Faster HIV-1 Disease Progression among Brazilian Individuals Recently Infected with CXCR4-Utilizing Strains

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Abstract

Introduction: Primary HIV infection is usually caused by R5 viruses, and there is an association between the emergence of CXCR4-utilizing strains and faster disease progression. We characterized HIV-1 from a cohort of recently infected individuals in Brazil, predicted the virus’s co-receptor use based on the env genotype and attempted to correlate virus profiles with disease progression.

Methods: A total of 72 recently infected HIV patients were recruited based on the Serologic Testing Algorithm for Recent HIV Seroconversion and were followed every three to four months for up to 78 weeks. The HIV-1 V3 region was characterized by sequencing nine to twelve weeks after enrollment. Disease progression was characterized by CD4+ T-cell count decline to levels consistently below 350 cells/μL.

Results: Twelve out of 72 individuals (17%) were predicted to harbor CXCR4-utilizing strains; a baseline CD4<350 was more frequent among these individuals (p = 0.03). Fifty-seven individuals that were predicted to have CCR5-utilizing viruses and 10 individuals having CXCR4-utilizing strains presented with baseline CD4>350; after 78 weeks, 33 individuals with CCR5 strains and one individual with CXCR4 strains had CD4>350 (p = 0.001). There was no association between CD4 decline and demographic characteristics or HIV-1 subtype.

Conclusions: Our findings confirm the presence of strains with higher in vitro pathogenicity during early HIV infection, suggesting that even among recently infected individuals, rapid progression may be a consequence of the early emergence of CXCR4-utilizing strains. Characterizing the HIV-1 V3 region by sequencing may be useful in predicting disease progression and guiding treatment initiation decisions.

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Introduction

HIV-1 disease progression, as reflected by either CD4+ T cell decline or opportunistic diseases, may be related to host and/or virus characteristics. The observation of the natural history of HIV-1 infection in well-characterized cohorts established before the antiretroviral treatment era indicates that the mean time of disease progression to AIDS is 10 years, although AIDS can develop in as little as two years in a proportion of patients [1]. On the other hand, a proportion of HIV-infected individuals, so-called elite controllers, will not show any CD4 decline over time due to extremely low levels of virus replication, which does not necessarily prevent the HIV-related cell activation or an accelerated aging process [2].

Certain host features are recognized as the main driving force behind disease progression or virus evolution. For example, the CCR5 allele polymorphism in individuals presenting heterozygous deletion of 32 nucleotides (delta 32) is associated with slower disease progression [3] and even better immunologic response to antiretroviral treatment [4]. The same low rates of CD4+ T cell decline are observed in individuals presenting the CCR2-64I mutation [5] or other specific class-I and -II HLA alleles that may have a negative or positive impact on HIV-1 disease progression [6]. Other host-related factors associated with HIV-1 disease
progression include a polymorphism at the SDF1-3’A conserved segment of the 3’ untranslated region of the SDF-1 structural gene transcript, which, in homozygous individuals (SDF1-3’A/3’A), is associated with delayed onset of AIDS [7]. Less clear however, is the relationship between co-infections with some other pathogens that may either increase or decrease cell activation, although co-infection with GBV, which is associated with decreased cell activation [8], is more clearly associated with slower disease progression [9] and better rates of antiretroviral response [10].

