Prevalence of Transmitted Drug Resistance and Impact of Transmitted Resistance on Treatment Success in the German HIV-1 Seroconverter Cohort

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Abstract

Background: The aim of this study is to analyse the prevalence of transmitted drug resistance, TDR, and the impact of TDR on treatment success in the German HIV-1 Seroconverter Cohort.

Methods: Genotypic resistance analysis was performed in treatment-naïve study patients whose sample was available (N = 158/1,276; 12.4%; CIwilson 10.7–14.3; p for trend = 0.25). NRTI resistance was predominant (7.5%) but decreased significantly over time (CIwilson: 6.2–9.1, p for trend = 0.02). NNRTI resistance tended to increase over time (NNRTI: 3.5%; CIwilson: 2.6–4.6; p for trend = 0.07), whereas PI resistance remained stable (PI: 3.0%; CIwilson: 2.1–4.0; p for trend = 0.24). Resistance to all drug classes was frequently caused by singleton resistance mutations (NRTI 55.6%, PI 68.4%, NNRTI 99.1%). The majority of NRTI-resistant strains (79.8%) carried resistance-associated mutations selected by the thymidine analogues zidovudine and stavudine. Preferably 2NRTI/1PIr combinations were prescribed as first line regimen in patients with resistant HIV as well as in patients with susceptible strains (susceptible 45.3%; 173/382 vs. resistant 65.5%; 40/61). The majority of patients in both groups were treated successfully within the first year after ART-initiation (susceptible: 89.9%; 62/69; resistant: 7/9; 77.8%).

Conclusion: Overall prevalence of TDR remained stable at a high level but trends of resistance against drug classes differed over time. The significant decrease of NRTI-resistance in patients newly infected with HIV might be related to the introduction of novel antiretroviral drugs and a wider use of genotypic resistance analysis prior to treatment initiation.

Introduction

Testing for HIV resistance to antiretroviral drugs in plasma is a rational supplement to guide clinicians in their therapeutic choice for patients receiving antiretroviral therapy, ART. Problems of drug failure related to drug resistance as a consequence of individual pharmacogenetics and adherence problems rendering drug levels suboptimal remains a major obstacle for clinicians to choose an effective antiretroviral regime. Once resistant strains of HIV are selected in the host population, they can be transmitted to new hosts and spread in the population at risk. Resistance-associated mutations of HIV have been documented for each of the different classes of antiretroviral drugs [1,2,3]. After a continuous increase of transmitted drug resistance (TDR) over several years in Western Europe stable or decreasing trends of TDR were reported [4,5,6,7,8].

Studies of the impact of primary resistance on treatment response [9,10] are limited by the fact that many have been undertaken before routine baseline resistance testing became common, and therefore inactive drugs were often utilised, and associated with either reduced treatment response, or no effect. Data on the impact of genotypic resistance testing on the composition of first-line therapy are scarce [11,12,13]. Some studies indicate variation among treatment strategies of physicians. Differences in HIV treatment practices have been attributed not only to medical factors but also to non-medical factors such as
patient adherence [14,15,16]. The aim of this study was to investigate the prevalence of TDR and the impact of the infection with resistant HIV on treatment success in patients with a known or well estimated date of seroconversion as best approximation for the date of infection in the German HIV-1 Seroconverter cohort.

Methods

Study population

The study was approved by the local ethical committee of the Charité, University Medicine, Berlin, Germany. Written informed consent was given by all patients. The HIV-1 Seroconverter Study is a nationwide multicentre observational study based on an open cohort of HIV-1-infected persons for whom the date of seroconversion is known (1. acute seroconverters) or can be estimated (2. documented seroconverters).

1. Acute seroconverters: laboratory diagnostic criteria for an acute seroconversion were i) detectable HIV-1 RNA (or p24 antigen) combined with a negative or indeterminate ELISA result or ii) reactive HIV-1-ELISA combined with a negative or indeterminate immunoblot-result followed by confirmation of complete seroconversion within 6 months. In persons recruited during seroconversion the date of the first reactive test was used as an approximation for the infection date.

2. Documented seroconverters: individuals with a last negative and a first positive HIV-antibody test with a maximum three year interval were recruited for the study. In documented seroconverters the mid-point between the dates of the negative and the first positive HIV-antibody test was used to estimate the time of infection.

