Identification of the Critical Sites of NNRTI-Resistance in Reverse Transcriptase of HIV-1 CRF_BC Strains

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Abstract

Background: The polymorphisms involved in drug resistance to non-nucleoside reverse transcriptase inhibitors (NNRTIs) in HIV-1 CRF_BC, the most prevalent HIV-1 strain in China, have been poorly characterized.

Results: To reveal the drug resistance mutations, we compared the gene sequences of pol region of HIV-1 CRF_BC from 631 treatment-naïve and 363 treatment-experienced patients using the selection pressure-based method. We calculated an individual Ka/Ks value for each specific amino acid mutation. Result showed that eight polymorphic mutations (W88C, K101Q, I132L, R135L, T139K/R, H221Y and L228R) in RT for treatment-experienced patients were identified, while they, except for R135L, were completely absent in those from treatment-naïve patients. The I132L and T139K/R mutants exhibited high-level resistance to DLV and NVP and moderate resistance to TMC-125 and EFV, while the K101Q and H221Y mutants exhibited an increased resistance to all four NNRTIs tested. The W88C, R135L, and L228R may be RTI-induced adaptive mutations. Y181C+K101Q mutant showed a 2.5-, 4.4-, and 4.7-fold higher resistance to TMC-125, NVP and EFV, respectively, than Y181C alone mutant, while Y181C+H221Y or K103N+H221Y mutants had significantly higher resistance to all four NNRTIs than Y181C or K103N mutants. K103N+T139K and G190A+T139K mutant induce higher resistance (2.0–14.2-fold and 1.5–7.2-fold, respectively) to all four NNRTIs than K103N or G190A alone mutation.

Conclusions: I132L and T139K/R are rare but critical mutations associated with NNRTI-resistance for some NNRTIs. K101Q, H221Y and T139K can enhance K103N/Y181C/G190A-associated NNRTI-resistance. Monitoring these mutations will provide useful information for rational design of the NNRTI-based antiretroviral regimen for HIV-1 CRF_BC-infected patients.

Introduction

Human immunodeficiency virus type 1 (HIV-1) has been categorized into nine genetically distinct subtypes within the M group, including subtypes A, B, C, D, F, G, H, J, and K. Recombination between genomes of two viruses of different subtypes results in generation of a circulating recombinant form (CRF) [1]. The distribution of these subtypes and CRFs varies widely by region. HIV-1 CRF_BC recombinant that was derived from subtype B’ (Thailand B) and Indian subtype C lineages has resulted in epidemics among the injecting drug users (IDUs) in China since this recombinant was first reported in 1999 [2,3]. Currently, CRF_BC, which has been found in most parts of China, has become one of the most commonly transmitted HIV-1 subtypes across the country and was also found in other countries [4]. Rapid evolution and high mutation rate of HIV allow the virus to gain the ability of drug resistance. It is possible that HIV-1 genetic diversity may influence the type of resistance mutations that might eventually emerge upon drug exposure as well as the rate of emergence of resistance [5,6]. Most studies have focused on the mechanisms of drug resistance of the subtype B viruses, which comprise only about 12% of HIV-1 cases in the world [7]. The currently available reverse transcriptase inhibitors have been widely used in the world, including China, against both B and non-B HIV-1 strains; however, the polymorphisms involving in drug resistance to non-nucleoside reverse transcriptase inhibitors (NNRTIs) in HIV-1 CRF_BC pol region have been poorly characterized. Particularly, the mutation sites associated with NNRTI-resistance in RT of HIV-1 CRF_BC viruses have not been reported [6].

