

Next Generation Sequencing Analysis of HIV-1 Group O Reverse Transcriptase Residue 181C Prevalence and Evolution over Time, With or Without Antiretroviral Selection Pressure

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Next Generation Sequencing Analysis of HIV-1 Group O Reverse Transcriptase Residue 181C Prevalence and Evolution over Time, With or Without Antiretroviral Selection Pressure

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BACKGROUND

HIV-1 group O viruses are endemic in Cameroon and found sporadically in other countries. Their genetic divergence from pandemic HIV-1/M causes polymorphisms on residues associated to HIV-1/M antiretroviral resistance.

HIV-1	group	Μ
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HIV-1/0-H HIV-1/O-T **HIV-1 group O:**

* 181Y	→ ▼ 181C	——→ ▼ 181Y	X181C		X TOT I	
Susceptible to NNRTIs	Resistant to NNRTIs	Susceptible to NNRTIs	Resistant to NNRTIs	2	Susceptible to NNRTIs	
High replication fitness	Low replicative Fitness	High replication fitness	High replicative fitness	•	Low replicative Fitness	

Non-Nucleotide Reverse Transcriptase Inhibitors (NNRTI) resistance mutation Y181C can be selected in HIV-1/M but naturally present in HIV-1/O, and associated to the recently emerged HIV-1/O-H subgroup¹. A previous study suggested that residue 181C could confer a better replicative fitness to HIV-1/O in vitro and that signature residues were associated to 181Y-like or 181C-like lineages².

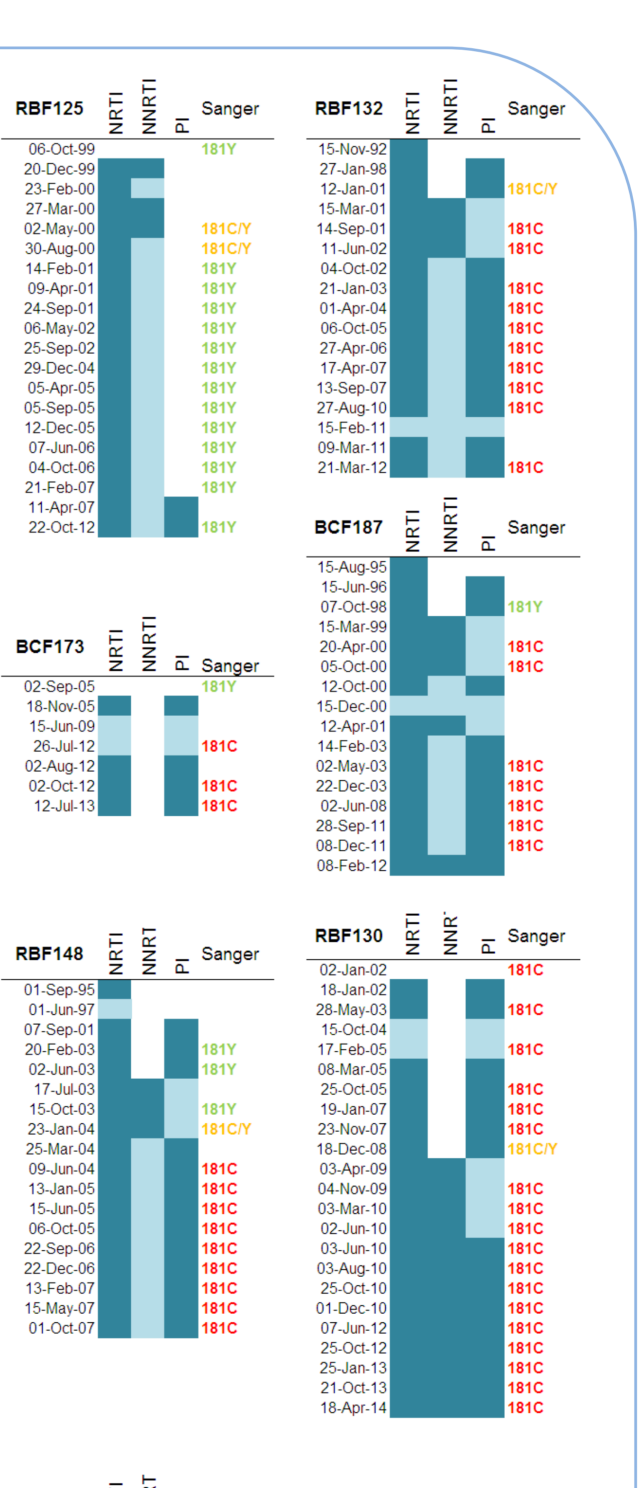
Here we aimed at exploring these hypotheses using in vivo samples and to investigate evolution of group O RT residue 181 under selection pressure due to NNRTI based treatment or not.