As we performed a long term observational study, various commercial kits to determine viral load (VL) with different detection limits were used over time. In this analysis viral suppression was determined as treatment success in case of viroaemnic suppression to at least 500 copies/ml including any category of low-level viral load measurement (<400, <50 and <40 copies/ml). Two consecutive viral load measurements within 5 to 12 months after initiation of first line therapy were mandatory for inclusion into the analysis of treatment success. Two or more consecutive viral load measurements above 500 copies/ml were categorised as virological failure. Therapeutic success was defined as viral suppression below the detection limit of at least ≤500 copies/ml. We defined 3 months as a minimum duration of first line treatment. All p-values are two sided, and a p-value of <0.05 was considered significant. All data were analysed using SPSS 17.0.

Results

Population characteristics

Of 1,564 HIV-positive patients with a known or estimated date of seroconversion 1,312 primary samples from drug-naïve patients were available and included into the analysis. All study participants seroconverted between 01.01.1996 and 31.12.2007. Genotyping was performed in 1,276/1,312 available samples of treatment naïve patients (97.3%). There was no bias between the seroconverters genotyped and the group of patients not included (no sample available or already treated at study entry with respect to exposure categories etc (data not shown). Nearly two thirds of patients were documented seroconverters (845/1,276; 66.2%; table 1). 33.8% were acute seroconverters (431/1,267) defined by laboratory criteria. 95.0% of the patients were male (1,212/1,267), 64 patients were female. In Germany data about route of transmission in HIV cases reported through the routine national surveillance system were documented in 85% of all patients newly diagnosed with HIV-1 infection (date of infection not known).

65% were MSM, 17% were heterosexuals, 12% of the individuals originated from high prevalence countries. Intravenous drug use was reported for 5% of all reported HIV cases in 2008 (Epidemiological Bulletin, May 2009, Robert Koch-Institute, Berlin, Germany). Concordantly with all reported HIV-1 cases in Germany, the predominant route of transmission in the German HIV-1 Seroconverter cohort was sex between men, MSM (85.2%). Hence, our study has a bias towards MSM comparing to the national HIV surveillance data. However, the proportion of the transmission route is comparable to the national surveillance data.

NonB subtype was observed in 7.4% of the study participants. CD4 cell counts/µl and plasma viral load measured at time of seroconversion were 510 cells/µl (median; IQR: 380–652) and 5.17 log₁₀/ml (median; IQR: 4.41–5.88), respectively (table 1). Antiretroviral therapy was initiated in 34.7% (443/1,267) defined by laboratory criteria. 95.0% of the patients were male (1,212/1,267), 64 patients were female. In Germany data about route of transmission in HIV cases reported through the routine national surveillance system were documented in 85% of all patients newly diagnosed with HIV-1 infection (date of infection not known).

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patients with a 2NRTI/1PIr regimen changed to another drug class. 30.3% (n = 33) of the 109 patients dropped out and no further information was available. The first-line regimen was censored by the date of data extraction for this manuscript in 11 cases (10.1%). For the remaining 1.8% (n = 2) no further treatment information was available.

Genotypic resistance analysis

Prevalence and epidemiological trends in transmission of drug resistance. Primary mutations according to the SDRM list of Bennett et al. [18,20,21] were identified in 158/1,276 (12.4%) of viral strains. Overall, the prevalence of transmitted drug resistance remained rather stable at 12.4% (CIWilson: 10.7–14.3) during the period of observation (\( p \) for trend over time 1996–2007 = 0.25) (fig. 1). NRTI associated resistance (single class resistance) was identified most frequently (6.3%; 80/1276; CI Wilson: 5.0–7.8). Followed by 2.4% NNRTI resistance (30/1276; CI Wilson: 1.6–3.4) and 2.1% PI resistance (27/1276; CI Wilson: 1.4–3.1). Dual- and multi-class resistance was only seen in 1.4% and 0.2% of patients with TDR, respectively. NRTI resistance reached a mean prevalence of 7.5% (CIWilson: 6.2–9.1), if the numbers were cumulated from all categories of transmitted resistance classes, e.g. mono-, dual- and multi-resistant HIV (NNRTI resistance 3.5% CIWilson: 2.6–4.6; PI resistance 2.9% CIWilson: 2.1–4.0). Prevalence of transmitted NRTI resistance dropped between 1999 and 2000 resulting in an overall declining trend (NRTI: \( p \) for trend over time 1996–2007 = 0.02) (fig. 2). In contrast, prevalence

