Prevalence of mutations and determinants of genotypic resistance to etravirine (TMC125) in a large Italian resistance database (ARCA)*†

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Objectives
To evaluate whether etravirine (TMC125) might be effective in patients failing therapy with current nonnucleoside reverse transcriptase inhibitors (NNRTIs), we analysed the prevalence of TMC125 mutations and the possible determinants of genotypic resistance to this drug among sequences reported to a large database in Italy [Antiretroviral Resistance Cohort Analysis (ARCA)].

Methods
We analysed the prevalence of TMC125 resistance-associated mutations (RAMs) and the TMC125 weighted genotypic score (WGS) together with the determinants of genotypic resistance. A total of 5011 sequences from 2955 patients failing NNRTI therapy were evaluated.

Results
Among the sequences in ARCA, 68% had at least one and 9.8% at least three TMC125 RAMs, whereas 31% had a WGS > 2. Frequent RAMs were Y181C, G190A, K101E and A98G, whereas V179F, Y181V and G190S appeared in < 5% of sequences. Multivariate analysis revealed a higher risk of developing at least three TMC125 RAMs associated with both nevirapine and efavirenz exposure, whereas CD4 counts ≥ 200 cells/µL retained their protective effect. An increased risk of WGS > 2 was linked to higher HIV RNA values (maximum risk at > 5 log10 copies/mL) and nevirapine exposure; CD4 counts ≥ 200 cells/µL were protective.

Conclusions
The prevalence of TMC125 resistance mutations in the ARCA cohort was 68%. The DUET studies showed that at least three TMC125-associated mutations were required to impair the efficacy of the drug and Y181C/V, V179F and G190S had the greatest effect on response. The prevalence of these mutations among the patients examined in our study was low. However, WGS > 2 was found for one-third of our sequences. Previous nevirapine exposure was associated with an increased risk of having WGS > 2 (adjusted odds ratio 1.76).

Keywords: ARCA, drug resistance, genotype, HIV-1, TMC125

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Introduction
HIV-infected patients who experience triple-drug class virological failure may be at increased risk of clinical progression and death [1]. Therefore, newer agents with activity against drug-resistant HIV-1 are needed [2].
Etravirine (TMC125) is a second-generation nonnucleoside reverse transcriptase inhibitor (NNRTI) with potent activity against wild-type and NNRTI-resistant HIV-1 and a high genetic barrier to the development of resistance [3,4]; its potent antiretroviral activity has been demonstrated in the phase III DUET studies [5,6].

Information on TMC125 resistance is still scarce: a set of 13 baseline reverse transcriptase mutations was previously identified in the DUET studies as having an effect on virological response to TMC125 [7–12]. Poveda et al. [12] suggested that efavirenz (EFV) might be less capable of inducing TMC125 resistance than nevirapine (NVP). Moreover, a longer duration of initial NNRTI treatment has been associated with increased evidence of in vitro TMC125 resistance [13] and the inclusion of NVP within the initial highly active antiretroviral therapy (HAART) regimen could result in a higher risk of virological failure and drug resistance compared with EFV [14]. This could limit the future use of TMC125 [15].

Recently, Tambuyzer et al. examined two TMC125 weighted genotypic scores (WGSs) [TBT (Tibotec, Mechelen, Belgium) and MGR (Monogram, San Francisco, CA, USA)], which produced similar results in defining susceptibility to TMC125 in treatment-experienced patients and were able to predict nonresponse to TMC125 in ~60% of subjects enrolled in the DUET trials [16]. Nevertheless, there is a difference between mutations associated with TMC125 use (L100I, E138G, V179F/I, Y181C/I and H221Y), i.e. mutations that emerge with use of TMC125, and mutations associated with an altered response to TMC125 (V90I, A98G, L100I, K101E/H/P, V106I, E138A, V179D/T/F, Y181C/I and G190A/S).

To evaluate whether efavirenz might be effective in patients failing therapy with current NNRTIs, we analysed the prevalence of efavirenz mutations and possible determinants of genotypic resistance to this drug among sequences reported to a large Italian database.

Materials and methods

We retrospectively considered HIV-1 reverse transcriptase sequences obtained from the Italian Antiretroviral Resistance Cohort Analysis (ARCA; available at www.hivarca.net) database for a total of 2955 patients experiencing therapy failure with an NNRTI-based regimen at the time of blood drawing, and with complete treatment history available. These subjects had been selected on the basis of having a resistance test while failing their antiretroviral regimen (viral load > 1000 HIV-1 RNA copies/mL). Patients were TMC125-naïve. Inclusion criteria were NNRTI-based regimen for at least 3 months, and an HIV RNA measurement and CD4 cell count available within 1 month.

Drug resistance mutations were interpreted following the latest International AIDS Society–USA (IAS–USA) panel list of mutations proposed to be TMC125-specific (www.iasusa.org; update in December 2008) [17]: V90I, A98G, L100I, K101E, K101P, K101H, V106I, E138A, V179D, V179F, V179T, Y181C, Y181I, Y181V, G190A, G190S and M230L. Given the high prevalence of mutation K103N among isolates resistant to other NNRTIs, we also added this mutation to our analysis.