In the present study, we compared the gene sequence of pol region of HIV-1 CRF_BC isolated from treatment-naïve and experienced patients, and then conducted the selection pressure analysis to identify rare but critical sites of mutations potentially
Polymorphism analysis of characteristics of the study populations

Results

Characteristics of the study populations

This study involved 994 HIV-1-positive patients, including 631 treatment-naïve patients (female: 29.6%; heterosexual contacts: 8.1%; intravenous drug use: 26.5%; unknown: 65.1%) and 363 ART-treated patients (female: 26.2%; heterosexual contacts: 19.8%; intravenous drug use: 29.2%; unknown: 51.0%). All the patients were identified to be infected by HIV-1 CRF_BC as determined by Neighbor-joining genetic analysis of pol sequences of the viruses obtained from plasma samples of the HIV-1-infected patients using PCR technique. The ART-experienced patients were receiving highly active antiretroviral therapy, including 2 NRTIs and 1 NNRTI. The NRTIs are lamivudine(3TC) plus zidovudine(AZT) or stavudine(d4T), while the NNRTI is either nevirapine(NVP) or efavirenz(EFV). Specifically, 13.5% of the patients had been treated with 3TC/AZT/EFV, 58.7% with 3TC/AZT/NVP, 15.7% with 3TC/d4T/NVP, and 6.1% with unknown regimen. The mean treatment time was 18 months, including 28.0% for 0–6 months, 11.0% for 7–12 months, 23.1% for 13–18 months, 13.5% for 19–24 months, 17.9% for >24 months and 6.1% for unknown time.

Polymorphism analysis of pol gene region of HIV-1 CRF_Bc from plasmas of treatment-naïve and treatment-experienced patients

We used the selection pressure-based method, an important way to explore the rare but critical sites of drug resistance [10,14–16], to investigate the association of these mutations with the drug resistance based on the criteria: (1) the Ka/Ks (the ratio of the number of non-synonymous substitutions per non-synonymous site (Ka) to the number of synonymous substitutions per synonymous site (Ks) and LOD (log odds ratio) value (confidence score to evaluate the significance of mutation or mutation pair) of the mutation in treatment samples was greater than 1 and 2, respectively, and the Ka/Ks of the mutation in treatment samples were larger than that in treatment-naïve samples; (2) Frequency of mutations in treatment was significantly larger than that in treatment-naïve samples; (3) the non-synonymous mutations with low frequency (<1% treatment samples) were excluded. By evaluating the first 330 amino acids in HIV-1 RT sequences (the similarity of RT amino acids 1–330 between subtype B pNL4-3 and CRF_BC is 94.3%), we found that the frequencies of 15 polymorphism sites in RT of CRF_BC strains isolated from the treatment-experienced patients were significantly different from those isolated from the treatment-naïve patients (Table 1). In addition to the three previously reported RTI resistance-related mutations (A98G, Y181C, and K103N), 12 polymorphic mutations at seven positions (W88C, K101Q, I132L, T139K/R, H221Y, and L228R) were presented in RT of CRF_BC strains isolated from the treatment-experienced patients, while they were completely absent in the RT of CRF_BC strains isolated from ART-naïve patients. Several mutations, including R135L, V179D, Y181C, and M184V, present in the treatment-naïve patients who were infected by HIV-1 CRF_BC strains, but their frequencies were significantly increased in ART-treated patients (P<0.01), while R135L isn’t reported to be associated with drug resistance. In order to ascertain these polymorphism sites selected by NVP or EFV, the frequency of these mutations in patients with regimen containing NVP or EFV was compared. Of these mutations, A98G, T139R, and L228R were solely selected by NVP, and Y181C has significantly higher frequency in NVP group than EFV group (P=0.0012, fisher exact test).

