Gag A01-R6K and Env A01-I6V escape mutations in the d165_2 haplotype disappeared from the population (Figure 5). Thus, several factors like the strength of each CTL response, breadth of CTL responses, and genetic drift, together appear to determine the relative

fitness of a haplotype. Most importantly, beneficial epitope escape mutations disappear due to higher relative fitness of other haplotypes present in the viral population, emphasizing that clonal interference plays an important role in acquisition of epitope escape mutations by HIV-1.

Selection of haplotype genomes

Figure 6 depicts the temporal dynamics of the reconstructed HIV-1 haplotype genomes. The dominant (black circle) and other less abundant (gray squares) haplotypes at each time point were connected to the preceding haplotype with the minimal non-synonymous Hamming distance (mentioned above each line in Figure 6). The haplotypes were thus connected with the preceding haplotypes based on their sequence similarity. In all six samples, the dominant quasiespecies seeded the subsequent dominant quasiespecies that was generally located at a lower mutational distance, whereas the other less prevalent quasiespecies were located at a higher mutational distance from the ancestral haplotype (Figure 6). The sequence of the dominant haplotype did not change between day 0 and day 3, but differed from its predecessor at all subsequent time points (Figure 6). The rate at which the dominant haplotype is replaced by the subsequent haplotype (denoted by red dashed lines in Figure 6) decreased over time from 0.17 to 0.005 per day (Figure 6, inset). The rates of replacement calculated using whole genome haplotypes better represent the evolution of HIV-1 under immune selection than the set of escape rates estimated on the basis of individual epitopes [11].

Genetic hitchhiking

As a consequence of clonal interference, the fixation of genomes with high relative fitness due to the presence of beneficial epitope escape mutations, may result in genetic hitchhiking of other mutations present in the same haplotype [26,30,31]. The linkage disequilibrium between the nucleotide positions in HIV-1 are shown as a network of sites that are genetically linked in the set of all 30 haplotype genome sequences (Figure 7). Different colors represent different HIV-1 genes and triangles denote the mutations in epitopes (Figure 7). There are two major clusters of linked sites, one containing sites from the *gag* gene only (cluster I), and another containing sites from several genes (cluster II). In addition there are several clusters containing less than 5 sites. Epitope escapes are typically linked with one or more sites in non-epitope regions and these links could reflect compensatory mutations (Figure 7). Interestingly, the two early epitope escapes from the dominant CTL responses against Nef A24-RW8 (T5M genomic site: 8495, green arrow) and Vif B38-WI9 (genomic site: 4544 and 4547, blue and red arrow) were linked only with very few other sites (Figure 7). In contrast, the late epitope escapes from sub-dominant CTL

		A24-RW8	B38-WI9	A01-GY9	A01-RY9	B44-EW9	Cw4-SF9	A24-RL9
	Freq	Nef	Vif	Gag	Env	Pol	Env	Env
d0_1	0.8695	-	-	-	-	-	-	-
d0_2	0.0656	-	-	R6K	-	-	-	-
d0_3	0.0349	-	-	-	-	-	-	-
d0_0	0.0301	-	-	-	-	-	-	-
d3_0	0.8361	-	-	-	-	-	-	-
d3_1	0.0822	-	-	-	-	-	-	-
d3_2	0.0817	-	-	-	-	-	-	-
d59_5	0.6393	T5M	-	-	-	-	-	-
d59_3	0.1097	-	S8A	E3A	-	-	-	-
d59_0	0.0945	F6L	-	-	-	-	-	-
d59_1	0.0729	F6L	V7A	-	-	-	-	-
d59_2	0.0508	F6L	-	-	-	-	-	-
d59_4	0.0328	F6L	-	-	R1Q	-	-	-
d165_0	0.6338	T5M	I9V	-	-	-	-	-
d165_3	0.1919	F6L	S8A	R6K	R1Q	-	-	-
d165_1	0.1156	F6L	V7A	-	-	-	-	-
d165_2	0.0587	-	I9V	R6K	I6V	-	-	-
d476_1	0.3696	T5M	I9V	-	-	N4K	-	-
d476_0	0.2452	T5M	I9V	-	-	-	-	-
d476_4	0.2016	Y2F	I9V	-	-	N4K	-	R4K
d476_3	0.1274	Y2F	I9V	-	-	-	-	-
d476_2	0.0561	T5M	I9V	-	-	N4K	-	-
d1543_5	0.2613	Y2F	I9V	-	R1Q	E2D	S1T	-
d1543_4	0.2062	T5M	I9V	Y9F	R1Q	E2D	S1T	-
d1543_2	0.1379	T5M	I9V	Y9F	R1Q	E2D	S1T	-
d1543_1	0.0886	Y2F	I9V	Y9F	R1Q	E2D	S1T	-
d1543_3	0.0841	Y2F	I9V	Y9F	R1Q	E2D	S1T	-
d1543_0	0.0839	Y2F	I9V	Y9F	R1Q	E2D	S1T	-
d1543_7	0.0791	T5M	I9V	-	R1Q	E2D	S1T	-
d1543_6	0.0589	T5M	I9V	Y9F	R1Q	E2D	S1T	-