The influence of HIV-1 genetic diversity on viral evolution and disease progression has also been recognized. Over time, there is an association between the emergence of CXCR4 tropic viruses and faster disease progression [11]. However, although primary infection is caused by viruses that exclusively use the CCR5 coreceptor, infection by dual-tropic viruses may be associated with rapid disease progression [12]. It has been reported that HIV evolves in a host-specific manner, and even among individuals infected with the same viral strain, disease progression may differ, with the emergence of CXCR4-tropic viruses being neither homogeneous nor predictable [13]. Although controversial, biological differences have also been demonstrated between HIV types/subtypes. It has been documented that HIV replication, transmission, cell activation, and disease progression are lower in HIV-2 infection compared to HIV-1 infection [14]. Interestingly, duplication in the NF-kB site, which has been associated with increased pathogenesis in HIV-1 subtype C [15], was also associated with rapid disease progression in one patient infected with HIV-2 [16]. Furthermore, it has also been reported that disease progression among individuals infected with subtypes D and C is faster than in those infected with subtypes A and A/V in Africa [17], and that subtype D infection leads to faster rates of CD4 cell decline and subsequent virological failure compared to infection with clade B and other non-clade B HIV strains in England [18]. It may be conceivable as well that the genetic diversity of the viruses may influence the pace of HIV-1 tropism change because the emergence of X4 viruses occurs very early among subtype D-infected individuals, and it occurs late in infections caused by subtype C viruses [19,20,21].

HIV epidemics in Brazil present co-circulating HIV-1 subtypes B, F and C, and few Circulating Recombinant Forms. According to predictions using Bayesian Markov chain Monte Carlo methods and the Reversible-jump MCMC method, HIV-1 subtype B emerged in 1971, subtype F emerged in 1981, BF recombinants emerged in 1989, subtype C emerged in 1987, and BC recombinants emerged in 1992 [22]. There is also a sub-lineage of subtype B harboring the GWGR motif instead of the GPGPR motif at the tip of the V3 loop, which was already detectable early in the course of the Brazilian epidemics, in 1983 [23]. Additionally, some reports have suggested that this Brazilian strain (subtype B+) will lead to a slower pace of disease progression than the usual North American/European subtype B present in Brazil [24,25].

In this study, we characterized HIV-1 from a cohort of recently infected individuals in Brazil and attempted to correlate the rate of CD4+ T cell decline with the genetic diversity of infecting HIV strains, including predicted co-receptor use and/or infection by subtype B+. It has been demonstrated that the specificity of genotype-based predictions of HIV-1 CXCR4 use is usually high [26,27], as is the specificity of predictions of virologic response to certain CCR5 inhibitors [28].

**Results**

Seventy-two individuals were initially included in the cohort. Seventy (94.6%) were male, 90% were men who have sex with men and none had acquired HIV through blood exposure. The mean age was 32.7 years, ranging from 20 to 56, the mean viral load was 72,552 (80,443) copies/mL (4.9 log_{10}), ranging from <400 to 703,000 copies/mL (<2.6 to 5.8 log_{10}), and the mean CD4+ T cell count was 573 cells/μL, ranging from 63 to 2,449 cells/μL. It is described here the analysis for the prediction of co-receptor use using false positive rate of 10%, since it correlated best with disease progression in this group of patients. According to the V3 region profile, 12 out of 72 individuals (17%) were predicted to harbor CXCR4-utilizing strains, as predicted by the bioinformatics tool Geno2pheno[coreceptor]. According to the lab tests results from the first visit, 5 out of the 12 patients who were predicted to be infected with CXCR4-utilizing HIV strains and 8 out of the 60 who were predicted to be infected with CCR5-utilizing strains presented with baseline CD4+ T cell counts lower than 350 cells/μL (p = 0.03, Fisher’s exact test). Table 1 depicts the demographic and virologic/immunologic characteristics of individuals infected with R5 or CXCR4-tropic viruses.

Five R5-infected individuals presented transmitted drug-resistant strains, compared to one CXCR4-tropic HIV-infected individual, for a prevalence of 8.3%. Mutations were as follows: K103N (1 X4), D30N (1), L10I/A71T+M184V+K103N (1), K103N/Y181C (1), M184I (1), K103n+M184V+L10I (1). As seen in Table 1, there were no differences in the mean HIV-1 baseline viral load, mean baseline CD4 count, age, HIV-1 subtype, gender, route of HIV-1 transmission, and transmitted drug-resistant strains between individuals infected by R5- or CXCR4-utilizing viruses.