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<th>Table 1. Characteristics of patients.</th>
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<td>Drug-naive patients, first sample available, no. (%)</td>
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<td>Median age at sc(^a) [years] (IQR(^b))</td>
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<td>Sex, no. (%)</td>
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<td>pol-subtype non-B, no. (%)</td>
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<td>Median viral load at sc [log(_{10})/ml] (IQR)</td>
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<td>Median duration of first-line regimen [days] (IQR)</td>
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<td>Documented seroconversion, no. (%)</td>
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<td>Median viral load at sc [log(_{10})/ml] (IQR)</td>
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<td>Median CD4(^+) cell count at sc [n/(\mu l)] (IQR)</td>
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<td>Acute seroconversion, no. (%)</td>
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<td>Median viral load at sc [log(_{10})/ml] (IQR)</td>
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<td>Median CD4(^+) cell count at sc [n/(\mu l)] (IQR)</td>
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\( ^a \)TDR according to Bennett et al, 2009,
\( ^b \)sc: seroconversion,
\( ^c \)CI: 95% confidence intervals,
\( ^d \)IQR: interquartile ranges;
\( ^* \)statistically significant results \( p < 0.05 \);
\( ^* * \)highly statistically significant results \( p < 0.01 \);
\( ^p \)Mann-Whitney - U Test;
\( ^n \)Fisher exact test.

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Figure 1. Overall prevalence of TDR in the German HIV-1 Seroconverter Cohort 1997–2007. The prevalence of TDR (%) is plotted by year of seroconversion (Bennett et al. 2009; 12.4%; CI 10.4–14.3; \( p \) for trend over time 1996–2007 = 0.02) (fig. 2). In contrast, prevalence

![Figure 1. Overall prevalence of TDR in the German HIV-1 Seroconverter Cohort 1997–2007.](image)
of NNRTI resistance tended to increase over time ($p_{\text{for trend}} = 0.07$). Overall PI resistance remained stable among the study population genotyped over the time ($p_{\text{for trend}} = 0.24$; figure 2).

**Factors associated with transmitted drug resistance.** The prevalence of TDR was highest among patients exposed to HIV-1 through sex between men (12.5%, CI: 10.7–14.9); heterosexuals 10.3%; CI: 5.3–19.0; HPL 3.9%; CI: 0.7–18.9; and IVDU 4.8%; CI: 0.9–22.7). Men acquired TDR more frequently than women (12.5%, CI: 10.7–14.4; vs. 10.9%, CI: 5.4–20.9), respectively. The median age at seroconversion did not differ between patients with or without TDR (33.5; IQR 27.8–33.3 vs. 33.0; IQR 28.0–38.0; $p_{\text{MWT}} = 0.7$) (table 1). In a univariate analysis, there was no evidence to suggest, that TDR was associated with sex (OR = 1.15; CI: 0.5–2.6; $p_{\text{Fisher Exact}} = 0.9$). Being infected with subtype B virus was associated with a significantly higher risk to acquire resistant HIV strains (OR = 3.06; CI: 1.16–8.08; $p_{\text{Fisher Exact}} = 0.009$) and the exposure category unknown is significantly more frequently reported in patients with resistant HIV strains (OR = 2.43; CI: 1.07–5.51; $p_{\text{Fisher Exact}} = 0.05$; table 1).

Transmitted drug resistance is frequently caused by single resistance mutations. TDR was mostly related to viruses carrying single resistance mutations (figure S4). The majority of NNRTI resistance, 99.1% was caused by a singleton resistance mutation, followed by 68.4% singleton PI resistance mutations and 55.6% singleton NRTI resistance mutations. TAM mediated NRTI resistance was due in 54.4% to more than one TAM present in the same viral genome.

**Dominance of NRTI resistance mutations, TAMs and T215 revertant substitutions.** The majority of the NRTI resistant strains (79.8%) carried thymidine analogue resistance mutations (TAMs) selected by zidovudine and stavudine. Among the TAMs, revertant amino acid substitutions at position 215 of the reverse transcriptase were most prevalent (74.7%). Despite the significant decrease of transmitted NRTI resistance there is no clear cut decrease of either the prevalence of TAMs or the T215 revertant substitutions (figure S1). T215 revertant substitutions were by far the most prevalent resistance mutations (3.8%) in the study population (figure 3).