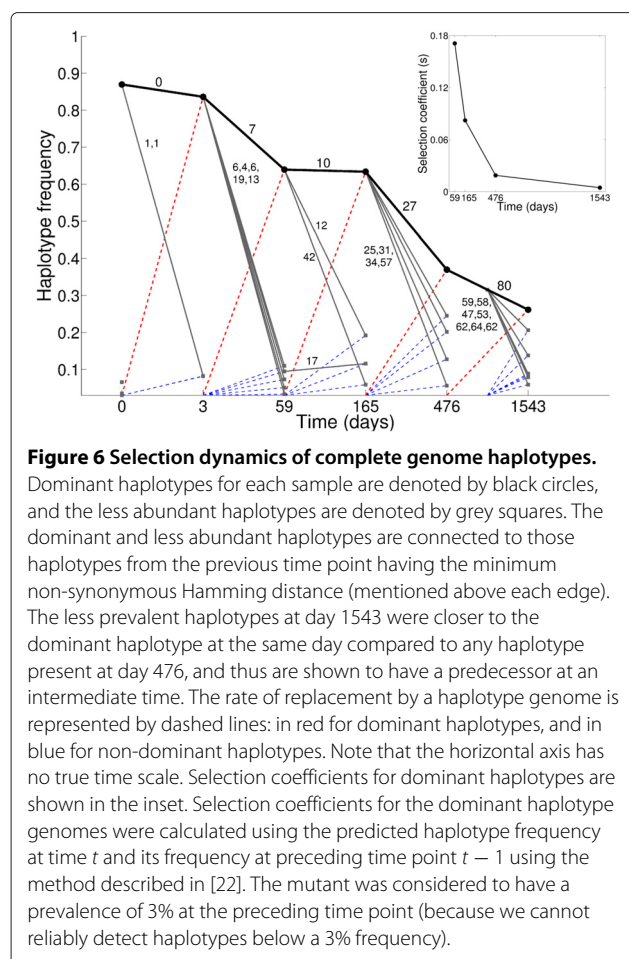
Figure 5 Epitope escapes in complete genome haplotypes. The escape mutations acquired in the seven epitopes identified by Henn et al. [11] are given per reconstructed haplotype for each time point. Haplotypes reconstructed from each temporal sample are given in a descending order of their frequencies. The dominant haplotype for each time point is depicted by the same colors as in Figure 3. Multiple escapes for the same epitope are depicted by different colors. Escape mutations for each epitope are denoted by a 3-letter code with the first letter as the original epitope amino acid, second letter as the position in the epitope and the third letter as the mutated amino acid. Mutations at anchor residues are highlighted.

responses like *pol* and *env* were genetically linked with several sites (cluster II, Figure 7).