As seen in Figure 1, 68 individuals were infected with subtype B, according to the V3 sequence, two were infected with subtype F, one with subtype C, and one with subtype A, which clustered closer to CRF_2 sequences (data not shown). Forty-two clade B-infected individuals harbored a proline at the tip of the V3 loop that may either increase or decrease cell activation, although co-

| Characteristics | RS (58) | CXCR4 using strains (4) | p |
|-----------------|--------|------------------------|---|
| Gender          | Male   | 58 (96.7%)             | 10 (83.3%) | 0.13 |
| Exposure        | MSM    | 53 (88.3%)             | 9 (75.0%)  | 0.35 |
| Age (mean)      | 32.65 (±1.06) | 33.08 (±3.30) | 0.88|
| Basal Viral Load (log mean) | 4.277 (±0.11) | 4.404 (±0.28) | 0.65 |
| Basal CD4 count (mean) | 566.0 (±27.50) | 617.7 (±184.70) | 0.65 |
| Tip of V3 loop  | B      | 40 (6)                 | 6 (0.44)   |
|                 | B*     | 20 (6)                 | 6 (0.44)   |
| Subtype         | B      | 56 (12)                | 12 (0.47)  |
|                 | non-B  | 4 (0)                  | 0 (0)      |
| CCR5 A32        | 4 (67%)| 1 (83%)                | 0.47 |
| GBV-C           | 17 (28.3%) | 2 (16.7%) | 0.47 |
| HLA Allele B    | 27 (5) | 8 (6.7%)               | 0 (0)      |
|                 | 57 (4) | 4 (6.7%)               | 2 (16.7%)  |
|                 | 58 (4) | 4 (6.7%)               | 2 (16.7%)  |
| NRTI and PI     | 5 (1) | 1 (0.74)               |

*Men who have sex with men. At the tip of V3 loop, strains were classified as B if they harbor a proline or B* if they harbor a tryptophan or a related amino acid.

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**Subtype B**

| # | ****§ | § | Tropism |
|---|-------|---|---------|
| 2018 | CTRPNNTRKSIHGPGRASYATGEIIGDIRQAHG | | R5 |
| 1111 | M. Q. | | R5 |
| 1119 | L. N. Y. | | R5 |
| 1057 | S. P. R. | | R5 |
| 1100 | L. D. | | R5 |
| 1080 | G. D. | | R5 |
| 2027 | D. N. | | R5 |
| 1052 | Q. D. K. | | R5 |
| 1013 | S. R. T. | | R5 |
| 1020 | H. A. D. | | R5 |
| 1022 | L. T. K.T. | | R5 |
| 1043 | K. T. D. N. | | R5 |
| 1054 | R. L. T. | | R5 |
| 1055 | M. K. | | R5 |
| 1060 | NM. TM. A. | | R5 |
| 1064 | GV.L. G. I.T. | | R5 |
| 1068 | G. R. SM. T. D. N. | | R5 |
| 1071 | QL. HT. K. | | R5 |
| 1072 | G. S. D. N. | | R5 |
| 1074 | G. A. R.D. | | R5 |
| 1075 | S.S.F. V. Y. | | R5 |
| 1076 | R. A. D. R. | | R5 |
| 1077 | QR. W.T. A. Y. | | R5 |
| 1078 | Q. P. L. Q. | | R5 |
| 1085 | VT. M. A. N. | | R5 |
| 1089 | L. KVL. K. | | R5 |
| 1090 | V. S. K. T. D. | | R5 |
| 1094 | P. FT. D. N. | | R5 |
| 1098 | LR. EETW. K. KK. | | R5 |
| 1107 | M. KS. D. | | R5 |
| 1110 | J.R. N. W.T. A. Y. | | R5 |
| 1112 | V. D. E. | | R5 |
| 1115 | L. A. A. A. | | R5 |
| 1117 | M. S.FT. D. | | R5 |
| 1121 | G. T. T. D. | | R5 |
| 2025 | R. N. AV. N. | | R5 |
| 2029 | R. L. KS.F. H. N. | | R5 |
| 2042 | A. G. | | R5 |
| 1046 | S.G. L. SV. R.Y. | | X4 |
| 1053 | K. QL. WVT. -KV. N.R.Y. | | X4 |
| 1082 | I. RGMTM. VYF. -K. K. | | X4 |
| 1097 | I. RG.TM. VYF. -K. K. | | X4 |
| 2022 | G. G. ED. | | X4 |
| 2033 | S. G. K.V. EK. N.K. | | X4 |
| 2015 | S. F. | | X4 |
| 1049 | QLVG. T.L. K.E | | X4 |
| 1084 | L.W. N. S. | | R5 |
| 1044 | L.W. T.L. N.Y. | | R5 |
| 1002 | M.W. T. | | R5 |
| 1048 | M.W. T.L. D.V. | | R5 |
| 1066 | M.W. | | R5 |
| 1067 | G. RG. MAW. D. | | R5 |
CD4 T cell decay during short term follow-up.

