Virus and host determinants of HIV-1 infection and AIDS pathogenesis

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Chapter 8

'Evolution of SI and NSI biological virus clones in relation to replication kinetics during the course of HIV-1 infection'

A.B. van 't Wout, H. Blaak, L.J. Ran, M. Brouwer, C. Kuiken, H. Schuitemaker


Determinants which govern these biological properties have been widely studied in the analysis of HIV-1, especially in the viruses (V) replication properties, although secondary properties like sur. off, che, and sur. have also been shown to influence replication rate in certain cell lines.
Evolution of syncytium-inducing and non-syncytium-inducing biological virus clones in relation to replication kinetics during the course of human immunodeficiency virus type 1 infection

To investigate the temporal relationship between human immunodeficiency virus type 1 (HIV-1) replicative capacity and syncytium-inducing (SI) phenotype, biological and genetic characteristics of longitudinally obtained virus clones from two HIV-1-infected individuals who developed SI variants were studied. In one individual, the emergence of rapidly replicating SI and non-syncytium-inducing (NSI) variants was accompanied by a loss of the slowly replicating NSI variants. In the other subject, NSI variants were always slowly replicating, while the co-existing SI variants showed an increase in the rate of replication. Irrespective their replicative capacity, the NSI variants remained present throughout the infection in both individuals. Phylogenetic analysis of the V3 region showed early branching of the SI variants from the NSI tree. Successful SI conversion seemed a unique event since no SI variants were found among later-stage NSI variants. This was also confirmed by the increasing evolutionary distance between the two subpopulations. At any time point during the course of the infection, the variation within the co-existing SI and NSI populations did not exceed 2%, indicating continuous competition within each viral subpopulation.

Human immunodeficiency virus type 1 (HIV-1) is susceptible to genetic recombination and has an error-prone reverse transcriptase enzyme, which combined with the absence of proofreading leads to a misincorporation rate of $10^{-4}$ to $10^{-5}$ per base, or approximately one misincorporation per genome per replication cycle. With $10^{10}$ new viruses produced each day and a half-life of approximately 6 h, an HIV-1-infected individual harbors a swarm of closely related viruses that comprise the so-called HIV-1 quasispecies. These HIV-1 variants have been shown to differ in biological properties such as replication rate, cell tropism, and syncytium-inducing (SI) capacity. Virus isolated in the early asymptomatic phase of infection is predominantly slowly replicating, macrophage-tropic, and non-syncytium-inducing (NSI) in vitro. During progression to disease rapidly replicating, more T-cell-tropic viruses appear, in about 50% of infected individuals associated with the emergence of SI variants.

Determinants which govern these biological properties have been mainly mapped to the envelope gene, especially to the variable (V) regions, although accessory genes like nef, vif, vpr, and vpu have also been shown to influence replication rate in certain cell types. Chimeric clones, constructed from an SI and an NSI molecular clone, showed that exchange of the gp120 V1 - V2 fragment together with the V3 fragment was sufficient to confer SI capacity. Sequence analysis of the V3 fragment of a large panel of HIV-1 isolates with distinct biological phenotypes demonstrated the presence of positively charged amino acids at either one or two fixed positions in the V3 loop of SI variants. During the transition from NSI to SI phenotype, the hypervariable V2 region is thought to undergo increases both in length, mainly through insertion of potential N-linked glycosylation sites, and charge.

Although replicative capacity and SI phenotype in general are coinciding biological features, their temporal relationship in the pathway of virus phenotype evolution is not known. SI variants could evolve from rapidly replicating NSI variants which are more likely to accumulate the mutations required for NSI to SI transition. Alternatively, SI variants could also initially replicate slowly but evolve to rapidly replicating HIV-1. When the V3 sequences of virus populations present around the time of NSI to SI conversion were analyzed, few V3 sequences intermediate between the NSI and SI variants were detected, in agreement with the hypothesis that less fit stages may have to be
crossed in order to reach the more fit SI stage\textsuperscript{(203)}.

In this study, we analyzed the temporal relationship between different biological properties of HIV-1. We studied replication kinetics of biologically cloned SI and NSI variants in two individuals who developed SI variants in the course of infection. From all virus clones, V3 sequences were determined and used for phylogenetic analyses. For comparison, viruses isolated around the time of AIDS diagnosis from two individuals harboring only NSI variants throughout infection were analyzed.