The difference in linkage profiles can be explained by the homogeneity of the HIV-1 population at the start of infection (Figure 2B). The effect of genetic hitchhiking is clearly demonstrated by the large linkage cluster comprised of links between sites from *gag* (red) to *nef* (green) which probably have little or no functional linkage (cluster II, Figure 7). The hitchhiking mutations can be mildly advantageous or neutral [30], or may even be potentially deleterious [31]. For example, 22 non-synonymous mutations and 2 synonymous mutations (probably neutral) in the *env* gene have hitchhiked with 1 beneficial *env* epitope escape mutation (purple circles in cluster II, Figure 7). Thus, several non-epitope mutations hitchhike with immune escape mutations. The presence of genetic hitchhiking leading to multiple co-evolving genetic loci may influence the phylogenetic signal, especially if the number co-evolving loci is high. In our analysis we find that the phylogenetic signal for *env*, *gag* and *pol* genes, containing most of the genetically linked sites, is supported by strong bootstrap values at the late time point branches confirming that the topology is robust.

The haplotype reconstruction pipeline, using a combination of read clean-up and PredictHaplo, highlighting the importance of clonal interference and genetic hitchhiking. Our study provides the first evidence of clonal interference in HIV-1 genomes, and is one of the few demonstrating clonal interference in viral evolution. Our results indicate that CTL selection is not the only driving force shaping the evolution of escape mutations. Clonal interference and the genetic background of escape mutations both play an important role in viral evolution, and should be considered to calculate CTL selection pressures and HIV-1 escape rates [46]. Several studies have demonstrated that different epitopes escape mutations evolve in the same or in different epitopes and get fixated in the population. However, this can be a result of either genetic drift or selection or clonal interference. Using whole genome haplotypes, we show that clonal interference plays an essential role in the evolution and fixation of escape mutations.

Our study extends the scope of analysis that can be conducted using next-generation sequencing reads, and unraveled novel within-host evolutionary dynamics. Our pipeline is not limited by the sequencing technology, and the longer reads obtained using existing sequencing



technologies, or new sequencing technologies like MiSeq (with 250 bp paired-end reads) and PacBio, can only improve the quality of haplotype reconstruction. The same approach can be applied to reconstruct haplotypes from MiSeq datasets (with 250 bp paired-end reads) by replacing the Roche specific read clean-up algorithm with a Illumina specific read clean-up algorithm. Henn et al. [11] provided assembled consensus sequences for each temporal sample which we used as reference sequences for our analysis. We performed the same analysis using the HXB2 reference sequence and found that this resulted in reduction of the number and quality of reconstructed haplotypes. Thus, proper read correction and clean-up is crucial for reconstructing global haplotypes.

A major drawback of this methodology may be that we miss low frequency recombinants in the population. Recombination during HIV-1 replication may facilitate the accumulation of multiple epitope escapes in single genomes. But during acute infection, the rate of recombination is known to be rather low [46-48], suggesting that most epitope escape mutations are sequentially acquired

by the quasispecies [46]. Note that recombinant haplotypes that become sufficiently prevalent in the population will be detected as novel unique haplotypes. We simulated a population containing two genomes (each present at 44.5% frequency) taken from LHV simulations and generated a recombinant genome (present at 11% frequency). Three haplotypes corresponding to 3 true genomes were reconstructed at frequencies comparable to the true frequencies. The reconstructed recombinant genome had a Hamming distance of 25 from the true recombinant genome and both the reconstructed parent genomes had a Hamming distance of 20 from the true parent genomes, confirming that our pipeline can reconstruct recombinant genome prevalent in the population. Moreover, most of the errors were found in the first and last 20 bases similar to other control simulations (Table 2). Other simulations with higher prevalence (33% and 55% frequency) of recombinant genome lead to similar results. One possible way to detect presence of recombinants in reconstructed haplotypes would be to apply recombinant detection tools like recombinant identification program (RIP) [49], and SplitsTree [50]. Using RIP, we did not find any convincing evidence of recombination between day 476 haplotypes that might have given rise to any of the day 1543 haplotypes.