METHODS

We used Next Generation Sequencing to study residue 181 distribution and associated signature residue polymorphisms in 75 NNRTI-naïve HIV-1/O patients (1). Evolution of residue 181 over time – under selection pressure due to NNRTI based treatment or not – was investigated by Sanger and confirmed by NGS for some samples in 8 patients (2). We compared the viral loads from 59 untreated patients depending on residue 181 (3).

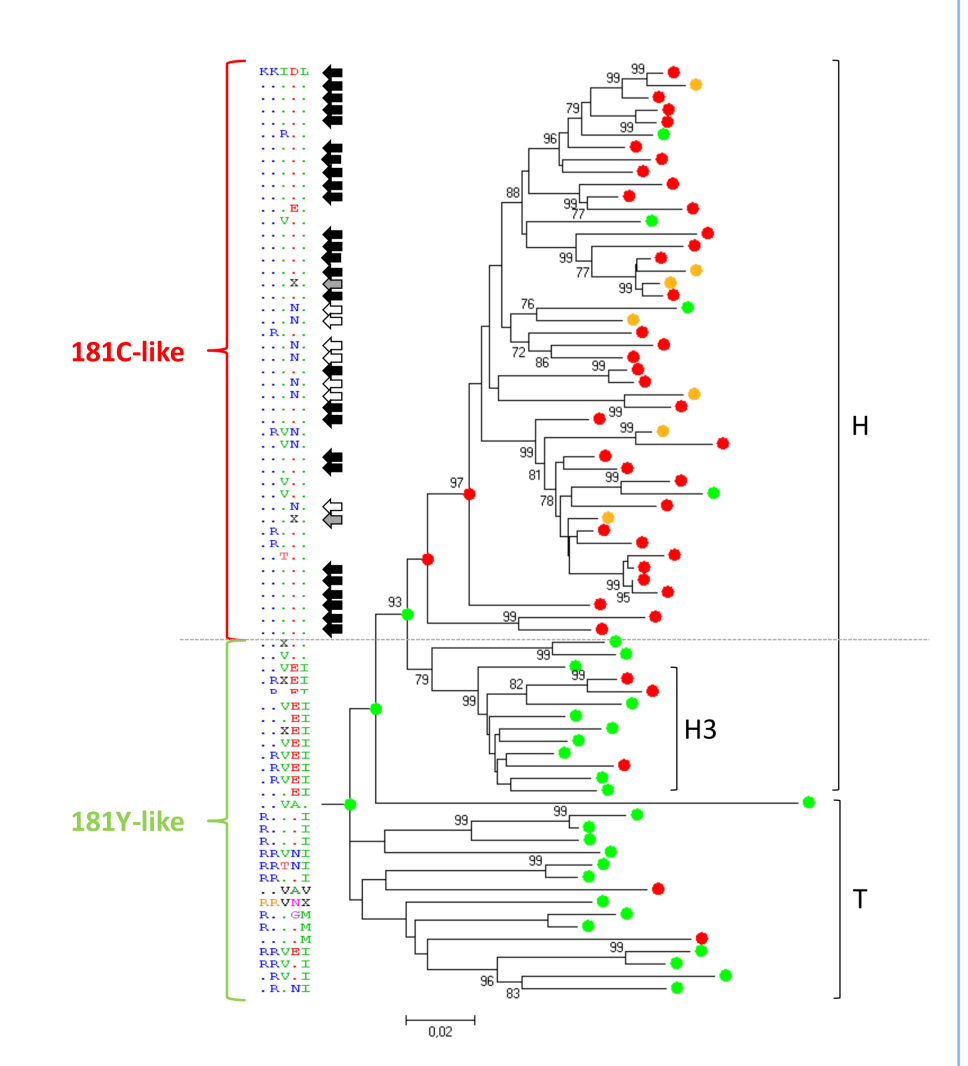
RESULTS (1) Residue 181C (red) was found in 40/75 NNRTI naive patients. Its association with HIV-1/O-H was confirmed (p<0.001). A 181C/Y mixture (orange) was found in 7 unlinked individuals.