**MATERIALS AND METHODS**

**Study subjects.** Four participants of the Amsterdam Cohort Studies on HIV infection and AIDS in homosexual men were analyzed. Characteristics of the four study subjects are depicted in Table 1. Three subjects (ACH0039, ACH0208, ACH0424) seroconverted at entry into the cohort studies and progressed to AIDS within 5 years thereafter. One subject (ACH0142) was seropositive at entry into the cohort studies and progressed to AIDS 9 years after seropositive follow-up.

**Limiting-dilution analysis.** To isolate biological virus clones and estimate the frequency of productively infected cells, limiting-dilution cultures were performed as described previously\textsuperscript{(194,201)} Briefly, participant peripheral blood cells (PBMC; 0.5 x 10\textsuperscript{6} cells per well; 24 to 96 replicates per concentration) were cocultivated with phytohaemagglutinin (PHA)-stimulated healthy donor PBMC (10\textsuperscript{6} per well) in 96-well microtitre plates. Every week for five weeks, 65 \mu{l} of each culture supernatant was transferred to 25-ml culture flasks, and 10\textsuperscript{5} fresh PHA-stimulated healthy donor PBMC were added to propagate the culture. The proportion of productively infected CD4\textsuperscript{+} T cells was estimated by the formula for Poisson distribution: $F = \frac{F_0}{F_0}$, in which $F_0$ is the fraction of negative cultures. PBMC from wells tested positive were transferred to 25-ml culture flasks containing 5 x 10\textsuperscript{6} fresh PHA-stimulated PBMC in 5 ml of culture medium to grow virus stocks. All viruses obtained from one individual were grown on target PBMC from one seronegative blood donor. Virus containing cell-free culture supernatant was stored at -70°C until further use, cells were frozen, and approximately 10\textsuperscript{5} cells were used for isolation of DNA. SI capacity of virus clones was determined by cocultivation with MT2 cells\textsuperscript{(39)}.

**In vitro characterization of virus replication rate.** Target PBMC from the same seronegative healthy CCR5 homozygous wild-type blood donor (determined as previously described\textsuperscript{(394)}) were used in all replication experiments. All viruses obtained from one individual were analyzed within the same experiment. The titer of the virus stocks was quantified by determination of the 50% tissue culture infectious dose (TCID\textsubscript{50}) in PHA-stimulated healthy donor PBMC. From each virus clone, 10\textsuperscript{2} and 10\textsuperscript{3} TCID\textsubscript{50} was added to 5 x 10\textsuperscript{6} 2-day PHA-stimulated PBMC derived from the same blood donor on which the titer was determined. Cells were incubated for 2 h at 37°C in a shaking water bath in 15-ml conical tubes in a 1.5-ml volume. PBMC were then washed twice, resuspended in 5 ml of culture medium containing reconstituent interleukin-2 (rIL-2; 20 U/ml; PROLEUKIN; Chiron Benelex B.V.), and cultured for 14 days in a 25-ml culture flask. After 5, 8, and 11 days, approximately 3 x 10\textsuperscript{6} fresh stimulated target cells were added in 3 to 4 ml of culture medium containing rIL-2 (30 U/ml). Samples (75 \mu{l}) for determination of p24 production were harvested every day after infection for 14 days and stored at -70°C. All samples obtained for virus clones from one individual were tested for p24 production at the same time, using an in-house p24 ELISA. Cultures were considered positive for virus production when p24 antigen levels exceeded background twofold, which equaled 31 ng p24/ml. p24 production per milliliter of supernatant was determined and corrected for the differences in volume of culture supernatants between the moments of sampling. Due to the relatively high detection limit of our p24 ELISA, the very early replication kinetics are not measured.

**DNA isolation, PCR and sequencing.** Total DNA from PBMC harboring the biological HIV-1 clones was isolated as described previously\textsuperscript{(474)}. Envelope V3 sequences from subject ACH0208 were amplified by PCR as described previously\textsuperscript{(474)}. PCR Products were purified with a GeneClean kit (BIO 101; Inc. Vista, CA) and sequenced directly using the dye-deoxy-chain termination method with Sequenase (U.S.Biochemical, Cleveland, OH), both as instructed by the manufacturers.

