Sequence Note

Near Full-Length Genome Analysis of HIV Type 1 CRF02.AG, Subtype C and CRF02.AG Subtype G Recombinants

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ABSTRACT

HIV-1 CRF02.AG strains are prevalent in west and west-central Africa, suggesting a longstanding presence of these subtype A/G recombinants in the global epidemic. Cocirculation of CRF02.AG strains with other group M subtypes may give rise to HIV-1 recombinants constituting a mosaic genome comprising fragments of three different subtypes. We report on the genetic analysis of the near-full-length genomes of such recombinants (VI1035 and VI1197) as well as CRF02.AG strains in Belgian individuals. VI1035 and VI1197 may be the result of successful “second-generation” recombinations of HIV-1 strains CRF02.AG with, respectively, subtype C (VI1035) and G (VI1197) strains in a dually infected individual.

The recognition that intersubtype recombination is an important source of HIV-1 variation has led to new HIV-1 classification criteria. A subtype is now defined as a monophyletic cluster of three epidemiologically unlinked variants. The monophyletic clustering must be maintained regardless of the subgenomic segment analyzed to fulfill the criterion for a separate subtype. HIV-1 group M is currently subdivided in subtypes A, B, C, D, F, G, H, J, and K. Following the same lines, circulating recombinant forms (CRFs) are defined as intersubtype recombinants for which at least three epidemiologically unlinked variants are monophyletic and thus have an identical mosaic genome structure. So far CRF01.AE, CRF02.AG, CRF03.AB, and CRF04.cpx fulfill these criteria.1 This classification forms a framework to identify new subtypes and intersubtype recombinants that are an important source of HIV-1 genetic variation. Whereas CRF03.AB and CRF04.cpx apparently still have a relatively limited spread, CRF01.AE and CRF02.AG have been well established into different populations most likely as a result of a founder effect. CRF01.AE is common in the Central African Republic, Thailand, and other Asian countries.2 Near-full-length CRF02.AG recombinants were identified in Nigeria, Djibouti, and Côte d’Ivoire.3 Based on genetic analysis of small genome fragments, a high prevalence of CRF02.AG was documented in west and west-central Africa suggesting a longstanding presence of these recombinants in the global epidemic. Indeed, reanalysis of env subtype A strains documented before the recognition of CRF02.AG, as well as analysis of newly documented samples indicate the epidemiological relevance of this CRF. Reanalysis of partial env sequences from Côte d’Ivoire and Cameroon, initially documented as subtype A, indicates that strains of HIV-1 similar to CRF02.AG are the most common form of subtype A in west and west-central Africa.3 We demonstrated by reanalysis of env sequences encoding the region C2V3 till the start of gp41 (900 base pairs) of subtype A samples from Côte d’Ivoire and Cameroon, in combination with gag HMA that CRF02.AG

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strains were already prevalent in these countries in the early 1990s. We recently documented by gag HMA\(^4\) that CRF02.AG is prevalent in Yaoundé (Cameroon) and Cotonou (Bénin), comprising about 40% of the documented samples. Another recent study on env C2V3 phylogenetic analysis of samples from the northern part of Cameroon documented one-third of the analyzed samples as CRF02.AG variants.\(^5\) The longstanding presence of CRF02.AG strains in regions where other group M subtypes cocirculate may give rise to a next generation of HIV-1 recombinants constituting a mosaic genome comprising fragments of three different subtypes. Here we report on the genetic analysis of the near-full-length genomes of such recombinants (VI1035 and VI1197) as well as CRF02.AG strains in Belgian individuals.

The Belgian individual infected with VI1035 (isolated June 30, 1993) was probably infected by her female Kenyan partner from the border region between Kenya and Tanzania. He later also had multiple heterosexual contacts in other parts of Africa and Asia. The Belgian individual infected with VI1197 (isolated February 1, 1994) was infected by blood transfusion in Belgium in June 1989. VII144, a follow-up sample of VI1035, was previously classified as subtype C by env HMA,\(^6\) but could not be classified by C2V3 phylogeny. VII1197 was previously classified as env subtype G by env HMA\(^6\) and phylogenetic analysis of a 900 base pair (bp) fragment encoding C2V3 until the start of gp41.\(^7\)

Near-full-length genomes of VI1035 and VII1197 were amplified by polymerase chain reaction (PCR) from DNA isolated from cocultured peripheral blood mononuclear cells (PBMC), cloned and sequenced as described previously.\(^9,10\) The lengths of the clones of VI1035 and VII1197 were 8945 and 8914 bp, respectively. All reading frames were open and of complete length. The nucleotide sequence data were deposited in the EMBL, GenBank, and DDBJ Nucleotide Sequence Databases under the following accession numbers: AJ276595 (VI1035) and AJ276596 (VI1197).

