Most env and gag Subtype A HIV-1 Viruses Circulating in West and West Central Africa Are Similar to the Prototype AG Recombinant Virus IBNG

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Summary: The genetic subtype was identified in gag and env of 219 HIV-1-positive samples collected in different African countries, 44 from Senegal, 55 from Cameroon, 82 from Gabon, and 38 from Djibouti. In total, 20 (9.1%) samples had discordant subtypes between gag and env, 6 of 44 (13.9%) in Senegal, 4 of 55 (7.2%) in Cameroon, 1 of 38 (2.6%) in Djibouti, and 10 of 82 (12.1%) in Gabon. Subtypes A and G were predominantly involved in the recombination events. Phylogenetic tree analysis of gag showed that an important number of the A sequences form a distinct subcluster with the AG-IBNG prototype strain (a complex A/G mosaic virus): 27 of 32 (84.3%) in Senegal, 12 of 17 (70.6%) in Nigeria, 24 of 39 (61.5%) in Cameroon, and 38 of 70 (54.3%) in Gabon. Full-length genome analysis of 3 and additional sequences in pol for 10 such strains confirmed that they have a similar complex A/G mosaic genomic structure. These data suggest that in West Africa, most probably between 60% and 84% of the subtype A viruses are recombinant AG-IBNG viruses. This finding has potential implications on future vaccine, diagnostic, and treatment strategies. The actual and future role of these viruses in the global pandemic must be monitored in all new molecular epidemiologic studies, a discrimination between subtype A and AG-IBNG-like viruses is necessary. Key Words: HIV-1 subtypes—env—gag—Recombination—AG-IBNG—Africa.

Phylogenetic analysis of HIV-1 viruses from different geographic locations has revealed that HIV-1 can be divided into at least three distinctive groups, M, N, and O (1). Group M, which is the major group, comprises most HIV-1 strains that cause AIDS worldwide. Full-length genome sequencing showed that group M HIV-1 viruses can be further subdivided into at least nine different, nonrecombinant, subtypes (A, B, C, D, F, G, H, J, and K) approximately equidistantly related with intrasubtype divergence up to 20% and intersubtype divergence between 25% to 35%, for the Env amino acid sequences (2–4). HIV-1 group O viruses have only 50% similarity with the HIV-1 group M isolates in the env gene (5,6). There are also many mosaic genomes of HIV-1, some of which are unique or restricted to one transmission cluster, and others that are major circulating forms (3). It was proposed to designate those recombinant viruses as...
“circulating recombinant forms (CRF),” by associating CRF with the name of the first full-length sequence of that form. There are currently 4 CRFs of HIV-1: AE-CM240 from southeast Asia, the AG-IBNG from Africa, the AG1-CY032 from Cyprus and Greece, and AB-KAL153–like viruses found in the epidemic among intravenous drugs users (IVDUs) in Kaliningrad in Russia (3).

The various HIV-1 groups and subtypes differ in their geographic dissemination, and so the subtype designations have been critical molecular epidemiologic markers for tracking the course of the global pandemic. HIV-1 group O seems to be endemic in Central West Africa, especially Cameroon, where the frequency of infection is estimated to be 2% and 5% of HIV-1–infected individuals (7–9). Group N viruses were recently reported from only 2 Cameroonian patients (1). Among HIV-1 group M viruses, geographic distribution of the different subtypes is also heterogeneous. In Europe and in North and South America, subtype B is largely dominant, whereas in South and East Africa and in India subtype C is spreading rapidly (10–12). In Southeast Asia, the circulating recombinant AE-CM240 is predominant (12,13). In Africa, all HIV-1 groups and HIV-1 group M subtypes have been documented; however, their distribution on the African continent is not homogeneous (14). Subtype A is predominant in West and Central Africa whereas subtype C is more common in East and South Africa. Subtype D is present in Central and East Africa with prevalences ranging from 5% to more than 40% of the circulating strains (15,16). Subtype G has been documented in many West and Central African countries, whereas subtype H was only found in Central Africa (2,15). In certain populations and regions where multiple HIV-1 subtypes co-circulate, many combinations of intersubtype recombinant viruses have been documented (A/C, A/D, B/F, A/G/W/1) (17–20), and even an intergroup M/O recombinant virus has been recently isolated from a Cameroonian patient (21). Recombinant viruses may have some advantages over the parental strain, including eventual modifications in tropism and replication efficiency (fitness). Therefore, recombination may play a significant role in global HIV evolution, creating novel viral genotypes within populations.