**Composition of antiretroviral first-line treatment in patients with susceptible and resistant HIV**

Analysis of treatment and treatment response was performed on the basis of resistance data according to the SDRM list of mutations. First-line therapy was initiated in 382/1,118 (34.2%) patients with susceptible strains and in 61/158 (38.6%) patients with TDR associated mutations.

First-line therapy of patients with susceptible genotype. First-line therapy was initiated in 382 patients with susceptible HIV strains and lasted for 182.0 days (median, IQR: 49.0–431.8). A Kaplan-Meier analysis revealed no significant difference in duration of first-line therapy comparing susceptible and resistant strains ($\chi^2_{\text{Log Rank}} = 0.004; p = 0.95$). Time between seroconversion and ART initiation was 327 days (median, IQR, 66.8–652.5) and Kaplan-Meier showed no significant difference in comparison to patients with resistant HIV strains ($\chi^2_{\text{Log Rank}} = 0.454; p = 0.50$). CD4 cell count measurements reported at the start of therapy were 335 cells/µl (median, IQR: 215.0–511.5) and plasma HIV-RNA measurement of 5.3 log 10/ml (median, IQR: 4.7–5.8). Any PI containing first line therapy was prescribed in 60.5% (231/382) of the patients with susceptible genotype. Predominantly, a 2 NRTI/1 PIr regimen was used as first-line therapy in 45.3% (173/382) of the patients. 34.3% (131/382) of individuals with susceptible strains received a NRTI/NNRTI first-line therapy. Any NNRTI comprising first-line therapy was initiated in 38.7% (146/382) of the patients. Efavirenz was preferred to nevirapine (92/148; 62.2% vs. 55/148; 37.2%, respectively).
First line therapy of patients with resistant genotype.

First line therapy was initiated in 61 patients (61/158) with TDR. Time between seroconversion and start of first-line therapy was 448 days (median, IQR: 162–660). The durability of the first-line regimen in this group was 144 days (median, IQR: 48–365). CD4 cell counts measured at time of antiretroviral treatment initiation were 312 cells/μl median and did not differ significantly to the susceptible group (IQR: 200–458; p MWT = 0.48). Also, plasma viral load was not significantly lower in patients with resistant strains compared to initial viral load measurements in the susceptible group (median, log 10/ml: 5.2; IQR: 4.6–5.5; pMWT = 0.36). Comparing VL between acute seroconverters with and without TDR, no significant difference was observed. First-line regimen containing any PI were used in 37/61 (60.7%) patients with resistant HIV. Any NNRTI was used in 24/61 (39.3%) patients in the TDR group. Efavirenz was again the preferred NNRTI compared to nevirapine in patients with TDR (16/24; 66.7% vs. 6/24; 25.0%).

Treatment success in patients with susceptible and resistant HIV

Viral suppression in patients with susceptible phenotype. Data about any viral load measurement during first year of first-line therapy were available for 271/382 (71%) of the patients with susceptible genotype. After applying the minimum duration of first-line therapy, e.g. 3 months, the patients included were 184/252 (73%). Consecutive viral load measurements within 5 to 12 months after treatment initiation were scarce. Data eligible for the analysis of treatment success were available only from 37/61 (60.7%) patients with resistant HIV. Any NNRTI was used in 24/61 (39.3%) patients in the TDR group. Efavirenz was again the preferred NNRTI compared to nevirapine in patients with TDR (16/24; 66.7% vs. 6/24; 25.0%).

Viral suppression in patients with resistant phenotype. Data about any viral load measurement data during first year of first-line therapy – taking the minimum duration of first line therapy into account - were available for 9/40 (22.5%) of patients with resistant HIV (table 2). Therapeutic success was achieved in the majority of treated patients (7/9, 77.8%). Two patients did not reach viraemic suppression (≤500 copies/ml) between 5–12 months after start of therapy.

Time trend in treatment response in patients with susceptible phenotype. Referring to the subpopulation entered into the analysis of treatment success, we performed an analysis on time trends in response to therapy solely in the group of patients infected with susceptible HIV strains (69 patients). We excluded the group of patients infected with resistant HIV due to small numbers (9 patients were entered into analysis on treatment success).