Envelope V3 sequences from subject ACH0039 were amplified by PCR using primers Seq1 (5'-TGATACATGTTG66CCACACAT166C-3'), nucleotide positions 6417-6443, sense) and Seq2 (5'-TCTTCTCATATCTCTCTCTCAGTC-3', positions 7629-7653, antisense) in the presence of 3 mM MgCl\textsubscript{2}, in the first reaction and primers Seq5 (5'-GTCACTACTGCTGTTAATG66C-3', positions 6988-7012, sense) and Seq6 (5'-ACTCCACATTATCCACCTG666A666-3', positions 7532-7556, antisense) in the presence of 3 mM MgCl\textsubscript{2} in the nested reaction. For amplification the following PCR amplification cycles were used: 5 min at 95°C once, 1.5 min at 95°C, 1 min at 55°C, and 1 min at 72°C, repeated 25 times. Following a
Table 1. Characteristics of the four study subjects under study*  

<table>
<thead>
<tr>
<th>Subject</th>
<th>CCR5 genotype</th>
<th>Time of:</th>
<th>AIDS defining events</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACH0142</td>
<td>wt</td>
<td>Nov. 1984 (E)</td>
<td>NA</td>
</tr>
<tr>
<td>ACH0424</td>
<td>wt</td>
<td>Aug. 1988 (S)</td>
<td>NA</td>
</tr>
</tbody>
</table>

*wt, wild type; hz, heterozygous; NA, not applicable; S, seroconversion; E, entry

5-min extension at 72°C and subsequent cooling to 4°C. Nested PCR products were purified with QIAquick PCR purification kit (Qiagen, Hilden, Germany). Dye terminator cycle sequencing with AmpliTaq DNA polymerase (Perkin-Elmer, Applied Biosystems Division, Foster City, CA) was performed with the following amplification cycles: 30 s at 92°C, 15 s at 50°C, and 4 min at 60°C, repeated 25 times with the sense nested PCR primer Seq5. Sequence analysis was performed on ABI 373S automated sequencer according to the manufacturer's protocol.

Phylogenetic and sequence data analysis. Alignment of the sequences was straightforward and done with the Pileup program, checked manually and keeping codons intact. Phylogenetic analyses were done with the neighbor-joining program (490) as implemented in the PHYLIP package (490). Bootstrap resampling was used to assess the strength of support for each branch of the phylogenetic trees (490). For bootstrapping, the SEQBOOT, DNADIST, and CONSENSE programs from this package were used. PHYLIP's DRAWTREE program was used to produce the plots. For direct comparison of nucleotide sequences, the number of mismatches as a proportion of sequence length (Hamming distance) was used (490). The distance matrix input for the neighbor-joining analysis was generated using Kimura's two-parameter estimation for nucleotides (492). Estimation of the number of silent and nonsilent substitutions was done according to Nei and Gojobori's method (493) as implemented in MEGA (494). In the statistical analysis of the data, to avoid the problem of dependence between data points, instead of the commonly used pairwise Hamming distance between clones (which results in n(n-1)/2 data points for n sequences), we calculated the distance between all clones and their consensus sequence, which results in n data points for n sequences, thus evading the problem of correlated observations. The significance of differences between the variation of groups of clones was evaluated using the test.

Nucleotide sequence accession numbers. All newly generated sequences have been deposited in GenBank under the following accession numbers: for ACH0039, AF021478, AF021494, AF021495, AF021499, AF0214503, AF021505, AF021508, AF021510, AF021513, AF021514, AF021516, AF021517, AF021518, AF021521, AF021522, AF021523, AF021524, AF021532, AF021533, AF021536, AF021537, AF021540, AF021542, AF021607, AF021608, AF021610, AF021611, AF021612, AF021613, AF021614, AF021616, AF021617, AF021618, AF021620, AF021622, AF021646, AF021647, AF021650, AF021651, and AF021652.