Individuals with higher viral loads followed by partial recovery of CD4+ T cells suggest that the initial impact on the CD4 T cell count among individuals infected with CXCR4-utilizing strains. The Cox proportional-hazards regression revealed a risk ratio for CD4 T cell decline over time, which is in accordance with the range described in the literature (4% to 15.9%) [31,32,33]. It has been consistently reported that individuals are initially infected with CCR5-utilizing strains. It has been recognized using mathematical models that 75% of individuals will be infected by only one HIV-1 strain whereas the remaining 25% will be infected by two to five HIV strains [34]. This genetic restriction occurs in spite of the transmission route, and it is so significant that only one variant may be selected from several quasispecies from different blood donors after the transfusion of platelets coming from different HIV-infected individuals [35]. Therefore, we believe that the V3 sequence profiles characterized in our study are an accurate representation of the homogeneous quasispecies present at that stage of HIV infection. It is also known that the change in HIV-1 tropism from R5 to X4 receptors generally occurs late in the disease when viruses start to preferentially replicate in the thymus rather than in the gastrointestinal tract, and the detection of viral tropism is indicated to the right of each sequence (CXCR4-using strains are named X4 for the sake of simplicity).

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Figure 1. Amino acid alignment of V3 sequences. Subtype B sequences were aligned with the consensus sequence. Amino acid positions appear at the top of each alignment, with dots indicating identities whereas dashes indicate deletions. # indicates the N-linked glycosylation site, * indicates the GPGR motif at the tip of the V3 loop, and † indicates positions 316 and 323 (HXB2 positions), which are associated with in vitro selection of maraviroc resistance (316T and 323V substitutions are shaded) [29]. Inferred tropism is indicated to the right of each sequence (CXCR4-using strains are named X4 for the sake of simplicity).

(GPG motif), whereas 18 harbored a tryptophan (GWG, the so called B* lineage), and eight other samples presented different amino acids, including phenylalanine (2), leucine, glycine, methionine, threonine and valine (1 each). All these alternative amino acids are encoded by nucleotide triplets that are closer to the one encoding tryptophan (TGG) than the ones that encode proline at this position in this specific group of patients (CCA/G/T), thus suggesting that they are derived from the B* lineage. The A316T substitution was detected in 15 individuals, whereas the I323V substitution was detected in only two subjects; both mutations were selected for by maraviroc in vitro, which lead to a plateau in the PhenoSense HIV Entry Assay (Monogram Biosciences, CA), preventing maximal inhibition by maraviroc [29] (Figure 1). Interestingly, 12 out of the 15 cases presenting A316T substitution appeared among the 26 B*-related strains, compared to three out of the 42 clade B strains (Fisher’s exact test p = 0.0003).