Table 1. The changes of significant mutations between treatment-naïve and experienced patients.

| Mutationsa,b | Treatment-naïve patients | Treatment-experienced patients | P valuec |
|--------------|--------------------------|-------------------------------|----------|
|              | Frequency (n) Ka/Ks LOD  | Frequency (n) Ka/Ks LOD       |          |
| W88C         | 0.00% (0) - -            | 2.20% (8) 8.00 7.74          | <0.001   |
| A98G         | 0.00% (0) - -            | 1.93% (7) 26.82 6.38         | <0.001   |
| K101Q        | 0.00% (0) - -            | 1.65% (6) 4.38 2.64          | 0.001    |
| K103N        | 0.95% (6) 1.85 2.23      | 18.73% (68) 32.29 64.51      | <0.001   |
| I132L        | 0.00% (0) - -            | 1.65% (6) 6.00 6.02          | 0.001    |
| R135L        | 0.32% (2) 0.67 >2.00     | 3.31% (12) 12.00 >2.00       | <0.001   |
| T139K        | 0.00% (0) - -            | 2.20% (8) 13.14 6.62         | <0.001   |
| T139R        | 0.00% (0) - -            | 1.10% (4) 6.57 2.33          | 0.008    |
| V179D        | 1.43% (9) 1.00 1.84      | 4.68% (17) 3.76 9.45         | 0.002    |
| Y181C        | 0.63% (4) 0.18 0.02      | 10.47% (38) 4.22 13.37       | <0.001   |
| M184V        | 0.16% (1) 1.00 0.31      | 20.39% (74) 74.00 37.00       | <0.001   |
| V188L        | 0.00% (0) - -            | 1.38% (5) 1.25 >2.00         | 0.003    |
| G190A        | 0.00% (0) - -            | 7.16% (26) 11.07 24.78        | <0.001   |
| H221Y        | 0.00% (0) - -            | 6.61% (24) 2.40 6.65         | <0.001   |
| L228R        | 0.00% (0) - -            | 1.65% (6) 3.00 3.71          | 0.001    |

Note:

aThe bold mutations are those that have not been reported to be associated with drug resistance.

bThe reference strain is CRF_Bc.CN.CN54. The mutations listed denote the reference amino acid from HIV-1 subtype CRF07_BC.
Susceptibility to NNRTIs against HIV-1 CRF_Bc strains with the newly identified mutations in RT

To investigate the contribution of these mutations to NNRTI resistance, the sensitivities of the viruses with WT and MT in RT to each NNRTI used, including Etravirine (TMC-125), Rescriptor (DLV), Viramune (NVP), and Sustiva (EFV), were determined. CRF_Bc strains with K103N and Y181C in RT were included as controls. The K101Q, I132L, and T139K/R mutants exhibited significant (2–28-fold) increases in resistance to all the four NNRTIs tested (P<0.05), and H221Y mutant had a moderate increase (approximately 2-fold) of resistance to these four NNRTIs (P<0.05), while the W88C, R135L and L228R mutations had no significant effect on the viral resistance to RTIs (Table 2). Besides K101Q and H221Y, the other three mutants I132L and T139K/R were rarely reported to associate with drug resistance. We found that HIV-1 subtype B viruses with I132L and T139K/R mutations were also resistant to NNRTIs, although their resistant level is relatively lower than that of HIV-1 CRF_Bc viruses with these mutations (Table 2).

Characterizing the mutation relationship based on predicted drug resistance mutations interaction network

To determine the influence of the mutation of one site to another, a conditional selection ratio was computed. If the conditional selection ratio of X to Y (X→Y) is greater than 1 and LOD is greater than 2, the influence of X to Y (X→Y) was considered significant. Then, software Cytoscape was used to construct the relationship among these predicted drug resistance mutations as reported [10].

The network represents the comprehensive relationship among the predicted drug-resistance mutations, and the arrows from the source node to the target node indicate the influence of one to another. In the network, the size of the node represents the mutation frequency of that site from one amino acid to another. In the network, the width of line represents the influence strength between two mutations. As shown in Figure 1, the network contained 15 mutation sites which have 40 interaction relationships (Table S1). In the network, mutations with higher frequency, such as M184V and K103N, were more likely to influence the other mutations. For example, M184V and K103N had 12 (A98G, K101Q, K103N, I132L, R135L, T139K, T139R, Y181C, Y188L, G190A, H221Y, and L228R) and 6 (R135L, T139K, T139R, Y181C, H221Y, L228R) target mutations, respectively.