RESULTS

Replicative capacity of longitudinally obtained biological virus clones. From two participants (ACH0039 and ACH0208) from the Amsterdam cohort, biological HIV-1 clones were isolated at several time points between seroconversion and AIDS diagnosis. Individual clones were compared for SI/NSI phenotype and analyzed for replication kinetics. It is important to note that not all virus clones detected in the limiting-dilution experiment were used in the replication experiments, but only a random sample of 6 to 10 virus clones from each time point. The NSI clones detected in the 17 months sample from subject ACH0039 were lost due to technical difficulties. In each experiment, the viral replication kinetics after infections with 100 TCID50 confirmed the observations after infection with 1,000 TCID50 (data not shown). Therefore, only the results for the inoculation with 1,000 TCID50 are shown in the figures. Four slowly replicating NSI variants were isolated 10 months after seroconversion of subject ACH0039, but the majority of the isolated NSI virus clones already had rapid replication kinetics (Fig. 1, top left panels). The first and rapidly replicating, SI variants were obtained 17 months after seroconversion. Throughout follow-up, co-existing SI and NSI variants that had similar rapid replication kinetics
Evolution of SI and NSI variants

Fig. 1. Replication kinetics of SI (•) and NSI (o) HIV-1 clones during the course of HIV-1 infection in two subjects harboring both SI and NSI virus variants. From each virus clone, $10^3$ TCID$_{50}$ was added to $5 \times 10^6$ 2-day PHA-stimulated PBMC derived from the same donor on which the titer was determined. After 5, 8, and 11 days, approximately $3 \times 10^6$ fresh stimulated target cells were added. Samples (75 µl) for determination of p24 production were harvested every day after infection for 14 days. Each line represents the results obtained for one virus clone, * indicates NSI clone 12B3 with arginines at positions 10 and 25. AIDS diagnosis for ACH0039 and ACH0208 was at 37 and 54 months respectively, post-seroconversion (post sc).

were isolated (Fig. 1, top right panels).

In both the early and the late phases of infection, different replication patterns were observed in subject ACH0208 (Fig. 1, bottom panels). During the first 9 months after seroconversion of subject ACH0208, only slowly replicating NSI viruses were obtained (Fig. 1, bottom left panels). After 15 months the first, slowly replicating SI variants were detected: these were replaced within 4 months by rapidly replicating SI variants. The SI variants obtained from ACH0208 gained increasing replication kinetics over time, while the co-existing NSI variants maintained slow replication kinetics (Fig. 1, bottom right panels). At the time of AIDS diagnosis, faster-replicating NSI variants were isolated, although replication was still markedly slower than of the co-existing SI variants.

The viruses isolated around the time of AIDS diagnosis from these two patients were compared with those obtained from two individuals harboring only NSI variants throughout infection, ACH0142 and ACH0424. In general, viruses obtained from all four individuals had rapid replication kinetics around the time of AIDS diagnosis (Fig. 2). Vanishingly fast-replicating NSI variants were isolated in both subjects harboring only NSI variants throughout infection. ACH0142 was diagnosed with AIDS at 36 months post-AIDS, while ACH0424 was diagnosed with AIDS at 9 months post-AIDS.

Fig. 2. Replication kinetics of SI (•) and NSI (o) HIV-1 clones in four subjects at the time of AIDS diagnosis. From each virus clone, $10^3$ TCID$_{50}$ was added to $5 \times 10^6$ 2-day PHA-stimulated PBMC derived from the same donor on which the titer was determined. After 5, 8, and 11 days, approximately $3 \times 10^6$ fresh stimulated target cells were added. Samples (75 µl) for determination of p24 production were harvested every day after infection for 14 days. Means and standard errors of the results for 4 to 5 clones per phenotype are shown.
diagnosis, irrespective of the presence of SI variants (Fig. 2).

**Contribution of phenotypically different HIV-1 clones to virus load.** To analyze the significance of slowly replicating co-existing NSI variants, we calculated the contribution of SI and NSI virus clones to the total cellular infectious load in the limiting-dilution analysis. For each time point analyzed, the contributions of SI and NSI HIV-1 variants to the total infectious cellular load are shown in Figure 3. SI variants were first detected 17 and 15 months after seroconversion in ACH0039 and ACH0208, respectively. The equally rapidly replicating co-existing SI and NSI variants in subject ACH0039 contributed equally to the infectious cellular load (Fig. 3, top left). However, also in subject ACH0208 the NSI variants constituted 50% of the total virus population throughout follow-up despite their slow replication rate (Figure 3, top right). The two NSI-harboring individuals, ACH0142 and ACH0424, also showed a gradual load increase over time, reaching viral loads similar to the two SI- and NSI-harboring individuals around the moment of AIDS diagnosis.