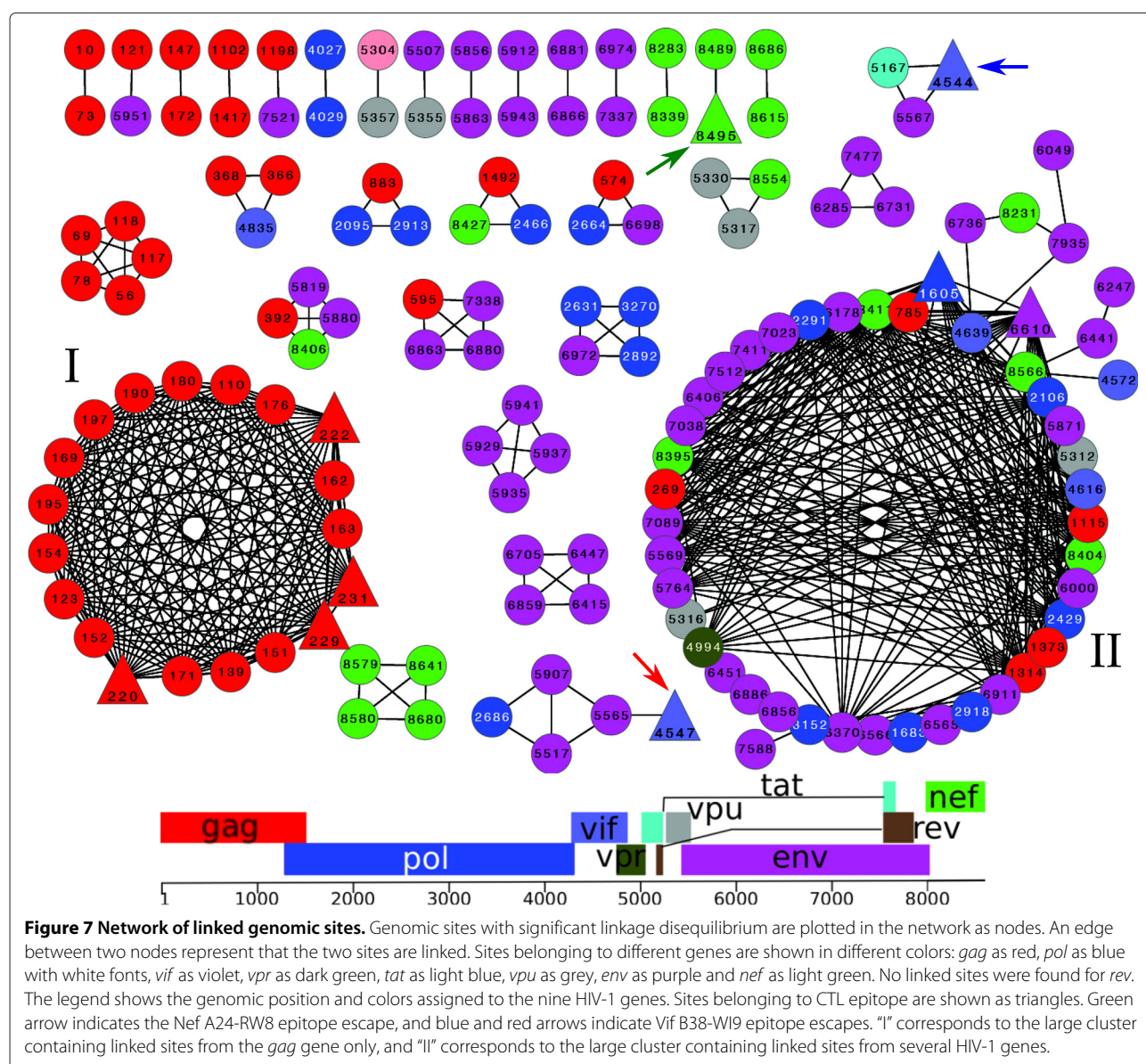
Conclusions

We have shown that whole genome HIV-1 haplotypes can be reconstructed from short 454-sequencing reads with high fidelity. Reconstruction of genome haplotypes provides an opportunity to study the interaction between epitope escapes. Several epitope escapes evolve in single haplotypes, and different combinations of multiple epitope escapes become prevalent in the viral population. As a result of clonal interference, the fate of each escape mutation, also depends upon the fitness of other immune escape mutations prevalent in the viral population. The long range linkage disequilibrium between genomic sites suggests that clonal interference between HIV-1 genomes results in the fixation of several non-epitope mutations via genetic hitchhiking.