Phylogenetic relationships using all of the samples revealed only one pair of patients presenting evidence of clustering based on the low bootstrap values (below 98%, as previously suggested) [30]. This indicates that the majority of individuals analyzed were epidemiologically unrelated, and clear evidence of convergence towards CXCR4+ sequences was not detected (Figure 2).

Overall, the mean and median CD4+ T cell counts were 551.5 and 529 cells/μL, respectively at baseline, and 537.9 and 475 cells/μL, respectively at visit 6. These CD4+ T cell counts represented a mean and median decline of −13.6 and −54 cells from baseline over this 78-week period (visit number 6). Specifically, the mean CD4 decline was −5.6 cells/μL for R5-infected individuals, compared to −168.9 cells/μL among individuals infected with CXCR4-utilizing strains. The Cox proportional-hazards regression revealed a risk ratio for CD4+ T cell count decline to a level below 350 cells/μL of 0.99 for higher baseline CD4 T cell count (protective effect, p = 0.99) and 5.1 for X4 strains (p = 0.007). When age was kept in the model, the risk ratio for high baseline CD4 remained at 0.99 (p = 0.007), thus suggesting that age enhances the effect of X4 strains on CD4 decline suggesting that age enhances the effect of X4 strains on CD4 decline. Risk ratio for baseline CD4 T cell count (protective effect, p = 0.0001) and 5.1 for X4 strains (p = 0.007). When age was kept in the model, the risk ratio for high baseline CD4 remained at 0.99 (p = 0.007), thus suggesting that age enhances the effect of X4 strains on CD4 decline. Risk ratio for high baseline CD4 T cell count (protective effect, p = 0.0001) and 5.1 for X4 strains (p = 0.007). When age was kept in the model, the risk ratio for high baseline CD4 remained at 0.99 (p = 0.007), thus suggesting that age enhances the effect of X4 strains on CD4 decline.
of X4 strains is correlated with rapid disease progression. (reviewed in reference [33]) Detection of CXCR4-tropic strains early in the course of disease may be attributed to faster evolution from initial R5 to X4 viruses or transmission of dual-tropic viruses, and we believe that these two possibilities may justify the high proportion of CXCR4-tropic viruses found in our cohort.

Not surprisingly, we were able to observe the association between the presence of CXCR4-utilizing strains and faster disease progression as characterized by CD4+ T cell counts below 350 cells/μL. Although baseline CD4+ T cell count was not statistically different between the two groups, the decrease to levels below 359 cells/μL could somewhat be biased by the slightly lower baseline levels in subjects with predicted CXCR4 using viruses (566 versus 611 cells/μL). It is unclear whether the association of faster disease progression with CXCR4-using strains represent a cause or a consequence of HIV-1-related immune suppression. However, the detection of CXCR4-using strains during recent infection in a group of patients with relatively higher CD4+ T cell levels and the observation of the steepest CD4+ T cell decline among these individuals may indicate that this phenomenon is more likely to be the cause rather than a consequence of immune suppression. Therefore, our findings stress the possibility of the presence of strains with more alleged in vitro pathogenicity during the early phases of HIV infection, and also suggest that, even

Figure 2. Phylogenetic relationships between analyzed samples. D85_40 at the top is the North-American/European B consensus sequence. * indicates CXCR4-using strains. Bootstrap values above 85% are indicated at the intersections of branches.
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Disease Progression and HIV Recently Infection
among recently infected individuals, rapid progression may be a consequence of the early emergence of either X4 or dual-tropic HIV strains.