**Phylogenetic analysis.** To understand the phylogenetic relationship between the co-existing SI and NSI variants in the two SI-harboring subjects, a region encompassing the V3 domain of the gp120 envelope molecule was sequenced, 330 nucleotides for ACH0039 and 195 nucleotides for ACH0208. When comparing the deduced 65 amino acids sequenced for both subjects, the V3 sequences were distinct, grouping separately into two groups, with a mean Hamming distance between the two groups of 15%. Bootstrap resampling supported the distinction of two separate groups, 100 of 100 bootstraps. The deduced amino acid sequences for the two subjects are shown in Figure 4.

The mean synonymous substitution rate ($d_S$) in the V3 region was 0.025 (± 0.015) for ACH0039 and 0.032 (± 0.030) for ACH0208 over 27 and 40 months of follow up, respectively. The mean rates of nonsynonymous substitution ($d_N$) were 0.036 (± 0.018) and 0.040 (± 0.023), respectively, which in both cases seems higher than the rate of synonymous substitutions but not significantly different due to the large standard deviations. When comparing $d_S$ and $d_N$ for each time point separately, only at the last time point (37 months) for subject ACH0039 did we observe a significant difference (Student's t test $P = 0.004$). Figure 5 shows the changes in $d_S$ and $d_N$ detected during follow-up. Variants in subject ACH0039 had a stable rate of synonymous substitutions (time effect analysis of covariance, $P = 0.23$), whereas the rate of nonsynonymous substitutions increased sharply during follow-up ($P < 0.0001$), resulting in a decreasing $d_S/d_N$ ratio. In subject ACH0208, both the rate of silent substitutions and the rate of nonsynonymous substitutions increased over time (time effect, $P = 0.005$ and $P < 0.0001$, respectively).

Early after seroconversion both subjects harbored a rather homogeneous group of NSI V3 sequences (mean Hamming distance of 1.2 and 0.8%, respectively). For both individuals, the first SI variants isolated were different from the co-existing NSI variants at at least 4 amino acid positions of the 35 amino acids in the V3 loop (mean Hamming distances were 3.8 and 3.9%, respectively at the first time point of detection of SI) (Fig. 4). To strengthen this observation, neighbor-joining trees with bootstrap resampling were constructed (Fig. 6). Indeed, in both patients the SI and NSI variants from all time points were separated but not with very high bootstrap values.
Fig. 4. Deduced amino acid sequences of the V3 region. The sequences are aligned with the consensus sequence of the variants present in the first sample for each patient. Amino acid positions involved in SI capacity are marked (4). Dashes indicate identity with the reference sequence. --, silent mutation compared with reference sequence. (a) Alignment of V3 sequences from virus clones obtained during the course of infection of participant ACH0039. Position 1 corresponds to amino acid 268 of the HXB2 envelope protein. (b) Alignment of V3 sequences from virus clones obtained during the course of infection of participant ACH0208. Position 1 corresponds to amino acid 296 of the HXB2 envelope protein.
Months after seroconversion

Fig. 5. Plots of synonymous ($d_s$) and nonsynonymous ($d_N$) substitution rates in the V3 region of the virus clones obtained in the course of HIV-1 infection from two subjects ACH0039 (a), and ACH0208 (b). Estimation of the number of silent and nonsilent substitutions between all sequences from one time point and the consensus sequence of that time point was done according to Nei and Gojobori’s method (reference 493) as implemented in MEGA (reference 494). Numbers of synonymous (○) and nonsynonymous ($) substitutions from the same time point are shown. *, $P<0.004$.