Methods

Simulated datasets generated for validation

To validate the haplotype reconstruction pipeline, we created multiple simulated HIV-1 populations. For each of these simulated datasets, we first generated 9 mutated haplotype genomes from a reference HIV-1 genome (GenBank accession number: JQ403055) by randomly selecting r sites such that $n/2 \leq r \leq n$, where n represents either 2, 4 or 10% sites of the total number of sites. Three data sets U2, U4 and U10 (with varying nucleotide diversities of 2, 4 or 10%) were made with a uniform frequency distribution, i.e. all haplotypes were present at the same frequency



(1/9) in the population. Three other data sets were made with a log-normal frequency distribution, with the same nucleotide diversities: 2, 4 and 10% for L2, L4 and L10, respectively. To implement a log-normal frequency distribution, some haplotype genomes were duplicated multiple times in the population to increase their frequencies. Thus, a typical population with a log-normal distribution consisted of 9 haplotypes present at the following copy numbers: 20, 12, 4, 2, 1, 1, 1, 1, 1, respectively. This generated a quasispecies with 5 minority haplotypes present at a frequency below 2.5%, and one dominant haplotype with a prevalence exceeding > 45%. To model heterogeneity in mutation rate over HIV-1 genome, we added 3 hypervariable regions to genomes with 1% nucleotide diversity in the LHV data set. In the hypervariable regions,

50% of nucleotides had 10% nucleotide diversity, while the remaining sites had a 1% nucleotide diversity. We used a log-normal frequency distribution, similar to the L1 data set.

Additionally, we simulated 10 populations with a quasispecies structure. For each "quasispecies data set" (LQ4_i), we generated nine master genomes at a 4% diversity (similar to those above). After creating the 9 master genomes, we created mutated copies with at most 0.1% nucleotide diversity from the corresponding master sequence. This created a cloud of genomes around each master genome. The frequency of the nine quasispecies were distributed log-normally in the LQ4 dataset at the following copy numbers: 50, 24, 8, 4, 2, 2, 2, 2, 1 (as shown in Figure 1A).

We used the ART_454 software [42] to simulate 454 sequencing errors and generate *in silico* reads with an average coverage of 500 reads per base, mimicking the average depth obtained in Henn et al. [11]. The *in silico* reads obtained were analyzed using the haplotype reconstruction pipeline. A lower average coverage of 200 reads hardly changed the number of reconstructed haplotypes (results not shown).

Longitudinal dataset used to study the within-host dynamics of HIV-1 haplotypes

The 454 sequence reads from six datasets V4137, V4136, V4139, V4140, V4676 and V4678 sampled from subject 9213 at day 0, 3, 59, 165, 476 and 1543 were downloaded from the NCBI SRA database [51]. As described in Henn et al. [11], the sequences were obtained by amplifying four overlapping PCR amplicons which span the entire protein coding region of HIV-1 genome. Subject 9213 was HLA-typed, and expressed the A01, A24, B38, B44 and Cw04 alleles. Henn et al. [11] recognized dominant and sub-dominant CTL responses against 7 HIV-1 epitopes in subject 9213 and described the immune escapes. We study the dynamics of these 7 escape mutations using the reconstructed haplotype genomes.

Haplotype reconstruction pipeline

The haplotype reconstruction pipeline consists of two major steps: 1) Read clean-up, and 2) Prediction of haplotypes. Reads obtained from 454 NGS are known to contain three major types of process errors: a) carry forward and incomplete extension errors, b) homopolymer miscall errors, and c) InDels in non-homopolymer regions [11,36]. These 454 sequencer specific errors were corrected using the default parameter settings of ReadClean454 v1.0 (or RC454) software which uses *Mosaik* aligner [52] to align sequences to a reference genome sequence [11]. The RC454 cleaned reads were then subjected to PredictHaplo (version 0.5) [53] for haplotype reconstruction. We used the sample's consensus assembly sequence obtained from Henn et al. [11], as the reference genome sequence for RC454 and PredictHaplo. We tested haplotype reconstruction at different parameter settings for PredictHaplo, and the default settings gave the most reliable results (results not shown). The cleaned-up reads obtained with the RC454 package were made compatible with PredictHaplo using customized Python scripts.