VI1035 and VII1197 were compared with a panel of 24 near-full-length HIV-1 including group M subtypes A (UG037, Q23), B (MN, SF2), C (ETH2220, BR025), D (ELI, UG114), F (V1850, BR020), G (HH8793, SE6165), H (VI997, 90CF056.1), J (SE9280, SE9173), K (MP535, EQTB11), and CRF01.AE (CM240, CF402), CRF02.AG (InNG, DJ263), and CRF04.cpx (CY032.3, PMVY) reference sequences in order to verify their recombinant status. Near full-length HIV-1 sequences were aligned automatically using CLUSTAL W,\(^11\) and the resulting alignment was refined manually with the alignment editor DCSE.\(^12\) Regions that could not be aligned unambiguously due to excessive length or sequence variation were omitted from the analysis. Similarity plot and bootscanning were performed using a sliding window of 300 base pairs (bp) advanced in 10-bp increments using SimPlot software.\(^13\) Bootscanning phylogeny involved generation of 1000 replicates, distance calculation by Kimura (2-parameter), transition/transversion ratio 2.0, and tree construction by neighbor joining. Plots of similarity were generated for identification of reference sequences having the highest similarity to (parts of)

![Bootscanning phylogeny](image-url)

FIG. 1. (a) Bootscanning analysis of the complete genome of VI1035. Genome alignment positions and bootstrap values (%) are depicted in the x- and y-axis, respectively. VI1035 is compared with representatives of subtype B (MN; outgroup), C (IN21068), and CRF02.AG (DJ264). White regions I, IV, and VII cannot be assigned to a particular subtype or CRF by phylogenetic analysis.
FIG. 1. Continued.
VI1035 and VI1197 (results not shown) and used for bootscanning analysis. Similarity plot analysis reveals mosaic structures including genome segments that are CRF02.AG like and subtype C (VI1035) or G (VI1197) segments; the graph of bootscanning analysis identifies the breakpoints (Figs. 1a and 2a). To confirm these results, the software package TREECON was used for distance calculations (Kimura), tree construction (neighbor joining), and bootstrap analysis (Figs. 1b and 2b). The genome structure of VI1035 was as follows (Figs. 1a and b). A segment comprising the gag encoding region up to half of p24 clustered with subtype A, CRF01.AE, CRF02.AG, and CRF04.cpx [I; alignment position (AP) 1–875; genome position (GP) 1–886].

A BLAST search (15; http://hiv-web.lanl.gov/) did not reveal any significant relationship of (parts of) this fragment to subtype A, CRF01.AE, CRF02.AG, or CRF04.cpx. The second half of the gag p24 region clusters with subtype C (II; AP 876–1375; GP 887–1383). The pol gene starts VI1035 turns to CRF02.AG for a fragment encoding protease, reverse transcriptase, and most of RNase H (III; AP 1376–3375; GP 1384–3396). A segment of the pol region encoding the remaining part of the RNase H as well as part of the integrase remains unclassified (IV; AP 3376–3775; GP 3397–3795). The remainder of the pol gene and vif cluster with CRF02.AG (V; AP 3776–4875; GP 3796–4894). A region comprising vpr, tat (exon 1), rev (exon 1), vpu, and an env gp120 segment encoding V1 and V2 is subtype C (VI; AP 4876–6150; GP 4895–6315). The env C2V3 region remains unclassified (VII; AP 6151–6450; GP 6316–6585). A fragment encoding the remaining part of env gp120, the external part and the membrane spanning domain of gp41, is subtype C (VIII; AP 6451–7525; GP 6586–7758). The env encoding internal part of gp41, the second exons of tat and rev, nef, and part of LTR cluster with CRF02.AG (IX; AP 7526–8689; GP 7759–8945).

The genome structure of VI1197 was as follows (Figs. 2a and b, and 3). A segment comprising the gag gene and the protease encoding part of the pol gene clustered with CRF02.AG (I; AP 1–1900; GP 1–1707). Part of the pol reverse transcriptase encoding region is subtype G (II; AP 1901–2300; GP 1708–2306). A pol gene segment encoding part of reverse transcriptase, RNase H, and part of integrase clusters with CRF02.AG (III; AP 2401–4100; GP 2407–4102). The remainder of the pol gene of VI1197 clusters with subtype G (IV; AP 4101–4400; GP 4103–4401). Most of the vif gene remains unclassified (V; AP 4401–4800; GP 4402–4801). A segment including the terminal part of the vif gene, vpr, tat (exon 1), rev (exon 1), vpu, env gp120, and the external part of gp41 encoding regions is classified as subtype G (VI; AP 4801–7300; GP 4802–7489). A segment encoding the env membrane spanning domain, part of the internal part of gp41, the second exon of tat, and part of the second exon of rev, remains unclassified (VII; AP 7301–7700; GP 7490–7884). The remaining part of the env gp41 and the rev second exon encoding regions, nef, and part of LTR cluster with subtype G and CRF02.AG (VIII; AP 7701–8689; GP 7885–8914).