We have not been able to correlate the pace of disease progression and the presence of the Brazil-specific HIV-1 B strains harboring a tryptophan residue at position 16 at the tip of the V3 loop (the so-called B° strains), as had been previously suggested [24,25]. It is interesting to note that B° strains emerged very early in the Brazilian epidemics, accounting for approximately 50% of subtype B strains in Brazil [ref] [23]. It is also interesting that the prevalence of B° strains is lower among patients in lower CD4+ T cell count strata, whereas the prevalence of the classical GPGR strains remains unchanged and the prevalence of strains presenting other amino acids at the tryptophan position increases (V3 position 16). Sera sample from patients harboring most of these strains harboring other amino acids at position 16 will react with specific B° peptides and not with GPGR peptides, thus suggesting that they have evolved from B° strains [23]. As tryptophan is coded by the TGG nucleotide triplet, it is conceivable that the common hypermutation process observed in HIV (G to A substitution) may lead to a disappearance of tryptophan over time because TGA, TAG and TAA are stop codons; therefore, tryptophan may be replaced by other related amino acids. One possible explanation for the sustained prevalence of B° strains in the Brazilian population may be related to the fact that most HIV transmission occurs during the first years of infection, a time when B° strains are still present in an infected individual. We therefore think that the lack of correlation between slower disease progression and B° strains found in our study is accurate because by the time the disease progresses in B°-infected individuals, the tryptophan residue may have already been replaced by other amino acids. One interesting finding was the high prevalence of A316T substitution at the V3 region (HXB2 position) in this set of samples. As previously mentioned, these mutations have been selected by maraviroc in cell culture, leading to decreased susceptibility to this drug [29]. The impact of these mutations has been confirmed by reverse mutation, which

Figure 3. Associations between inferred tropism and time to CD4 decline to levels below 350 cells/μL (survival probability). There was a statistically significant association between the presence of CXCR4-using viruses (X4) in the genotype at baseline and an earlier time to CD4 decline below 350 cells/μL (log rank P = 0.0450). The mean time to CD4 decline below 350 was 51.31 ± 3.66 and 34.14 ± 10.61 months for R5- and X4-infected individuals, respectively.
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Figure 4. CD4 decline over time in individuals infected with R5 or CXCR4-using viruses (mean initial CD4+ T cell counts of 558.5 and 481.4 respectively).
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restored wildtype susceptibility to maraviroc. However, this mutation has been reported to be exceedingly rare in clade B viruses, accounting for only two out of 23,000 sequences in the Los Alamos HIV sequence database [29]. Another interesting finding was the association of A316T substitution with the Brazilian clade B strains (B’), which are extremely common in Brazil and rare elsewhere. Therefore, the phenotypic impact of this substitution on maraviroc susceptibility in B’ strains deserves further confirmation.

We also investigated the influence of other co-factors related to HIV-1 disease progression such as the CCR5 polymorphism, GBVc virema, and presence of “protective” HLA profiles such as HLA B*27, B*57 and/or B*58 and were not able to find any statistic correlation with CD4+ T cell decline, perhaps due to the small number of patients here analyzed.

We recognize that this study may lack sensitivity and/or specificity in detecting patients infected by X4 or dual-tropic/mixed HIV populations because phenotypic assays were not used in this study. However, we achieve the objective of predicting genotypic correlates of HIV-1 disease progression using available tools. Our findings confirm the similar results of another recent study [36], and we believe that our findings demonstrate the necessity of trying to predict HIV-1 co-receptor use among recently infected/diagnosed naïve individuals because it may predict which individuals will be more likely to progress faster in their HIV-related immune deficiency, and it may perhaps suggest that a safer approach in dealing with these individuals could be an earlier initiation of antiretroviral therapy. For the purpose of this sort of HIV-1 characterization, genotype-based techniques may prove to be more cost-effective than phenotypic assays because they are less cumbersome, easily available, and less expensive.

Methods

Ethics Statement

Informed written consent was obtained from all the patients and the study was approved by the Ethics Committees and the Institutional Review Board of the Federal University of São Paulo (#0919/01).

Cohort characterization

A cohort of individuals recently infected with HIV was initiated in May 2002. Individuals seeking free and anonymous testing and counseling services offered by the City of São Paulo Health Department and identified as infected with HIV-1 were offered to undergo the Serologic Testing Algorithm for Recent HIV Seroconversion (STARHS) after signing an initial IRB-approved informed consent. They were invited to join the cohort if recent HIV infection was detected (see below). As of May 2002, we were able to recruit 72 individuals in which env V3 characterization was possible. Recruited individuals were followed every three to four months after the initial clinical visit [37].