Recombination can be missed if large portions or different regions of the viral genome are not examined. The AG-IBNG recombinant virus has been documented in several African countries (22); however, the real prevalence of this virus in the global epidemic is still unknown. This virus has a complex mosaic genomic structure involving subtype A and G sequences with multiple breakpoints, but given that both gag and env derive largely from subtype A, these viruses were initially classified as subtype A (23,24). However, they form a significant subcluster within subtype A in the gag and env regions and can therefore be recognized even by using partial sequencing of familiar regions (22).

In this study, we tried to identify the proportion of recombinant HIV-1 strains by identifying the genetic subtype in gag and env of more than 200 HIV-1 strains collected in different geographic regions in Africa. We also estimated the proportion of AG-IBNG–like viruses among gag or env subtype A viruses.

MATERIALS AND METHODS

Study Samples

In total, 219 randomly chosen HIV-1–positive samples of African origin were tested, 55 from patients living in Cameroon, 44 from Senegal, 38 from Djibouti, and 22 from Gabon. All samples from Senegal have been collected among patients seen at one of three major hospitals in Dakar between 1997 and 1998. All Cameroonian samples were collected between 1995 and 1997 from patients visiting the military hospital in Yaounde. The samples from Djibouti were all from citizens of metropolitan France, as previously described (25), who had become infected with HIV-1 after overseas deployment in this country. Most remained asymptomatic but had been infected between 1987 and 1997. The samples from Gabon have been collected in 1997 among different population groups.

DNA was extracted from the primary or short-term cultured peripheral blood mononuclear cells (PBMCs) using Qiagen (Qiagen S.A., Courtaboeuf, France) DNA isolation kit.

Genetic Subtyping in the gag Region

A 700-bp fragment, corresponding to the p24 region from the gag gene, was amplified with previously described primers, G00-G01 and G60-G25 (26). The conditions for polymerase chain reaction (PCR) were as follows: a first denaturation step for 3 minutes at 92°C, followed by 30 cycles of 92°C for 10 seconds, 55°C for 30 seconds, and 1 minute at 72°C, with a final extension for 7 minutes at 72°C, in a final volume of 50 μl. The reaction mixture consists of 50-mmol/l KCl, 10-mmol/l Tris-HCl (pH 9), 0.1% Triton X-100, 1.4-mmol/l MgCl2, 10-nmol each primer, 0.2-mmol/l each dNTP, and 2.5-U Taq polymerase. Moreover, 1 μl of this amplified product was used for the second round using the same reaction mixture and PCR conditions for 40 cycles, in a final volume of 100 μl. The PCR amplification products were detected by electrophoresis on a 1% agarose gel and visualized by ethidium bromide staining.

Nucleotide sequences were obtained by direct sequencing of the PCR products. The amplified DNA was purified using a QiAquick gel extraction kit (Qiagen S.A., France). Cycle sequencing was performed using fluorescent dye terminator technology (dye terminator cycle sequencing with AmpliTaq DNA polymerase FS (PE Biosystem) according to the manufacturer’s instructions. Electrophoresis and data collection were done on an Applied Biosystems 373A automatic DNA sequencer (stretch model, PE Biosystem, Courtaboeuf, France).