The risk of developing at least three TMC125 resistance-associated mutations (RAMs) [7,8] was assessed by means of binary logistic regressions models. Univariate and multivariate analyses were performed to estimate crude and adjusted relative risks (odds ratios, 95% confidence intervals and Wald statistic) for gender, age, HIV RNA, CD4 cell count; and NVP, EFV, protease inhibitor (PI) and enfuvirtide (T20) exposure. Moreover, we considered the number of NNRTIs received and the duration of NNRTI therapy. The level of statistical significance was set at $P = 0.05$. SPSS 15 for Windows was the statistical software package used for the analyses (SPSS, Chicago, IL, USA). Moreover, we conducted our analysis with the endpoint of having a TBT WGS > 2, which has been reported to predict poor virological response to TMC125 in treatment-experienced patients [16].

Results

A total of 5011 sequences obtained from 2955 patients were evaluated. Of these, 1241 subjects (42.0%) were exposed only to NVP, 1053 (35.6%) only to EFV, and 613 (20.7%) to both NVP and EFV.

Of these 2955 patients, 2153 (72.9%) presented with at least one TMC125 RAM. Among the sequences in ARCA, 68% had at least one and 9.8% at least three TMC125 RAMs, whereas 31% showed a WGS > 2.

Among the samples with at least one RAM for TMC125 ($n = 3407$), the mutations most frequently represented were Y181C (27%), G190A (22.8%), K101E (11.7%) and A98G (9.3%). K103N was present in 53.9% of sequences. V179F, Y181V and G190S were present in 0.3%, 1.0% and 4.9% of sequences, respectively. When at least three TMC125-related mutations were found ($n = 495$), the mutations most frequently represented were G190A (62%), Y181C (57.6%) and K101E (44%). K103N was present in 44.8% of sequences. V179F, Y181V and G190S were present in 1.4%, 1.0% and 13.5% of sequences, respectively (Fig. 1). Among the samples with TBT WGS > 2 ($n = 1553$), the most frequent mutations were Y181C (59.3%), G190A (26.7%) and K101E (17.8%); K103N was found in 40.1% of sequences.
We also analysed the association between TMC125 RAMs and exposure to NVP and EFV: these mutations appeared more frequently in NVP- than EFV-treated patients: 90.2% of sequences from patients exposed to NVP vs. 35.2% of sequences from patients exposed to EFV had mutation Y181C, and these percentages were, respectively, 84.7% vs. 43.1% for G190A, 72.7% vs. 49.8% for K101E, 100% vs. 23.5% for Y181V, and 66.7% vs. 33.3% for V179F. G190S appeared more frequently with exposure to EFV (81.4% of sequences) than NVP (43.3% of sequences).

Multivariate analysis revealed that male gender and being EFV- or NVP-experienced were associated with statistically significant increases in the risk of developing three or more TMC125 RAMs. CD4 values ≥200 cells/µL and greater age (for each additional 10 years) were statistically protective factors, whereas PI and T20 experience and HIV RNA values did not show any statistically significant associations. Considering the duration of NNRTI treatment, every additional year of therapy represented a slightly increased risk of developing at least three TMC125 RAMs, although this was not statistically significant; also, being exposed to two NNRTIs, but not to one, produced an adjusted odds ratio (AOR) of 1.93 (95% CI 0.74–5.07).

We then focused our attention on the risk of having a TBT WGS ≥2. As shown in Table 1, some differences were found in comparison to the analysis of at least three TMC125 RAMs. Of interest, a strong predictor of a decreased phenotypic susceptibility to TMC125 was a higher HIV RNA value (maximum risk at >5 log10 copies/mL), with the AOR increasing from 2.62 for HIV RNA (≤3.7 log10 copies/mL; 95% CI 1.35–5.10; \( P = 0.004 \)) to 3.99 for HIV RNA (≥5 log10 copies/mL; 95% CI 1.98–8.04; \( P < 0.001 \)). NVP exposure retained an increased risk of a TBT WGS ≥2 (AOR 1.76; 95% CI 1.42–2.18; \( P < 0.001 \)), whereas previous EFV treatment did not. Duration of NNRTI therapy and previous exposure to one NNRTI did not have any significant effect, whereas exposure to two NNRTIs still had a significant effect, with an AOR of 2.26 (95% CI 1.05–4.88; \( P = 0.038 \)).

**Conclusions**

The prevalence of TMC125-related mutations in the ARCA cohort was 68%. According to the DUET studies [7,8],

![Fig. 1](image-url) Drug resistance mutations were interpreted following the latest International AIDS Society-USA (IAS-USA) panel list of mutations proposed to be etravirine (TMC125)-specific (www.iasusa.org; update in December 2008) [17]. There were 3407 sequences with at least one TMC125 resistance-associated mutation (RAM) and 495 sequences with at least three TMC125 RAMs. Mutation K103N was pictured as reference NNRTIs resistance mutation.