The table below shows the fold changes in EC50 values of different mutations in relation to WT virus for each NNRTI.

### Table 2: Sensitivity and resistance of different mutation sites in HIV-1 CRF_Bc pol region to NNRTIs using an in vitro phenotypic assay.

| Mutations | TMC-125 | DLV | NVP | EFV |
|-----------|---------|-----|-----|-----|
|           | EC50a (nM) | Fold changeb | EC50 (µM) | Fold change | EC50 (µM) | Fold change | EC50 (nM) | Fold change |
| WT (BC)   | 1.03 ± 0.10 | - | 0.09 ± 0.00 | - | 0.10 ± 0.01 | - | 1.713 ± 0.180 | - |
| W88C (BC) | 1.09 ± 0.02 | 1.06 | 0.05 ± 0.00 | 0.51 | 0.09 ± 0.01 | 0.89 | 0.98 ± 0.06 | 0.57 |
| K101Q (BC) | 3.61 ± 0.35 | 3.50 | 0.15 ± 0.01 | 1.71 | 1.30 ± 0.14 | 12.57 | 5.25 ± 1.80 | 3.06 |
| K103N (BC) | 1.400 ± 0.06 | 1.35 | 7.35 ± 0.23 | 82.56 | 17.23 ± 1.36 | 167.23 | 97.48 ± 4.06 | 56.91 |
| I132L (BC) | 5.21 ± 0.78 | 2.55 | 1.72 ± 0.12 | 19.35 | 4.30 ± 0.26 | 28.05 | 10.52 ± 1.22 | 6.13 |
| R135L (BC) | 1.36 ± 0.15 | 1.32 | 0.07 ± 0.00 | 0.75 | 0.12 ± 0.00 | 1.13 | 1.87 ± 0.13 | 1.09 |
| T139R (BC) | 3.73 ± 0.62 | 1.82 | 0.42 ± 0.02 | 4.69 | 2.59 ± 0.53 | 25.12 | 3.23 ± 0.38 | 1.89 |
| T139K (BC) | 4.82 ± 0.96 | 4.67 | 0.33 ± 0.02 | 3.66 | 0.76 ± 0.02 | 7.35 | 5.65 ± 0.12 | 3.30 |
| M184V (BC) | 0.69 ± 0.04 | 0.67 | 0.03 ± 0.00 | 0.31 | 0.05 ± 0.01 | 0.44 | 0.70 ± 0.05 | 0.41 |
| Y181C (BC) | 5.91 ± 1.70 | 5.73 | 4.53 ± 0.13 | 50.92 | 15.84 ± 1.42 | 153.74 | 3.22 ± 0.14 | 1.88 |
| H221Y (BC) | 1.42 ± 0.16 | 1.38 | 0.18 ± 0.01 | 2.04 | 0.21 ± 0.03 | 2.02 | 3.02 ± 0.51 | 1.76 |
| L228R (BC) | 1.02 ± 0.10 | 0.99 | 0.04 ± 0.001 | 0.47 | 0.11 ± 0.00 | 1.03 | 0.79 ± 0.04 | 0.46 |
| WT (B)    | 1.66 ± 0.35 | - | 0.11 ± 0.00 | - | 0.19 ± 0.01 | - | 3.26 ± 0.25 | - |
| I132L (B) | 6.51 ± 1.19 | 3.91 | 0.51 ± 0.02 | 4.61 | 1.20 ± 0.03 | 6.38 | 11.61 ± 0.19 | 3.56 |
| T139K (B) | 5.21 ± 1.03 | 3.13 | 0.200 ± 0.01 | 1.78 | 0.43 ± 0.03 | 2.26 | 6.87 ± 0.79 | 2.10 |
| T139R (B) | 9.64 ± 0.87 | 5.79 | 0.44 ± 0.01 | 3.99 | 1.09 ± 0.05 | 5.78 | 8.97 ± 0.63 | 2.75 |

Note:
- The unit for the EC50 values of TMC-125 and EFV is nM, while that for the EC50 values of DLV and NVP is µM. Data were presented as the mean ± standard deviations of three separate determinations.
- WT: HIV-1 CRF_Bc strain N-14-1.
- Fold change was determined by calculating the ratio of EC50s for mutations and WT viruses.