JAIDS Journal of Acquired Immune Deficiency Syndromes, Vol. 23, No. 5, April 15, 2000
The amplified fragments were purified with the Qiaquick gel extraction kit (Qiagen) and then directly sequenced on an automatic sequencer (Applied Biosystems 373A stretch model, PE Biosystems) using dye terminator technology on an automated DNA sequencer (ABI 373A stretch, Applied Biosystems). The sequenced fragments were analyzed by a series of overlapping nested PCRs amplifying fragments of 2800 bp with an approximate 700 bp overlap.

We used the Boehringer Long Template Expand DNA polymerase (Boehringer Mannheim, Laval, Qc, Canada), according to the instructions of the manufacturer. The overlapping PCRs fragments from each patient were either cloned and sequenced or the PCR product was directly sequenced. Sequencing was performed by using cycle sequencing and dye terminator technology on an automated DNA sequencer (ABI 373A stretch, Applied Biosystems). The sequenced fragments were reanalyzed and assembled into contiguous sequences by using the Seqed program (Applied Biosystems).

### TABLE 1. Distribution of HIV-1 genetic subtypes in gag (p24) and env (V3-V5) genes among samples collected in various African countries

<table>
<thead>
<tr>
<th>Country</th>
<th>Genotype</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
<th>H</th>
<th>Discordant samples</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Senegal (n = 44)</td>
<td>gag</td>
<td>32 (72.7)</td>
<td></td>
<td></td>
<td>5 (11.3)</td>
<td>4 (9.1)</td>
<td></td>
<td></td>
<td>2 (4.5)</td>
<td>1 (2.2)</td>
<td>5 (11.4)</td>
</tr>
<tr>
<td></td>
<td>env</td>
<td>33 (75.0)</td>
<td></td>
<td></td>
<td>6 (13.6)</td>
<td>4 (9.1)</td>
<td></td>
<td></td>
<td>1 (2.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cameroon (n = 55)</td>
<td>gag</td>
<td>39 (70.9)</td>
<td></td>
<td></td>
<td>1 (1.8)</td>
<td>2 (3.6)</td>
<td>9 (16.3)</td>
<td></td>
<td>4 (7.2)</td>
<td></td>
<td>4 (7.2)</td>
</tr>
<tr>
<td></td>
<td>env</td>
<td>39 (70.9)</td>
<td></td>
<td></td>
<td>1 (1.8)</td>
<td>2 (3.6)</td>
<td></td>
<td></td>
<td>11 (20)</td>
<td>2 (3.6)</td>
<td>4 (7.2)</td>
</tr>
<tr>
<td>Djibouti (n = 38)</td>
<td>gag</td>
<td>6 (15.7)</td>
<td>4 (10.5)</td>
<td>27 (71)</td>
<td>1 (2.6)</td>
<td>1 (2.6)</td>
<td></td>
<td></td>
<td></td>
<td>1 (2.6)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>env</td>
<td>6 (15.7)</td>
<td>4 (10.5)</td>
<td>27 (71)</td>
<td>1 (2.6)</td>
<td>1 (2.6)</td>
<td></td>
<td></td>
<td></td>
<td>1 (2.6)</td>
<td></td>
</tr>
<tr>
<td>Gabon (n = 82)</td>
<td>gag</td>
<td>70 (85.3)</td>
<td></td>
<td></td>
<td>5 (6.1)</td>
<td>1 (1.2)</td>
<td>1 (1.2)</td>
<td>4 (4.8)</td>
<td>2 (2.4)</td>
<td>10 (12.1%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>env</td>
<td>69 (81.4)</td>
<td></td>
<td></td>
<td>5 (6.1)</td>
<td>1 (1.2)</td>
<td>4 (4.8)</td>
<td>6 (7.3)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total (n = 219)</td>
<td>gag</td>
<td>147 (67.1)</td>
<td>4 (1.8)</td>
<td>35 (15.1)</td>
<td>12 (5.5)</td>
<td>10 (4.6)</td>
<td>12 (5.5)</td>
<td>9 (4.1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>env</td>
<td>147 (67.1)</td>
<td>4 (1.8)</td>
<td>39 (17.5)</td>
<td>8 (3.7)</td>
<td>1 (0.45)</td>
<td>12 (5.5)</td>
<td>9 (4.1)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Sequence Analysis of the pol Gene