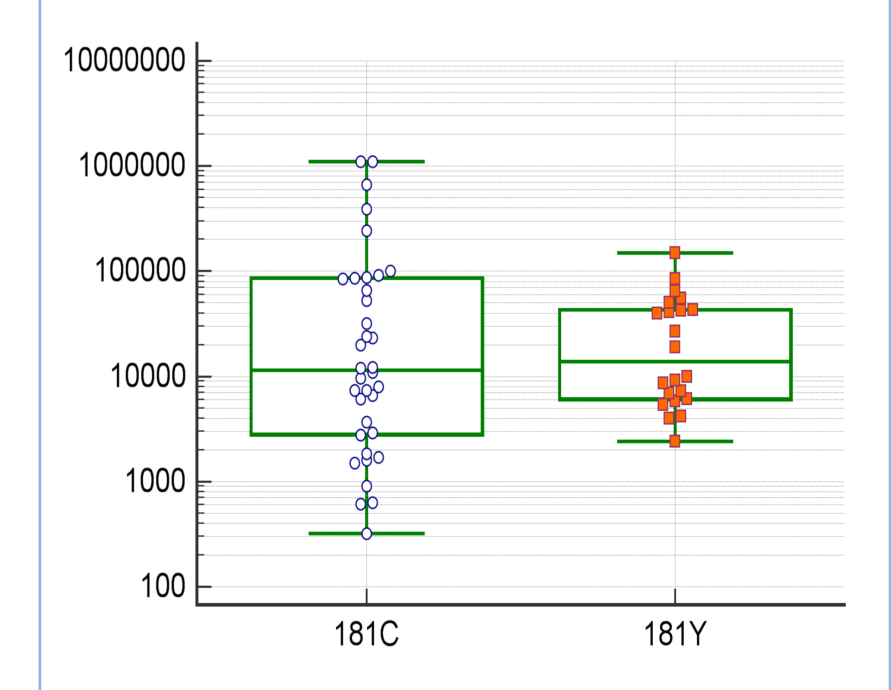
RESULTS (2) Evolution of residue 181 was similar to what observed under NNRTI in HIV-1/M for one 181Y HIV-1/O-T virus (RBF125). Three HIV-1/O-H viruses selected 181C due to NNRTIs, but conserved it several years after NNRTI interruption (RBF132, BCF187, RBF148). Four HIV-1/O-H viruses evolved without NNRTIs (C/Y => C, n=2, C)=> C/Y, n=2).



RESULTS (3)

The viral load range in untreated patients infected by 181C viruses (min: 2.5log cp/ml ; max: 6.0log cp/ml; N=37) was larger than that of 181Y virus infected patients (min: 3.4log; max: 5.2log; N=22).





The mean viral load was higher in the 181C group (5.1log, median: 4.1log) than in the 181Y group (4.5log, median: 4.2log), but not significantly according to Student t test (p=014).

Residues at signature positions were diverse in 181Y viruses (green) but a 28K-103K-142I-174D-178L pattern was highly conserved in 181C viruses (black arrows).

ARV naive
Under ARV treatment
Previously treated
*Confirmed by NGS

BCF177	NR	NNF	_ Sanger	RBF145	L L	NRT	
29-Apr-97					Ż	z	Sanger
31-Jan-01			181Y	11-Dec-02			181C
16-May-07				17-Jun-03			181C
09-Jan-08				09-Apr-06			181C
04-Mar-10				28-Feb-07			181C/Y
11-Oct-11			181C/Y	08-Apr-08			181C/Y
02-Jul-12			181C	30-Oct-08			181C/Y
31-Oct-13			181C	06-Oct-10			

CONCLUSIONS

Mutation 181C presence and evolution in HIV-1/O is **linked to the virus genetic background**.

It is associated to the emergent HIV-1/O-H subgroup where it can be naturally present, or conserved after NNRTI selection, possibly due to a favourable pattern (28K-103K-142I-174D-178L) on associated signature residues.

BCF1

However, no replicative advantage is observed for 181C viruses in vivo.

¹Leoz et al., PLOS Pathogens 2015. ²Tebit et al., AIDS Research and Human Retroviruses, 2016.