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Critical Sites of NNRTI-Resistance in HIV-1 CRF_Bc
Susceptibility to NNRTIs of HIV-1 CRF_BC strains with the newly identified in combination with the well-known Y181C, G190A or K103N mutation

We next examined the effect of the single mutation sites listed above in combination with Y181C or K103N on viral resistance to NNRTIs. The pNL4-3 clone containing HIV-1CRF_BC pol with mutations Y181C, G190A or K103N was constructed through site-directed mutagenesis with or without the newly identified mutations in this study. We tested the phenotypic resistance of these combinations Y181C, G190A or K103N with different mutation sites in RT of HIV-1 CRF_BC to NNRTIs using an in vitro phenotypic assay. As shown in Table 3, Y181C+K101Q mutant showed a 2.48-, 4.37-, and 4.69-fold higher resistance to TMC-125, NVP and EFV, respectively, than Y181C alone mutant (P<0.05). Y181C+H221Y mutations resulted in signifi-
cantly higher resistance to all four NNRTIs than Y181C alone mutation, ranging in 3.00–4.24 FC (P<0.05). K103N+T139K mutant induce higher resistance to all four NNRTIs, with FC ranging from 2.00 to 14.15. K103N+H221Y mutations exhibited an increased (1.69- to 2.96-fold) resistance to the four NNRTIs tested (P<0.05), while K103N+K101Q mutants did not displayed a higher NNRTI-resistance than K103N alone mutant. G190A+T139K also showed a higher increased (1.48- to 7.21-fold) resistance to all four NNRTIs than G190A alone mutation.

**Discussion**

Most of the current anti-HIV drugs have not been tested in the clinical trials in China, drawing attention to the effectiveness of these drugs against the HIV-1 strains circulation in China. We recently have shown that Fuzeon and Maraviroc, the only two HIV entry inhibitors approved for clinical use by the US FDA, are much less effective against the HIV-1 subtypes circulating in China than the B subtype predominating in the United States and Europe [5]. Therefore, it is essential to study the effectiveness of a new class of antiretroviral drugs, such as NNRTIs, before they are introduced into China.

At present, the antiretroviral drugs have been used not only for treatment, but also for prevention of HIV infection/AIDS [19]. HIV clinical trials revealed the magnitude of benefit when using antiretroviral drugs to prevent sexual transmission or mother-to-child transmission of HIV-1 [20,21], suggesting the new use of antiretroviral drugs for pre- and post-exposure prophylaxis [22]. Therefore, analysis of the drug-resistance becomes more and more important for rational design of therapeutic and prophylactic regimen.