The protease and RT genes were amplified with seminested PCR using G25rev (GCAAGAGTTTGGCTGAAAGCAATGAG) and IN3 (TCTATCCATCTAAAGAATAGTACACCTGATTCC) as outer primers and AV150 (GTTGGGAAGGACACAAATGGAAG) and IN3 as inner primers. The PCR parameters for the two rounds were as follows: a first denaturation step for 5 minutes at 92°C, followed by 40 cycles of 92°C for 20 seconds, 58°C for 30 seconds, and 72°C for 2 minutes. PCR reactions were done with the TTh DNA Polymerase. from Perkin Elmer (Perkin Elmer-Cetus, Norwalk, CT, U.S.A.) according to the instructions of the manufacturer. The amplified fragments were purified with the Qiagen gel extraction kit (Qiagen) and then directly sequenced on an automatic sequencer (Applied Biosystems 373A stretch model, PE Biosystems) using the ABI Prism Big Dye Terminator cycle sequencing technology.

### Phylogenetic Analysis

Nucleotide sequences were aligned using CLUSTAL W (28) with minor manual adjustments, bearing in mind the protein sequences. Regions that could not be aligned unambiguously, due to sequence variability, were omitted from analysis. Phylogenetic trees using the neighbor-joining method and reliability of the branching orders using the bootstrap approach were implemented using CLUSTAL W. Genetic distances were calculated with the Kimura's two-parameter method, (ratio Tt = 2.0) (29).

### TABLE 2. The different discordant gag/env subtype profiles observed in this study and the number of samples seen in each of the countries studied

<table>
<thead>
<tr>
<th>Subtype (gag-env)</th>
<th>Senegal</th>
<th>Cameroon</th>
<th>Djibouti</th>
<th>Gabon</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-G</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>A-C</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>A-B</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>A-F</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>D-A</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>D-B</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>G-A</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>H-A</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
<tr>
<td>H-G</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1</td>
</tr>
</tbody>
</table>
Intersubtype Recombinant Analysis

To analyze whether the viruses were recombinant in the sequenced regions, several additional analyses were performed. Diversity plots, using the DIVERT program available online (accessed April 27, 2000; available at http://193.50.234.246/~beaudoin/anrs/Diversity.html), determined the percentage of diversity between selected pairs of sequences by moving a window of 300 bp along the genome alignment in 20-bp increments. Divergence values for each pairwise comparison were plotted at the midpoint of the 300-bp segment.

Simpplot 2.5 software (30), kindly provided by Stuart Ray, was used to calculate bootstrap plots. For the bootstrap plots, the Simplot software performed bootscanning on neighbor-joining trees by using SEQBOOT, DNADIST (with Kimura's parameter method and a transition:transversion ratio of 2.0), NEIGHBOR, and CONSENSUS from the Phylip package for a 500-bp window moving along the alignment in increments of 50 bp (this approach is similar to the bootscanning method described by Salminen et al. (31)). We evaluated 100 replicates for each phylogeny. Bootstrap values for the studied sequences were plotted at the midpoint of each window.
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TABLE 3. The number and percentages of the subtype A samples clustering with the AG-IBNG prototype strain in gag (p24), and the genetic distances observed in the different subtype A clusters