Characterization of recent HIV-1 infection using the Serologic Testing Algorithm for Recent HIV Seroconversion (STARHS)
The Vironostika® HIV-1 Micro-ELISA System (bioMérieux Inc., Durham, NC, USA) was used to retest the original HIV-positive samples. The Vironostika LS/EIA tests a 1:20,000 dilution of the specimen under modified incubation conditions. Specimens found to be positive on the S/EIA and negative on the LS/EIA are considered to represent recent infection [38,39]. The method is based upon the slow rate of increase in antibody titers observed during the early period of infection, as well as on data from study subjects with known dates of seroconversion. Based on the calibration of the assays and on a defined threshold, the serologic testing algorithm for recent HIV seroconversion (STARHS) classifies HIV infection as recent or long-standing depending on the differential HIV antibody titer. The assay performs uniformly on HIV-1 subtype B, and the mean window period between seroconversion on the S/EIA and seroconversion on the LS/EIA is 170 days, with a 95% confidence interval (95% CI) of 145–200; we therefore estimated incidence based on the 170-day period. The STARHS was repeated in samples collected six months after the initial visit to document full seroconversion and to confirm the true positive nature of recent HIV infections the enrolled individuals, since 2% of the cases can be falsely classified to be seroconverters. Additionally, false recency rate (FRR) is population specific occurring among AIDS cases, elite controllers, and HAART suppressed patients in the population under study, which were not the case the studied population [40,41]. All testing was performed at the Retrovirology Laboratory of the Federal University of São Paulo, located in São Paulo, Brazil. Volunteers were also seen by an Infectious Diseases physician to determine whether the clinical and laboratory evaluation was compatible with a recent infection.

HIV extraction, amplification, and sequencing

We selected proviral DNA rather than RNA given that, for HIV-1 tropism evaluation, proviral DNA may constitute a more sensitive method to detect CXCR4-utilizing strains [26]. Furthermore, as the patients in this study were evaluated at a time point close to primary infection, the divergence between these two compartments would not have been significant. Samples analyzed were from the second visit, which occurred approximately 9 to 12 weeks after enrollment in the cohort. Proviral HIV-1 DNA was purified using the QIAamp blood kit (Qiagen, Santa Clarita, CA, USA), in accordance with the manufacturer’s instructions. To analyze the V3 region of gp120 I, a 1.2-kb fragment of V1–V5 was amplified in the first round and a 350-bp fragment of C2V3C3 was amplified in the second PCR, as previously described [42]. The protease and reverse transcriptase regions of the pol gene were also amplified to assess primary HIV-1 resistance to antiretrovirals, as previously described [43]. Purified PCR products were sequenced bi-directionally with an ABI PRISM BigDye Terminator Cycle Sequencing Ready Reaction Kit and AmpliTaq DNA Polymerase (Applied Biosystems, Foster City, CA, USA). Sequences were corrected and assembled using the Sequencher 4.0 program (GeneCodes, Ann Arbor, MI, USA).

Sequencing analysis and prediction of co-receptor use

The computer-modeled amino acid sequences were aligned and used as a guide for manual editing of nucleotide sequence alignments. The alignments were generated using the CLUSTAL X program, version 3.0 [44]. For each alignment, phylogenetic analyses were performed using the PHYLYP program package, version 3.57. The DNAdist program was used to calculate distance matrices based on the maximum-likelihood model, and neighboring trees were generated using the Neighbor and Consense programs. Statistical significance was assessed with bootstrap tests in a total of 100 replications. (GenBank accession numbers pending). Co-receptor use was predicted using the bioinformatics tool Geno2pheno[coreceptor] [false positive rate of 3, 10, and 20% - Table S1], coreceptor.bioinf.mpi-inf.mpg.de [27]. We chose to pursue the analysis using false positive rate of 10% which best correlated with disease progression in this group of patients.
Antiretroviral resistance analysis

Mutations related to resistance to NRTI, NNRTI, and PI antiretrovirals were evaluated according to published guidelines, which excluded common polymorphisms [45]. The prevalence of mutations A316T and I323V in the V3 loop of gp120 (HXB2 positions), which display selection by treatment with maraviroc in vitro and are related to decreased susceptibility to this inhibitor in R5 strains, was also evaluated [29].