| Table 1 | Determinants of genotypic resistance [Tibotec (TBT) score 0–2 or >2] to TMC125 |
| --- | --- | --- | --- | --- | --- |
| Category | TBT score 0–2 | TBT score >2 | AOR | 95% CI | P |
| Age (years) | 42.8 (42) | 41.4 (40) | 0.80 | (0.72–0.88) | <0.001 |
| Sex | | | | | |
| Female | 504 (65.6) | 264 (34.4) | 1 |
| Male | 1168 (62.2) | 698 (37.4) | 1.26 | (1.05–1.51) | 0.015 |
| CD4 count | | | | | |
| <200 cells/µL* | 298 (56.4) | 204 (33.3) | 0.70 | (0.54–0.90) | 0.005 |
| 200–399 cells/µL | 408 (66.7) | 204 (33.3) | 0.70 | (0.54–0.90) | 0.005 |
| ≥400 cells/µL | 450 (71.4) | 180 (28.6) | 0.57 | (0.44–0.74) | <0.001 |
| ND | 516 (59.7) | 348 (40.3) | 0.99 | (0.76–1.29) | 0.964 |
| HIV RNA | | | | | |
| <0.7 log10 copies/mL* | 69 (86.3) | 11 (13.8) | 1 |
| 0.7–3.7 log10 copies/mL | 476 (68.8) | 206 (30.2) | 2.62 | (1.35–5.10) | 0.004 |
| 3.7–5 log10 copies/mL | 585 (60.5) | 382 (39.5) | 3.67 | (1.90–7.10) | <0.001 |
| >5 log10 copies/mL | 138 (54.5) | 75 (45.5) | 3.99 | (1.98–8.04) | <0.001 |
| ND | 404 (62.2) | 248 (38.0) | 2.92 | (1.49–5.71) | 0.002 |
| EFV-experienced | 1008 (66.7) | 504 (33.3) | 0.96 | (0.78–1.17) | 0.665 |
| NVP-experienced | 974 (58.5) | 690 (41.5) | 1.76 | (1.42–2.18) | <0.001 |
| PI-experienced | 1325 (62.7) | 789 (37.3) | 1.10 | (0.89–1.37) | 0.369 |
| T20-experienced | 54 (66.7) | 27 (33.3) | 0.73 | (0.45–1.19) | 0.207 |

This multivariate analysis was conducted with the endpoint of having a TBT score > 2.

Values in the TBT score columns are n (%), with the exception of mean (median) for age. AOR for age is for each additional 10 years.

*Reference category.

AOR, adjusted odds ratio; CI, confidence interval; ND, not determined; EFV, efavirenz; NVP, nevirapine; PI, protease inhibitor; T20, enfuvirtide.

Y181C, G190A, K101E and A98G were the mutations more frequently represented. The DUET studies showed that at least three TMC125-associated mutations were required to impair the efficacy of the drug [7,8]. In our cohort, only 9.8% of sequences showed at least three TMC125-associated mutations, suggesting that the existence of this condition is infrequent even in patients with evidence of resistance to the other NNRTIs. V179F, Y181V and G190S, which have the most pronounced effect on the response, were present in <5% of sequences. When at least three TMC125 RAMs were present, the mutations most frequently represented were confirmed to be Y181C, G190A and K101E, but not A98G. In this setting, the prevalence of V179F, Y181C and G190S also increased.

Y181C, a common mutation which confers resistance to other NNRTIs and to TMC125 when associated with two or more TMC125 RAMs and which was highly prevalent (32.2%) in the Tibotec data set [16], was associated with at least two mutations in a higher percentage of sequences in this study than found in the DUET studies (27% vs. 15%, respectively) [7], but it was present with V179F and G190S in <5% of sequences. The association of Y181C with G190A, K101E and A98G was statistically significant. The prevalence of V179F was low, but when associated with at least two mutations was present in 57% of sequences and was associated most frequently with Y181C, but was never associated with G190S or Y181V. Y181V and G190S, the other mutations with a large impact on response, were associated with at least two mutations in a low percentage of sequences.

The multivariate analysis of determinants of TBT WGS >2 demonstrated a protective effect of higher CD4 counts and a negative effect of higher HIV-RNA values, as expected. The use of EFV resulted in less NNRTI resistance than did the use of NVP. The pattern of resistance mutations suggests that subsequent virological suppression with TMC125-containing regimens may be more successful if previous treatments included EFV rather than NVP. The difference between exposure to NVP and EFV might be relevant in resource-limited settings where NVP is often used. The long-term use of NVP without optimized nucleoside reverse transcriptase inhibitor (NRTI) background therapy could lead to an accumulation of resistance mutations. This is of particular relevance in situations where second-line HAART regimens are difficult to obtain.

The use of both NNRTIs, rather than the duration of NNRTI exposure, had an impact on the occurrence of TBT WGS >2. As many HIV-positive patients still initiate therapy with an NNRTI, it is particularly important to take this evidence into consideration. Of note, lower CD4 counts (<200 cells/µL) and higher HIV RNA loads (>3.7 log_{10} copies/mL) were related to a greater risk of a TBT score >2.

The judicious examination of subjects’ therapeutic histories and the use of TBT WGS were found to be effective in predicting resistance to TMC125. The adoption of such tools is recommended for evaluating new antiretrovirals for clinical use.

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References