<table>
<thead>
<tr>
<th>Country</th>
<th>IBNG n (%)</th>
<th>non-IBNG n (%)</th>
<th>Genetic distances (mean and extreme values)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Senegal</td>
<td>27/32 (84.3)</td>
<td>5/32 (15.7)</td>
<td>IBNG: 4.3 (1.7–7.7) non-IBNG: 7.5 (3.9–10.9) IBNG + non-IBNG: 6.2 (1.7–11.5)</td>
</tr>
<tr>
<td>Cameroon</td>
<td>24/39 (61.5)</td>
<td>15/39 (38.4)</td>
<td>IBNG: 4.8 (2.1–7.7) non-IBNG: 8.3 (2.9–10.6) IBNG + non-IBNG: 7.3 (1.7–11.2)</td>
</tr>
<tr>
<td>Djibouti</td>
<td>—</td>
<td>6/6 (100)</td>
<td>IBNG: — non-IBNG: 7.2 (1.8–9.6) IBNG + non-IBNG: 7.2 (1.0–11.7)</td>
</tr>
<tr>
<td>Gabon</td>
<td>38/70 (54.3)</td>
<td>32/70 (55.7)</td>
<td>IBNG: 5.0 (1.6–9.9) non-IBNG: 7.5 (1.0–11.7) IBNG + non-IBNG: 7.2 (1.0–13.5)</td>
</tr>
</tbody>
</table>

Other subtypes: 5.4 (0.9–10.6).

Genetic distances were calculated with the Kimura two parameter method, using the same alignment as used for the construction of the phylogenetic trees in Figure 1.

Genbank Accession Numbers


The pol sequences are available in Genbank with the accession numbers AJ286134 to AJ286143. The near-full-length genome sequence accession numbers are: AJ286133 for 9CM-807, AJ251056 for 98SE-MP1211, and AJ251057 for 98SE-MP1213.

FIG. 2. Phylogenetic tree of the near-full-length sequences of the three new HIV-1 viruses with reference strains from each of the known subtypes. The analysis was performed as described in Materials and Methods and using the same references sequences (except K-V1325) as for the gag trees in Figure 1. Bootstrap values <70% are not shown.
RESULTS

Genetic Subtypes in the env and gag Genes

Table 1 shows the genetic subtypes identified in the V3 to V5 region of the envelope by HMA and the numbers and percentages of the genetic subtypes identified in the p24 region of the gag gene by sequencing followed by phylogenetic tree analysis, for the corresponding strains in the different countries. In the West and Central West African countries subtype A was predominant in gag and env, whereas in the East African country, Djibouti, subtype C was the classification of most subtype samples in the sequence, as commonly documented among circulating recombinant AE strains.

Among the 44 samples from Senegal, 5 (11.4%) had discordant subtype designations between envelope and gag. In Cameroon, 4 (7.2%) of 55 samples, and in Gabon 10 (12.9%) of 82 samples were discordant. In Djibouti, only 1 discordant sample was seen and this corresponded to an env subtype E sample, with a gag subtype A sequence, as commonly documented among circulating recombinant AE strains.

Table 2 shows the different discordant subtype profiles observed between gag and env in the different countries. Overall, among the total 20 discordant samples, 9 different profiles were seen and in 7 of them subtype A sequences, either in gag or env, were involved. In only 3 of 20 discordant samples, no A sequences were involved and among the other 17 samples, 9 were A in gag and 8 were A in the envelope.

Phylogenetic Analysis of the gag (p24) Sequences

Figure 1 shows the phylogenetic tree analysis of the gag sequences for the different geographic regions. These phylogenetic trees show that among the subtype A sequences, distinct subclusters can be identified, more specifically, an important number of the gag A sequences form a distinct subcluster with the AG-IBNG prototype strain (a complex mosaic virus [22]). These AG-IBNG clusters are supported with high bootstrap values of 97% in Cameroon and only 45% and 47% in Gabon and Senegal, respectively. None of the subtype A samples from Djibouti clustered with the AG-IBNG.

Table 3 summarizes the number of subtype A samples studied that may form a separate cluster with the AG-IBNG prototype strain. Overall, 89 of 147 (60.5%) subtype A gag sequences are close to AG-IBNG in the phylogenetic tree. In Senegal, 27 of 52 (51.9%) of the env subtype A viruses are IBNG-like in gag, whereas in Cameroon and Gabon, 24 of 39 (61.5%) and 38 of 70 (54.3%), respectively, of the subtype A strains cluster with AG-IBNG.