We used a logistic regression equivalent to the Cox-Armitage trend test. In this logistic regression analysis we applied as dependent variable binary coded treatment success and as covariant ‘Year of start of first line therapy’ (categories were: 1997–2000 = 0; 2001–2004 = 1; 2005–2007 = 2). The result indicates that with the higher the category the more likely to have treatment success ($b = 0.32; p = 0.12; OR = 1.37; CI: 0.93–2.04$). However note, that the result is not significant at the alpha = 0.05 level.

Discussion

In contrast to trends of TDR reported in other European cohorts [4,6,22,23,24,25,26], prevalence of TDR was stable at a high level during a decade of observation (1997–2007) in the German HIV-1 Seroconverter cohort [18,19]. Estimates of the prevalence of TDR may vary in cohorts because of differences in the study design, geographical location, definitions and classifications of TDR and composition of the study population. Prevalence of resistance is higher in patients with a known date of infection than in patients with an established but unknown date of HIV-infection. This was supported by lower prevalence of TDR in...
transmitted drug resistance

Table 2. Treatment success of patients with susceptible and resistant HIV 5–12 months after ART initiation.

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<tr>
<th>Characteristics of patients on first line therapy (FLT)</th>
<th>Predicted phenotype</th>
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<tr>
<td>Viral load (VL) data available within 12 months of FLT and &gt;1 VL report</td>
<td>Susceptible</td>
</tr>
<tr>
<td>Failure: 2 or more VL &gt;500 copies/ml</td>
<td>69/184 (37.5%)</td>
</tr>
<tr>
<td>Success: VL constant ≤500 copies/ml</td>
<td>2/69 (2.9%)</td>
</tr>
<tr>
<td>Any detectable viral load measurements: 1 VL &gt;500 copies/ml</td>
<td>62/69 (89.9%)</td>
</tr>
<tr>
<td>Any detectable viral load measurements: 5/69 (7.2%)</td>
<td>none</td>
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doi:10.1371/journal.pone.0012718.t002

Patients chronically infected with HIV as reported in a prospective multi centre cohort in the state of Nordrhein-Westfalia in Germany (9%; 95% CI 7.1–10.9) [27]. Reversion to wild type of resistance mutations affecting viral fitness from the dominant quasispecies over time is supposed to be the reason for the lower prevalence of TDR in chronically infected patients. As an example the mutations M184V and T215Y/F known to strongly reduce viral replication efficiency were never observed in chronically infected patients [30] in contrast to the population of newly infected HIV-1 seroconverters analysed in this study [M184V n = 10 and T215YF n = 10, respectively; figure 3].

The significant decrease of NRTI-related resistance over time (p = 0.02) in this population might have been caused by several factors. The reduction of the intensive use of mono- and dual therapies in previous years and the reduced replication capacity of viruses harbouring NRTI and multi drug resistance are associated with the reduced prevalence of NRTI resistance [28,29]. Additionally, improved treatment strategies, low tolerance of detectable viral load, and prompt management of treatment failure by the physicians might have lead to a reduction of NRTI resistance in treatment experienced persons subsequently influencing the spread of TDR in patients recently infected with HIV. Indeed, the decrease of NRTI resistance coincided with a decrease of the prevalence of the lamivudine selected mutation M184V and the TAMs T215YF, K219Q, K70R, in patients newly infected with HIV (figure S2). In contrast, some distinct NNRTI mutations (P225H, G190AS and Y181C) and also HIV strains carrying more than one NNRTI mutation were only observed since 2003/2004 and might contribute to the increase in NNRTI-resistance (figure S3), whereas the prevalence of the prominent NNRTI mutation K103NS (2.3%) is stable during the observation period (figure S3).

The overall prevalence of viruses with genotypic resistance in recently HIV-infected individuals did not vary over time [30], although the distribution of resistance between individual drug classes has changed as also shown in our study. The most common NRTI-resistance mutations detected in treatment-naive patients were M184V/I and TAMs, reflecting the extensive use of 3TC, ZDV and d4T and the use of mono and dual NRTI-therapy in the pre- and early HAART era. The T215 revertant was the most prevalent NRTI related mutation among patients with primary resistance [13,31]. These strains are phenotypically susceptible to zidovudine and stavudine in vitro but are able to acquire T215Y more rapidly than the wild type under the selective pressure of zidovudine [32–33]. Evidence exists that T215 revertants replicate as efficiently as the wild-type virus or even gained improved fitness and growth advantage compared to T215Y/F [34,35,36].