Some in vitro and in vivo observations suggest that the various subtypes may respond differently to NNRTIs [23]. The frequency and pattern of mutations conferring resistance to these drugs differ among HIV-1 subtypes and can influence the outcome [24]. CRF_BC strain accounted for more than half of HIV-1 infection among HIV-1 subtypes and can influence the outcome [24]. The frequency of NNRTI resistance of CRF_BC strain was significantly different from those isolated from the treatment-naïve patients. Especially, seven mutations at six positions (W88C, K101Q, I132L, R135L, T139K/R, H221Y and L228R) were completely absent in the RT of CRF_BC strains isolated from drug-naïve patients. In contrast, their frequencies in strains isolated from ART-treated patients were significantly increased, suggesting their specific association with ART treatment. Since the ART regimen of these patients contained two NRTIs and one NNRTI, in vitro experiments were tested for susceptibility to 3TC, d4T, AZT, TFV. The results demonstrated that these mutations were not associated with the resistance to NRTIs (Table S2). We postulate that these mutations may have effect on their sensitivity to NNRTIs. Five mutants (K101Q, I132L, T139K/R and H221Y) among these eight mutants exhibited an increased resistance to the four NNRTIs tested. According to Stanford HIV resistance database, the mutations of I132L, T139K and T139R were rare events (0.11%, 0.57% and 4%, respectively) in B subtype under treatment, which may indicate the higher genetic barrier for these three mutations in B subtype than CRF_BC. Although it is reported that K101Q and H221Y may belong to the ETR RAMs [26], and H221Y was a mutation responsible for drug-resistance to Rilpivirine [27], our study has shown for the first time that both K101Q and H221Y mutations are associated with the increased resistance to all the four NNRTIs tested. Our study has demonstrated that the viruses with I132L and T139K/R mutations that exhibited high-level resistance to NNRTIs are

| Mutations       | TMC-125 | DLV | NVP | EFV |
|-----------------|---------|-----|-----|-----|
|                 | EC50a (nM) | Fold change | EC50 (µM) | Fold change | EC50 (µM) | Fold change | EC50 (µM) | Fold change |
| Y181C           | 5.91±1.70 | - | 4.53±0.13 | - | 13.95±0.44 | - | 3.22±0.14 | - |
| Y181C+K101Q     | 14.67±0.36 | 2.48 | 7.26±0.24 | 1.60 | 60.95±2.42 | 4.37 | 15.11±0.18 | 4.69 |
| Y181C+H221Y     | 19.19±0.29 | 3.25 | 13.60±0.82 | 3.00 | 59.17±8.93 | 4.24 | 11.76±1.67 | 3.65 |
| Y181C+L228R     | 6.34±0.92 | 1.07 | 3.50±0.22 | 0.77 | 26.35±1.88 | 1.89 | 3.37±0.55 | 1.05 |
| K103N           | 1.40±0.06 | - | 7.35±0.23 | - | 17.23±1.36 | - | 97.48±4.06 | - |
| K103N+K101Q     | 2.14±0.16 | 1.53 | 7.07±0.48 | 0.96 | 25.20±0.52 | 1.46 | 124.19±14.47 | 1.28 |
| K103N+H221Y     | 3.96±0.46 | 2.83 | 21.75±0.84 | 2.96 | 25.85±0.49 | 1.79 | 164.41±9.02 | 1.69 |
| K103N+L228R     | 1.78±0.18 | 1.28 | 5.23±0.16 | 0.71 | 17.60±0.97 | 1.02 | 153.94±1.10 | 1.59 |
| K103N+T139K     | 2.80±0.14 | 2.00 | >100.00 | >14.15 | 82.08±8.20 | 4.77 | 408.95±52.69 | 4.20 |
| G190A           | 1.81±0.53 | - | 0.02±0.01 | - | 6.11±0.13 | - | 8.31±0.72 | - |
| G190A+T139K     | 2.92±0.44 | 1.61 | 0.03±0.00 | 1.48 | 44.01±1.70 | 7.21 | 15.68±1.45 | 1.87 |

**Note:**
- The unit for the EC50 values of TMC-125 is nM, while that for the EC50 values of DLV, NVP and EFV is µM. Data were presented as the mean ± standard deviations of three separate determinations.
- Fold change was determined by calculating the ratio of EC50s for mutations and WT viruses.
- Each sample was tested in triplicate, and each experiment was repeated twice. EC50 (mM, except nM for maraviroc) data are presented as means ± standard deviations.
- For NNRTIs, fold change was calculated by measuring the effect of mutations on drug-resistance.
- For NRTIs, fold change was calculated by measuring the effect of mutations on drug-sensitivity.
- The EC50 values for TMC-125, DLV, NVP and EFV were obtained from three independent experiments performed in triplicate.
Critical Sites of NNRTI-Resistance in HIV-1 CRF_BC

the rare but critical mutants associated with NNRTI-resistance in both CRF_BC and B subtype.