FIG. 2. (a) Bootscanning analysis of the complete genome of VI1197. Genome alignment positions and bootstrap values (%) are depicted in the x- and y-axis, respectively. VI1197 is compared with representatives of subtype B (MN; outgroup), G (SE6165), and CRF02.AG (D1264). White regions V and VII cannot be assigned to a particular subtype or CRF by phylogenetic analysis. (b) Phylogenetic classification of genome segments based on an alignment of HIV-1. The phylogenetic position of VI1197 is indicated in italic; 2000 bootstrap samples were analyzed. Bootstrap values are given in percentages at the internodes if they exceed the 70% level. The tree is rooted arbitrarily between subtypes.
The long-standing presence of non-B strains in HIV-1-seropositive Belgian individuals was documented earlier. Generally these infections were linked to regions in the world where these strains are prevalent. To verify to what extent CRF02.AG variants were introduced into the Belgian population, we screened previously documented HIV-1 \textit{env} subtype A-infected individuals for actual CRF02.AG infections by phylogenetic analysis including CRF02.AG references, as well as by \textit{gag} HMA. Four of 29 “A” variants, introduced to the Belgian population 8 to 10 years ago, were classified as
CRF02.AG by gag HMA, of which three phylogenetically clustered with env C2V3 CRF02.AG references supported by bootstrap analysis (results not shown; Table 1). These findings indicate a longstanding presence of HIV-1 CRF02.AG strains in Belgium.

It was previously recognized that the frequency of documenting HIV-1 recombinants in seropositive individuals correlated with cocirculation of multiple HIV-1 subtypes in populations in particular geographic regions. Since HIV-1 CRF02.AG strains are prevalent in west and west-central Africa, it may not be surprising that CRF02.AG variants contributed to “second-generation” recombinants comprising fragments of three different subtypes, that are transmitted into the population. In addition to the mosaic CRF02.AG-C (VI1035) and CRF02.AG-G (VI1197) genomes documented in this study, we previously identified one isolate of gag/env A/CRF02.AG, CRF02.AG/F, and CRF02.AG/G recombinants in Côte d’Ivoire, and Bénin, respectively.

VI1035 and VI1197 may be the result of a successful recombination of HIV-1 strains of CRF02.AG with, respectively, subtype C (VI1035) and G (VI1197) strains in a dually infected individual. The evolution toward “second-generation” recombinants contributes to the unpredictability of evolutionary routes in the further evolution of HIV-1.

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**TABLE 1. Epidemiological Data of HIV-1 Seropositive Individuals**

<table>
<thead>
<tr>
<th>Isolate number</th>
<th>Source of infection</th>
<th>Time of infection</th>
<th>Risk behavior</th>
<th>Genetic classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>VI712</td>
<td>DRC</td>
<td>May 1992</td>
<td>Heterosexual</td>
<td>CRF02.AG CRF02.AG</td>
</tr>
<tr>
<td>VI860</td>
<td>Nigeria</td>
<td>Between December 1989 and December 1992</td>
<td>Heterosexual</td>
<td>CRF02.AG CRF02.AG</td>
</tr>
<tr>
<td>VI1007</td>
<td>Côte d’Ivoire</td>
<td>Before 1990</td>
<td>Heterosexual</td>
<td>CRF02.AG A+CRF02.AG</td>
</tr>
<tr>
<td>VI1035</td>
<td>Africa and/or Asia</td>
<td>Before 1993</td>
<td>Heterosexual</td>
<td>C U CRF02.AG-C</td>
</tr>
<tr>
<td>VI1144</td>
<td></td>
<td></td>
<td></td>
<td>C U CRF02.AG</td>
</tr>
<tr>
<td>VI1197</td>
<td>?</td>
<td>?</td>
<td>Blood transfusion</td>
<td>CRF02.AG G CRF02.AG-G</td>
</tr>
<tr>
<td>VI1320</td>
<td>West-Africa</td>
<td>May 1990</td>
<td>Heterosexual</td>
<td>CRF02.AG CRF02.AG</td>
</tr>
</tbody>
</table>

*The gag region as determined by gag HMA encodes from amino acid 132 of p24 until amino acid 40 of p7 (according to HIV-1 ELI; accession number K03454).

bThe env C2V3 region as determined by sequencing and phylogenetic analysis.
REFERENCES


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