Detection and quantification of GB virus type C RNA

Viral RNA was extracted and reverse transcribed and a fragment of the nonstructural 5a region (NS5a) was PCR amplified as previously [46,47,48] described. After amplification, 5 ml of the PCR product was used for electrophoresis analysis on a 2% agarose gel. The positive and negative samples were correborated with nested RT-PCR that amplified a fragment of 344 bp of the 50 noncoding region (50 NCR) as previously described. After amplification, 5 ml of the PCR product was used for electrophoresis analysis on a 2% agarose gel.

CCR5 polymorphism

We obtained genomic DNA samples extracted from 300 ml of buffy coat using a QIAamp Blood Kit (QIAGEN Inc, CA), using the methodology indicated by the manufacturer. The presence of CCR5Δ32, representing the heterozygous status of the allele by the host was determined by polymerase chain reaction (PCR) as previously described [49]. Subsequently, amplified products were separated with electrophoresis in a 3% agarose gel for 40 minutes at 110 mV and visualized with ethidium bromide under ultraviolet light. The expected PCR product size was 241 bp for the wild-type and 209 bp for the CCR5Δ32 alleles.

HLA class I genotyping

For each subject a blood sample was taken and genomic DNA was extracted using a QIAamp Blood Kit (QIAGEN Inc, CA), using the methodology indicated by the manufacturer. This assay provided results at an intermediate level (groups of alleles), with assignment of the four-digit allele in some cases.

Characterization of disease progression and statistical analysis

All included individuals were followed at the HIV outpatient clinics of the Federal University of São Paulo. Clinical data, RNA-HIV-1 viral loads and CD4+ T cell counts were collected every three to four months. Disease progression was characterized by the decrease of CD4+ T cell counts from above 350 cells/μL to levels consistently below 350 cells/μL, and/or the initiation of antiretroviral treatment. Antiretroviral treatment was initiated according to the Brazilian Guidelines for HIV-1 Therapy (www.aids.gov.br), which until 2009, recommended treatment for patients with CD4+ T cell counts below 350 cells/μL or in the presence of AIDS-defining conditions. The level of CD4 below 350 cells/μL was arbitrarily chosen since all individuals that reached these levels started with antiretroviral treatment, thus obscuring the disease progression end-point. On the other hand, in this specific group of patients, antiretroviral treatment was not initiated in any individual with CD4+ T-cell levels above 350 cells/μL. Statistical analyses were performed using the Chi-square and Fisher’s exact test. The influence of the HIV-1 V3 sequence on clinical outcomes was assessed by Kaplan–Meier analyses. Cox proportional hazard regression was used to calculate univariate and multivariate risk ratios (RR) and 95% confidence intervals (CI). Baseline variables examined included HIV-1 baseline viral load, CD4+ T cell count, age, HIV-1 subtype, gender, and route of HIV-1 transmission.

Supporting Information

Table S1 Tropism Prediction using genotropheno [coreceptor] with false positive rates (FPR) of 5, 10 and 20%.

Author Contributions

Conceived and designed the experiments: MCAS ECS LMJ RSD. Performed the experiments: MCAS MTMG SS RMC. Analyzed the data: MCAS RSD MTMG RMC. Contributed reagents/materials/analysis tools: MCAS RSD MTMG. Wrote the paper: EGK RSD. Handling samples: HT. Attending physician: MMS.

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