The potential mechanistic association between the NNRTI-resistance and the I132L and T139K/R mutations may be ascribed to the location of these mutation sites. All of the three mutations are located in the ββ/β0 loop (residues 132–140) of RT, which is involved in the formation of the base of the NNRTI-binding pocket [28,29]. Mutations of these residues may cause the conformation change of the pocket, resulting in the decreased binding between the NNRTI and the pocket in RT. It was also reported that T139K mutation could seriously impair catalytic activities of RT [30].

The increasing evidences suggest that in addition to those currently known mutations, more and more unidentified mutations may also be involved in the development of NNRTI resistance, which contribute to NNRTI therapy failure [6], and the development of resistance to NNRTIs may be more complex than the classical one-step model of significant resistance via a single mutation so far considered [31]. It has been reported that HIV can employ various combinations of mutations to resist drug treatments [32]. To further determine mutational interactions between the newly identified and unknown mutations in RT of CRF_BC strains, a conditional selection ratio were computed. We found that all mutations were connected together as a component and in the network, mutations of high frequency were more likely to influence the other mutations (Fig. 1). The relationship among mutations in the networks can give clues to the combinatorial mutation patterns responsible for HIV drug resistance within the network. Particularly, H221Y were associated with Y181C and/or K103N mutations in RT of CRF_BC strains isolated from the treatment-experienced patients and K101Q showed positive interaction with M184V. Others have also reported similar combinational mutations, although the effect of these combined mutations on drug-resistance has not been clearly defined [6,33].

To understand the effect of our newly identified mutations combined with those known mutations, we examined the effect of the single mutation sites in combination with Y181C, G190A or K103N on viral resistance to NNRTIs. The result showed that either Y181C+H221Y or K103N+H221Y mutants exhibited significantly enhanced resistance to all the four NNRTIs tested, compared with Y181C alone and K103N alone mutants. Y181C+K101Q mutants also showed higher resistance to TMC-125, NVP and EFV than Y181C alone mutant. K103N+T139K and G190A+T139K mutants increase an increased resistance to all four NNRTIs. These results suggest that K101Q, T139K and H221Y are able to enhance the NNRTI-resistance mediated by those well-characterized HIV-1 mutants. The positive interaction between K101Q and M184V is of interest and will be investigated in vitro in future time.

In summary, our data suggest that I132L and T139K/R mutations that exhibited high-level resistance to NNRTIs are the rare but critical mutants associated with NNRTI-resistance in RT of CRF_BC strains that are predominantly circulating in China, while K101Q and H221Y mutations are associated with the increased resistance to all the four NNRTIs tested, although at codons 101 and 221 were reported relating to NNRTI resistance. The co-presence of H221Y, T139K or K101Q with the well-known RTI-resistance mutations K103N, G190A or Y181C may strengthen the drug-resistance effect. Further study is needed to determine how these mutations and combined mutations affect the binding kinetics of NNRTIs. We suggest that these newly identified mutations should be considered for the improvement of algorithms that predict clinical responses to antiretroviral drugs and for assessing the efficacies of next-generation drugs. This information will aid in designing initial treatment strategies for persons infected with CRF_BC viruses and interpreting genetic resistance among the CRF_BC-infected patients whose antiretroviral therapy has failed.