Genetic distances have been calculated for the subtype A strains and the subclusters observed within subtype A and the other subtypes (B–H) in the various countries. Data are summarized in Table 3. Genetic distances observed for the AG-IBNG cluster are significantly lower than those calculated for the other subtype A strains: 4.3% in Senegal, 4.8% in Cameroon, and 5% in Gabon versus 7.5%, 8.3%, and 7.2%, respectively, for the non–IBNG-like subtype A strains. The average genetic distances within the other subtypes, B to H, is 5.4% and are generally lower than for subtype A. This was confirmed in all three West African countries studied.

Sequence Analysis of Near-Full-Length Genomes of gag AG-IBNG-Like Viruses

To study whether gag subtype A viruses that cluster with the AG-IBNG prototype strain virus have a similar complex mosaic genome, the near-full-length genome was sequenced for 3 subtype A viruses clustering with this strain in gag: 2 from Senegal (98SE-MP1211 and 98SE-MP1213) and 1 from Cameroon (97CA-MP807). We obtained 8851 bp for MP1211, 8858 bp for MP1213, and 8800 bp for MP807. The sequences start between the first and the tenth amino acid from the gag polyprotein, LTR extremities were not sequenced. All other genes were complete without premature stop codons and no major deletion or insertion events were present in the sequences of these strains.

To determine the phylogenetic relationships of the newly characterized viruses, we constructed evolutionary trees from the near-full-length genome. The phylogenetic tree shows that these 3 new strains form a separate clus-
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with the AG-IBNG virus and with the previously described AG-IBNG–like viruses, D1263 and D1264 from Djibouti supported by high bootstrap values (100%; Fig. 2).

To characterize more precisely whether the 98SE-MP1211, 98SE-MP1213 and 97CA-MP807 viruses have the same mosaic structure as the AG-IBNG prototype virus, we performed pairwise sequence comparisons between the near-full-length genome sequence of these isolates and with those available from the database, using the same alignment of nucleotide sequences used for the phylogenetic tree construction. Importantly, distance values were calculated only after removing all sites with a gap in any of the sequences, ensuring that all comparisons were made across the same sites. When the new sequences were plotted against references of all nonrecombinant subtypes, we observed for all 3 sequences similar disproportionate levels of sequence divergence with subtype A and C along their genome, consistent with the pattern observed when the AG-IBNG prototype virus is plotted against the same reference strains (data not shown).

To evaluate systematically the bootstrap values supporting independent monophyletic phylogeny in different parts of the genome, the alignment was bootscanned in regard to the clustering of the putative AG-IBNG–like viruses with a sliding window of 500 bp advancing with 50-bp increments. The same multiple-genome alignment generated for the phylogenetic analysis of the near-full-length genomes shown in Figure 2 was used to calculate the consensus reference sequences (50% threshold) for the 8 groups of nonrecombinant subtypes corresponding to subtypes A, B, C, D, F, G, H, and J. The magnitudes of the bootstrapping values supporting the clustering of our strains with consensus sequences from members of each subtype was performed. The results of the bootscan analysis are shown in Figure 3. The complexity of these new strains is readily apparent with alternating segments clustering with subtype A and G, similar to the complexity observed for the AG-IBNG prototype strain. When the 98SE-MP1211, 98SE-MP1213, and 97CA-MP807 strains were bootscanned against an alignment that includes the AG-IBNG strain, bootscanning results of the entire genomes revealed that they all were similar to the AG-IBNG isolate. The lower bootstrap results found in certain regions of the genome result from the simultaneous presence in the analysis of nonrecombinant subtype A sequences and the A sequences from the AG-IBNG strain.