(92 and 72%, respectively). Total Hamming distances between the variants present at each time point increased over time in both patients (Fig. 7). Variation within each phenotypic group at each time point was relatively stable (mean Hamming distance approximately 1.5%). However, the evolutionary distance between the two groups of variants increased steadily over time (mean Hamming distances were 6.7 and 7.0%, respectively, at the last time point available). Accordingly, Hamming distances were 3.2 and 4.9% between the late NSI variants and the early NSI variants, respectively, and were 3.9 and 4.5% between the late SI and the early SI variants, respectively, for both patients. To test if the course of the variation over time differs between SI and NSI variants, an analysis of covariance was done with the variation (distance of each clone to the consensus) as the dependent variable, the time since seroconversion as a continuous, and SI/NSI as a dichotomous predictor. The time effect was significant in both patients for the comparison between SI sequences and NSI consensus, or NSI sequences and SI consensus (ACH0039, $P = 0.0009$ and $P = 0.0001$, respectively; ACH0208, $P < 0.0001$, $P < 0.0001$, respectively), but only in patient ACH0208 a significant interaction effect was observed, indicating that the NSI and SI variation developed differently over time (ACH0039: NSI $P = 0.58$, SI $P = 0.70$; ACH0208 NSI $P = 0.0001$, SI $P = 0.0009$).

One of the NSI variants (12B3) isolated 15 months after seroconversion from subject ACH0208 had arginines at positions 10 and 25, two amino acids positions that are positively charged in the co-existing SI variants but not in the other NSI variants (Fig. 4). Interestingly, this NSI variant had the highest replication kinetics of the NSI variants present at that time point (Fig. 1, second bottom panel, line marked with 4). The NSI variants present in this individual at 46 months after seroconversion had certain amino

![Fig. 6](image_url)
Evolution of SI and NSI variants

Fig. 7. Plots of Hamming distances for the V3 region between the virus clones obtained in the course of HIV-1 infection from two subjects, ACH039 (a) and ACH0208 (b). Distances were calculated by using DNADIST as implemented in the PHYLIP program. Comparisons between NSI variants, between SI variants, and between SI and NSI variants from the same time point are shown.

...acid changes that were only detected in SI variants before that time (for example, the lysine's at positions 32 and 47, the isoleucine at position 37 and, the glutamate at position 41) (Fig. 4). When the four NSI variants were omitted from the data set the bootstrap value separating the SI and NSI branches of the phylogenetic tree increased from 72% to 91% (data not shown).

DISCUSSION

In this study, we analyzed the temporal relationship between changes in virus replication characteristics and the evolution of SI and NSI virus populations during the natural course of HIV disease. Interestingly, rapidly replicating SI virus clones had emerged more than 3 years before AIDS diagnosis in two individuals'. After the emergence of, and compared to SI variants, the co-existing NSI variants had slower replication kinetics in one subject and similar replication kinetics in the other. The similar replication kinetics of the co-existing SI and NSI variants with very different V3 regions in this latter subject again illustrate that regions other than V3 are important for replication rate.

When SI variants are present in patient PBMC, they always outgrow the co-existing NSI variants in in vitro bulk coculture of the patient PBMC with PHA-stimulated target PBMC (unpublished observations). In vivo, however, irrespective their replicative capacity, NSI and SI HIV-1 clones each constituted 50% of the total infectious cellular load, in agreement with our previous observations. The selective growth advantage for SI HIV-1 as observed in vitro is apparently not present in vivo. It is possible that the NSI variants detected in the periphery originate solely from compartments where virus production from macrophages predominates. Most primary SI variants are not able to infect macrophages in vitro, probably because their entry depends on much higher concentrations of CD4 on the cell surface than is expressed on macrophages. Therefore, it seems likely that virus production from macrophage compartments is predominantly of the NSI phenotype. The SI variants are apparently unable to compete with the direct spread of NSI variants from the macrophages to CD4+ T cells which may occur during the close cell-to-cell contact during antigen presentation.

Alternatively, the persistence of slowly replicating NSI variants may be explained by the existence of different target T cell populations for NSI and SI variants. Recently, several members of the CC and CXC chemokine receptor families have been shown to function as co-receptors for HIV entry. NSI variants primarily use the β-chemokine receptor CCR5, while primary SI variants were shown in vitro to use both CCR5 and the α-chemokine receptor CXCR4. The equal contribution to the infectious cellular load indicates the ability of the slow NSI variants to compete with the more rapid SI variants for the same CCR5-expressing targets. This may be mediated by increased affinity for the CCR5 and/or CD4 receptor, allowing replication in cells with lower expression of these molecules.