Methods

Study population

The study population included pre-selected HIV-1-positive patients with treatment-naive and experienced antiretroviral therapies, who participated in a multicenter AIDS Cohort Study including China Global Fund AIDS Program, and “Eleven Five” major projects in Xinjiang and Sichuan provinces of China during 2007–2011. The individuals who newly HIV-infection screened and confirmed were investigated without experiencing ART were chosen as the treatment-naive patients in Xinjiang and Sichuan province of China during that time. The HIV/AIDS patients who received ART with 2 NRTIs and 1 NNRTI regimen in the two provinces were investigated to detect viral load and CD4 count periodically. When the patients encountered virological failure during ART according to WHO ARV therapy failure criteria (the virological failure was defined as a viral load of ≥10 000 copies/ml) [8], they were recruited as the treatment-experienced patients. To obtain the CRF_BC recombinant representative isolates, 994 patients were chosen through sequence blastx on the website (http://www.hiv.lanl.gov/content/sequence/BASIC_BLAST/basic_blast.html). Furthermore, to confirm these sequences, we conducted a Neighbor-joining genetic analysis of pol sequences obtained from plasma samples of all HIV-1-infected patients using the PCR technique as previously described [9]. This study was approved by the Institutional Research Ethics Community, China CDC, and all subjects signed informed consent forms before blood collection.

HIV-1 pol sequence detection

HIV pol sequence was carried out by an in-house polymerase chain reaction protocol as previously described [9] Briefly, viral RNA was extracted from patient’s plasma using a QIApp Viral RNA Mini Kit (Qiagen Inc., Chatsworth, CA) and cDNA was generated using primer RT21 (CTGTATTTCTGCT-CAAGTCTTTTATGGG). A nested PCR was then employed using the generated cDNA as template. The nested PCR product was purified using a QIAquick Gel Extraction Kit (Qiagen Inc) and sequenced with the ABI 3100 DNA Sequencer.

Ka/Ks and Conditional selection ratio calculation

The Ka/Ks values for specific amino acid substitutions were determined as described by Chen et al [10]. To measure how a specific amino acid of one site X influences one in the other site Y. The ‘conditional selection ratio’ is defined as the ratio of Ka/Ks of Y when the amino acid is mutated at X (Ka/Ks)1/Y/Δx divided by the Ka/Ks of Y in the absence of any mutation at X (Ka/Ks)Y/Δx, and it was computed as follows:

\[ \left( \frac{K_a}{K_s} \right)_{Y/X} = \frac{\left( \frac{K_a}{K_s} \right)_{Y/\Delta x}}{\left( \frac{K_a}{K_s} \right)_{Y/\Delta x}} = \frac{N_{YXa}}{N_{YXo}} \]

Where NyXa is the number of samples with the same amino acid mutation both at site Y and X; and NyXo is the number of samples with a synonymous mutation at codon Y and an amino acid mutation at codon X. NyXa and NyXo are the number of samples with the amino acid mutation and a synonymous
Critical Sites of NNRTI-Resistance in HIV-1 CRF_BC

Phenotypic assay to HIV-1 NNRTIs based on TZM-bl cells

HIV-1 (HIV-1WT) and HIV-1 with the mutations (HIV-1MT) were generated by transfection of the plasmids into 293T/17 cells by using Fugene 6 Transfection Reagent (Roche Applied Science) according to the manufacturer’s instructions. The 50% tissue culture infectious dose (TCID₅₀) and the antiviral activity of NNRTIs were determined using TZM-bl cells as previously described [12,13]. The concentration of drug that effects 50% viral replication (EC₅₀) values was determined by nonlinear regression using GraphPad Prism 5.01. Mean EC₅₀ were calculated using all replicates for each virus and are expressed as mean ± SD. The Wilcoxon rank sum test was applied to pairwise comparisons to determine whether the observed differences between EC₅₀ for different site-mutations were statistically significant.

Supporting Information

Table S1 The Conditional selection ratio among drug resistance related mutations.

(DOC)

Table S2 Sensitivity and resistance of different mutation sites in HIV-1 CRF_BC RT region to NRTIs.

(DOC)

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Author Contributions

Conceived and designed the experiments: LM, YS. Performed the experiments: YH, ZL, YJ, YO, RA. Analyzed the data: LM, YH, ZL. Contributed reagents/materials/analysis tools: YH, ZL, YJ, YO, RA, LL. Wrote the manuscript: LM, YH, SJ, YS.

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