Sequence Analysis of the pol Region from gag Subtype A Viruses Including AG-IBNG–Like Viruses

The AG-IBNG HIV-1 virus is mosaic in the pol region. We therefore sequenced the pol region, covering the protease and reverse transcriptase (RT) genes for 10 subtype A samples in gag; among them 6 that had formed a separate cluster with AG-IBNG. Figure 4 shows the phylogenetic tree analysis of the 11 samples from the gag and pol sequences. Of the 6 samples clustering with AG-IBNG in gag, all formed a well-separated cluster with this strain in the pol region. For the other 4 subtype A strains, 1 was subtype A in pol, 1 clustered with AG-IBNG, and 2 were subtype J in pol. Diversity plotting and bootscanning on the pol sequences of the AG-IBNG–like sequences showed that all these sequences have a similar mosaic structure in pol as the AG-IBNG prototype strain. In Figure 4C, bootscan analysis in the pol gene is shown for one isolate from the study (97CM-MP803), as well for the AG-IBNG strain in the same region.

Estimated Prevalence of Recombinant HIV-1 Viruses and AG-IBNG–Like Viruses

Full-length genome analysis showed that the 3 HIV-1 viruses clustering in gag (p24) with the prototype AG-IBNG virus have a similar complex A/G mosaic genomic structure. Additional sequence analysis of the pol region, showed that 6 viruses clustering in gag and pol with AG-IBNG had a similar A/G mosaic structure in this gene. These data allow us to extrapolate that in West and Central West Africa, most probably between 60% and 84% of the subtype A viruses are AG-IBNG–like viruses. Moreover, some of the subtype A gag sequences involved in recombination (different env and gag subtypes) cluster with AG-IBNG in the gag region. More
viruses that circulate may even be higher. In a previous study, ranging from 2.6% in Djibouti to 12.1% in Gabon. These results are due to coinfections with divergent viruses. Based on partial subtypes that cocirculated and most probably resulted from recombination between different subtypes.

Addition of the AG-IBNG viruses to the numbers of viruses for which discordant gag and envelope subtypes were observed made the overall minimal percentages of recombinant virus involved in the HIV-1 epidemic in Africa very high (Table 4); 75.0% in Senegal (6 discordant with 27 AG-IBNG), 51.0% in Cameroon (4 discordant with 24 AG-IBNG), and 54.7% in Gabon (10 discordant with 38 AG-IBNG).

### DISCUSSION

Until recently, based on partial sequences of different genes or full-length genome sequencing, it was estimated that up to 10% of the HIV-1 strains in the global pandemic are intersubtype recombinants (32–34). All these recombinant viruses originated from geographic regions where multiple subtypes cocirculated and most probably result from coinfections with divergent viruses. Based on the discordant subtype results between partial gag and env sequences, 20 (9.1%) of the 219 strains in our study were potential recombinant viruses, with percentages ranging from 2.6% in Djibouti to 12.1% in Gabon. These data correspond to the global estimates on recombinant viruses that are circulating.

However, in some regions the number of recombinant viruses that circulate may even be higher. In a previous study on the genetic subtype distribution in Nigeria, we documented 12 (32.4%) of 37 samples from that country with discordant gag and env subtypes (35). These high values were also reported in another study on Nigerian samples (36).