Interestingly, the subject that harbored rapid
SI and slow NSI variants was heterozygous for the
32-bp deletion (Δ32) in the CCR5 gene[291,292].
It is likely that the NSI viruses in this individual
have adapted to entry of target cells expressing
lower levels of CCR5, for example, by using other
coreceptors or developing higher affinity for
CCR5 itself. Indeed, we have recently found
evidence for adaptation of viruses from CCR5 Δ32
heterozygous individuals for growth in cells from
CCR5 Δ32 heterozygous blood donors (Blaak H,
unpublished observations). As the NSI virus
variants from subject ACH0208 have slower
replication kinetics than the NSI variants from
subject ACH0039, this adaptation may be
associated with decreased replication kinetics.

The fact that the evolutionary distance
between co-existing NSI and SI populations
increases over time may indeed indicate that in
general there is only little interaction between
these two compartments of HIV-1 replication. The
two compartments evolved away from the common
ancestral sequence with no sign of saturation,
which may indicate that SI variants are newly
generated in an individual rather than transmitted
together with the NSI variant, as the latter
scenario would result in two highly distinct co­
existing SI and NSI populations already at the
very first time point of detection of SI variants,
of which we find no evidence in these patients, nor
in several other cohort participants studied for
this phenomenon (data not shown). Only in one
case, where SI variants were detected
immediately after infection of a new individual did
we find co-existing SI and NSI populations that
were more than 5% apart, reflecting transmission
of both co-existing variants from the donor[193].

After the first emergence of the SI variants in
both subjects, no new SI variants generated from
late NSI variants were detected. Once SI
variants are present it may be more difficult for
newly arising switch variants to compete.
Alternatively, there may only be a certain type of
NSI variants from which SI variants can be
generated. When this stage in the evolution of
NSI variants has been passed, no new SI variants
will emerge or the new SI variants may not be able
to compete with the late NSI variants. Indeed, at
the time of AIDS diagnosis, high virus loads and
rapid virus replication kinetics were found in all
individuals, including the two individuals harboring
only SI variants throughout the course of infection. Apparently, the presence of rapid NSI
variants was sufficient to establish progression to
AIDS without the generation of SI variants.
Possibly, these late rapid NSI variants were
equally cytopathic as the late rapid SI variants in
the other two individuals[479]. Subject ACH0039,
however, harbored rapid NSI variants 10 months
after seroconversion. Still, in this subject SI
variants emerged. This may suggest that the
increased target cell repertoire of SI variants in
some individuals provides enough selective growth
advantage for SI variants to emerge in the
presence of rapidly replicating NSI variants.

In both subjects, the co-existing SI and NSI
populations each are surprisingly homogeneous
throughout the study period. In subject
ACH0039, the average Hamming distance rarely
exceeded 2%, and in subject ACH0208 only after
more than 30 months of infection did the average
Hamming distance per phenotype reach the 5%
level. The homogeneity per phenotypic population
may indicate fierce competition within the NSI
and SI HIV-1 populations, where only the fittest
variant in both groups evolves further, generating
the new virus population detected at the next
time point. The possible recombination observed in
subject ACH0208 suggests, however, that
interaction between the co-existing SI and NSI
variants may occur, which would imply at least
some overlap of target cell populations, in
agreement with the reported CCR5 usage of
primary SI variants[479]. Despite the homogeneity
of each phenotypic virus population at a particular
time point, there is a continued evolution toward
new variants reflected in the increasing Hamming
distance between the NSI and SI virus
populations and between each population and the
populations present at earlier time points.

Overall, sequence variation increased with time
in the two subjects, due to increases in both the
number of synonymous and nonsynonymous
substitutions in ACH0208 and due to increases in
the number of nonsynonymous substitutions only in
ACH0039. Nonsynonymous virus variation in both
p17 and V3 has been shown to be correlated with
immune selective pressures[479,201,487,495-497]. The
HIV-specific cytotoxic T-lymphocyte reactivity
against autologous virus epitopes in the V3 region
in these two subjects is currently under study.
Acknowledgments

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