Phylogenetic analysis

For phylogenetic analysis, genes and genomes from the reconstructed haplotypes were aligned using ClustalW [54]. The complete genome, *pol*, *env* and *gag* alignments, used in Figure 3, were 8878bp, 3040bp, 2645bp and 1546bp long, respectively. Phylogenetic trees were

reconstructed using the maximum likelihood (ML) method performed by PHYML (version 3.0) [55]. ML analysis was conducted using the general time reversible (GTR) model, with six substitution rate categories for the gene phylogenies, and nine for the complete genome phylogeny, with the proportion of invariant sites set to 0. The gamma distribution parameter and nucleotide frequencies were estimated from the dataset for the phylogenetic reconstruction. From the parsimonious starting tree, the best tree generated using both the nearest neighbor interchange (NNI) and the subtree pruning and regrafting (SPR) approaches is shown in Figure 3. 100 bootstrap replicates were performed to assess the support for the branches and bootstrap support values ≥ 90 are indicated on the branches. For Figure 1A and Figure 2A, neighbor-joining phylogenetic trees were reconstructed using ClustalW because of their high sequence similarity [54]. To distinguish between synonymous and non-synonymous substitutions, we generated phylogenetic trees using MG and YAP codon substitution models performed by CodonPhyML [56]; however, the topology of the codon based phylogenetic trees was similar to the nucleotide trees.

Selection and linkage analysis

To estimate the fitness and selection coefficients of mutated viral genomes, we used the method described by Maree et al. [22], which depends upon the frequency of wildtype and mutated viral genomes, and the time of evolution. To estimate selection coefficients, we considered that the frequency of a mutant haplotype rises from a value of 0.03 (frequency threshold below which we did not detect any haplotype) at $t - 1$ to the observed frequencies at time t replacing its predecessor [57]. Note that the slow down of the selection coefficients is likely to be biased by the long intervals of late time points.

To detect the extent of positive selection, we first codon aligned the gene sequences using CodonAlign tool [58] and then used the random effects likelihood (REL) method from Datamonkey server [59,60]. Linkage disequilibrium between all polymorphic genomic sites was calculated using DNAsp and linkage values only significant after Bonferroni correction ($p\text{-value} < 0.4 \times 10^{-7}$) were considered [61]. Linked sites were plotted as a network using Cytoscape (version 2.8) [62].

Additional file

Additional file 1: Figure S1. Sequence alignments of the reconstructed haplotype and the corresponding quasispecies. Sequence alignment of four most prevalent haplotypes (A) haplotype_4, (B) haplotype_0, (C) haplotype_3, and (D) haplotype_5 and the corresponding quasispecies. Mismatches between the simulated genomes and the reconstructed

haplotype are indicated as vertical lines with different colors corresponding to different nucleotides (A: Green, T: Red, G: Orange, C: Light blue and Gaps: Gray). The variation at each genomic site was present only in a subset of genomes demonstrating that the haplotypes are reconstructed as consensus sequences of the corresponding quasispecies.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

Designed and analyzed the experiments: AP and RDB. Performed the experiments: AP. Wrote the article: AP and RDB. Both authors read and approved the final manuscript.

Acknowledgments

We thank Thomas Cuypers, Chris van Dorp, Lidija Berke and Bram Gerritsen for discussions and carefully reading the manuscript. This study was also supported by the "Virgo consortium" funded by the Dutch government [Project number FES0908] and by the "Netherlands Genomics Initiative (NGI)" [Project number 050-060-452].

Received: 12 February 2014 Accepted: 24 June 2014

Published: 4 July 2014

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doi:10.1186/1742-4690-11-56

Cite this article as: Pandit and de Boer: Reliable reconstruction of HIV-1 whole genome haplotypes reveals clonal interference and genetic hitchhiking among immune escape variants. *Retrovirology* 2014 **11**:56.

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