Many treatment options are available for the initial antiretroviral regimen of patients infected with HIV [37,38]. Reports describing the inferior treatment outcome in patients harbouring TDR, were frequently published [2,22,39]. If resistance testing is not possible or the patients needs to commence treatment before resistance results are available in case of low CD4 numbers, it may be advisable to initiate treatment with a boosted PI-regimen [40]. Taking into account that boosted PIs have a higher barrier to develop resistance, an accumulation of mutations is required to induce PI-related resistance. As observed in this cohort, the majority of the patients either carrying susceptible or resistant HIV strains, was treated with any PI comprising regimen as the first-line regimen. Mostly, a ritonavir boosted lopinavir/NRTI first-line regimen was preferred, reflecting the extended therapeutic options since PIs of the second generation were available for treatment.

It was previously assumed, that TDR detected at HIV-diagnosis would not impact the course of infection as fitter wild-type strains would become the dominant quasispecies. It is now well known, that TDR can persist for years as dominant quasispecies [41] and for even longer in a minority of the viral quasispecies in plasma (RNA) and as archived resistance within proviral DNA copies in the genome of peripheral blood mononuclear cells (PBMCs) [34] and other target cells of HIV. Therefore treatment response might be influenced by TDR even after many years, resulting in ongoing viral replication under selective pressure and promoting the evolution of further resistant variants [42]. First results in this study showed that treatment response was successful in patients with TDR bearing in mind that only small numbers of viral load measurements after start of therapy were available [10].

More than half of the patients in both groups (resistant 77.8%; susceptible 89.9%) were reported to be under detection limit within 5–12 months (≤500 copies/ml) after start of antiretroviral treatment. A long-term observation of the changes in the composition of individual antiretroviral regimen and monitoring of the treatment outcome is necessary to assess treatment success adequately. As reported in other studies, the presence of revertants, which were identified at a high prevalence in this study, seems to have a negative impact on virological response [6,35,43]. However, the results of this study underline the fact that most persons with TDR have good treatment outcome by using resistance testing to guide the choice of a first-line regimen [15,44].

Our findings reinforce that a broad and representative HIV resistance surveillance network between virologists, practitioners, clinicians and patients has to be installed to intensify the epidemiological knowledge about the transmission of resistant HIV, genotypic resistance testing frequencies, and treatment response in patients carrying resistant strain.

Supporting Information

Figure S1  Prevalence of TAMs and NRTI resistance mutations by year of seroconversion. The prevalences of TAMs and NRTI resistance mutations (SDRM list) were calculated by year of seroconversion. Found at: doi:10.1371/journal.pone.0012718.s001 (0.23 MB TIF)
**Figure S2** Prevalence of NRTI resistance mutations by year of seroconversion. The prevalences of each of TAMS and other NRTI resistance mutations (SDRM list) were calculated per year of seroconversion. A TAMs. Never observed: K219R. B NRTI resistance mutations found in other than TAMS. Never observed: K65R, T69s, L74V, V73AMTS, F116Y, Q138M.

Found at: doi:10.1371/journal.pone.0012718.s002 (0.88 MB TIF)

**Figure S3** Prevalence of NNRTI and PI resistance mutations by year of seroconversion. The prevalences of each of the NNRTI and PI resistance mutations (SDRM list) were calculated by year of seroconversion. A NNRTI resistance mutations. Never observed: L100I, K101E, V106M, Y181I, G190EQ, M230L (P236L not included). B PI resistance mutations. Occurred once: L24V, V32I, G48V, G73ACST.

Found at: doi:10.1371/journal.pone.0012718.s003 (0.73 MB TIF)

**Figure S4** Prevalence of TDR caused by one or more resistance mutations in the HIV genome by year of seroconversion.

Found at: doi:10.1371/journal.pone.0012718.s004 (0.51 MB TIF)

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**Author Contributions**

Conceived and designed the experiments: BB CK OH. Performed the experiments: KK SS. Analyzed the data: CH JW CK. Contributed reagents/materials/analysis tools: HJ SD. Wrote the paper: BB.