In 1994, AG-IBNG, a new subtype A isolate from Ibadan, Nigeria, was reported on the basis of the gp120 sequence (23). Based on gag and env sequences, it was later described as a distinct subcluster within A subtype (24). With the recent availability of full-length subtype A and G sequences, it is clear that this strain is a complex mosaic virus with alternating subtype A and subtype G sequences (22). Geographic dispersion of the AG-IBNG subtype was already suggested by the presence of this isolate in Nigeria (in West Africa), and two viruses with a similar mosaic genome from Djibouti (DJ263 and DJ264) in East Africa (22). Additional full-length genome sequencing in our study from strains obtained in two African countries (two from Senegal and two from Cameroon) confirm the geographic spread of this AG-IBNG-like viruses. From our data and the previously described data on the Djibouti strains, we can extrapolate with a relative high probability that the viruses clustering with AG-IBNG in gag are predominantly AG-IBNG-like viruses. Our study shows on a large number of samples that in West and Central West Africa, AG-IBNG viruses comprise most of the env and gag subtype A strains currently circulating in these countries; 84.0% in Senegal, 61.0% in Cameroon, and 54.3% in Gabon. In a previous study on Nigerian samples, we have shown that most subtype A viruses from this country clustered with AG-IBNG: 71% based on p24 gag sequences and 64% based on env sequences (35). Genetic distances observed among the different AG-IBNG viruses indicate that they have been spreading for some time. The high prevalences of these particular A/G recombinant in Senegal, Gabon, Cameroon, and Nigeria are an additional factor in considering these viruses as a separate and distinct group in the global epidemic.

Virtually any isolate, including recombinant viruses, may compete more efficiently in a given country than the original genotype if it acquired some selective advantage. However, it remains striking to see how efficiently recombinant viruses have spread among different population groups in different geographic regions (suggesting that they could have a better viral fit than the parental nonrecombinant strains). In Thailand, where subtypes E (an A/E recombinant) and B were initially introduced, subtype E became predominant (13,37). In China, subtype B and C viruses have been introduced, and a recombinant B/C virus is spreading now through various parts of the country. Several reports documented the introduction of subtypes A and B among intravenous drug users in Kalingrad, Russia, and a recombinant A/B virus is rapidly spreading there (39,40). In West Africa, subtype A is predominant with prevalences ranging from 70% to >85% of the strains circulating (15) and, as shown in this study, AG-IBNG viruses comprise most envelope subtype A viruses in different countries.

Recombinant viruses are already contributing substan-

**TABLE 4. Estimation of the numbers and percentages of recombinant HIV-1 viruses circulating in West Africa**

<table>
<thead>
<tr>
<th>Country</th>
<th>n tested</th>
<th>gag/env subtype n (%)</th>
<th>AG-IBNG n (%)</th>
<th>Total n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Senegal</td>
<td>44</td>
<td>5 (13.9%)</td>
<td>27 (61.4%)</td>
<td>33 (75.0%)</td>
</tr>
<tr>
<td>Cameroon</td>
<td>55</td>
<td>4 (7.2%)</td>
<td>24 (43.6%)</td>
<td>28 (50.9%)</td>
</tr>
<tr>
<td>Gabon</td>
<td>82</td>
<td>10 (12.1%)</td>
<td>38 (46.3%)</td>
<td>48 (58.3%)</td>
</tr>
<tr>
<td>Total</td>
<td>181</td>
<td>19 (10.5%)</td>
<td>89 (49.2%)</td>
<td>119 (65.7%)</td>
</tr>
</tbody>
</table>

JAIDS Journal of Acquired Immune Deficiency Syndromes, Vol. 23, No. 5, April 15, 2000
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...ially to the global pandemic, and the likelihood of generating recombinant viruses will only continue to increase as the different HIV-1 subtypes spread to all continents (25,41,42). In a previous study, we documented that even the complex AG-IBNG-like viruses are implicated in this part of the world can have implications for future vaccines, as well as diagnostic, and treatment strategies. Especially with the recent introduction of antiretroviral treatment in Africa, one must be aware that all the AG-IBNG-like viruses are subtype G in the protease gene, and recently a naturally occurring lower susceptibility for HIV-1 subtype to protease inhibitors has been shown (43). The current and future role of these viruses in the global pandemic must be monitored in all new molecular epidemiological studies and discrimination between subtype A and AG-IBNG-like viruses is necessary.

Acknowledgments: This work was cosponsored by grants from the European Union (INCO-DC No. IC18CT97-0216 and INCO-DC No. IC18-CT96-0110), and the Agence Nationale de Recherches sur le SIDA (ANRS, Projet Sidak). C. Montavon holds a doctoral fellowship from ANRS.

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