Reliability of Rapid Subtyping Tools Compared to That of Phylogenetic Analysis for Characterization of Human Immunodeficiency Virus Type 1 Non-B Subtypes and Recombinant Forms

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Received 17 March 2008/Returned for modification 17 May 2008/Accepted 30 September 2008

Human immunodeficiency virus type 1 (HIV-1) subtyping is often estimated on the basis of pol sequences by using online websites instead of phylogenetic analysis (phy). We evaluated the reliability of distinct rapid subtyping tools versus phy with a large panel of HIV-1 non-B subtypes and circulating recombinant forms (CRF). pol sequences (277 protease [PR] and 171 reverse transcriptase [RT] sequences) previously assigned by phy to eight distinct HIV-1 non-B subtypes were obtained from 277 HIV-infected patients. Phy was run again to identify CRF. Subtyping was then performed using three rapid tools (the Stanford, NCBI, and REGA online tools). Thirty-three additional clade B sequences were tested as controls. New phylogenetic analyses reclassified two-thirds of pol sequences previously assigned to HIV-1 non-B clades as CRF. CRF02_AG variants were correctly assigned by the Stanford and NCBI tools for 92 to 97% and 96 to 99% of PR-RT sequences, respectively, while they were correctly assigned by the REGA tool for only 18 to 32% of PR-RT sequences. The Stanford, NCBI, and REGA tools failed to assign pure non-B clades correctly for 24 to 33%, 35%, and 57 to 64% of PR-RT sequences, respectively. For PR-RT sequences from CRF other than CRF02_AG, discrepancies occurred in 98 to 100%, 18 to 43%, and 80 to 87% of sequences, respectively. The concordance between those tools and phy was almost complete for subtype B assignment. Rapid subtyping tools show relatively low agreement with phy in identifying HIV-1 non-B clades and CRF other than CRF02_AG. The Stanford tool shows the best concordance with phy for the assignment of pure non-B clades, while the NCBI tool performs better at identifying CRF. Before entering routine clinical use, rapid subtyping tools should be optimized and updated periodically. Larger numbers of different non-B subtypes and CRF sequences should be included.

Both the high error rate of reverse transcriptase (RT) and the occurrence of recombination events contribute to the expansion of the genetic heterogeneity of human immunodeficiency virus type 1 (HIV-1) in vivo (11). Phylogenetic analysis (phy) permits the identification within HIV-1 group M of nine subtypes (A, B, C, D, F, G, H, J, and K) (13), at least 43 major circulating recombinant forms (CRF) (http://www.hiv.lanl.gov/content/sequence/HIV/CRFs/CRFs.html), and multiple unique recombinant forms. CRF are defined as intersubtype recombinants for which at least three epidemiologically unlinked variants are monophyletic and share an identical genetic structure. Unique recombinant form variants are widely distributed worldwide, with recombination breakpoints different from those of CRF, and they have been found in fewer than three infected persons. Recombination events result from coinfection and/or superinfection of cells with different HIV variants; they occur especially in populations where multiple subtypes cocirculate, such as in sub-Saharan Africa (18). Among the HIV-1 recombinant viruses, the CRF02_AG variant accounts for more than half of the infections in West and Central Africa (16, 17). This variant infects several million people and is the predominant strain among the population from these African countries living in Southwest Europe but infected in their native countries. An increasing number of non-B subtypes have been reported throughout Europe and in areas of North America with large numbers of immigrants. Local transmission of non-B subtypes and CRF have already been documented (1, 5, 14). In Spain, almost all HIV-1 subtypes have been identified (7, 15), and multiple recombinants, mostly CRF02_AG variants, are currently in circulation (6, 8, 9).

Identification of non-B subtypes and CRF is important for confronting potential problems with the use of antiretroviral drugs and diagnostic tools, such as viral load and resistance tests, which have been designed mainly based on clade B sequences (3, 10, 19). Moreover, pathogenicity can differ across HIV-1 clades, and several studies have shown that subtype D may lead to more-rapid disease progression than other subtypes (2, 22). Thus, HIV-1 subtype characterization is becoming an important aspect of the adequate clinical management of HIV-infected persons (4).

Since phylogenetic analyses are not widely implemented in clinical settings due to their complexity, most clinicians perform HIV-1 subtyping by interpreting pol sequences obtained following drug resistance testing by using online websites instead of phy. Here we evaluate the reliability of the three rapid subtyping tools available (the Stanford, NCBI, and REGA tools).
tools) versus phy with a large panel of HIV-1 non-B subtypes and CRF from HIV-1-infected patients living in Spain.

MATERIALS AND METHODS

pol sequences (277 protease [PR] and 171 RT sequences) previously assigned to eight distinct HIV-1 non-B subtypes by phy (5, 7, 15) were obtained from 277 HIV-1-infected patients. Most of them were on regular follow-up at our institution, an HIV/AIDS and Tropical Medicine Reference Center. All patients were under epidemiological suspicion of carrying non-B clades, because they were either (i) foreigner coming from areas where non-B clades are endemic, (ii) natives who had traveled to countries where HIV-1 non-B subtypes are endemic and who admitted to having engaged in high-risk behaviors there, or (iii) natives who had sexual contacts with people coming from those areas.

The study population had a mean age of 40 years, and almost half (51.5%) were male. Among the patients in known risk groups, 89.5% and 1.9% had acquired HIV-1 infections through heterosexual and homosexual relationships, respectively, while 5.8% admitted intravenous drug use, 1.9% were infected by a vertical route, and 0.9% were infected by blood transfusion. This study was part of a project approved by a review board and the ethical committee from our institution. It was designed to protect the rights of all subjects involved under the appropriate local regulations. To maintain subject confidentiality, a unique identification number was assigned to each specimen.

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provided by the three rapid subtyping tools (Stanford, NCBI, and REGA). In more detail, subtyping by the three rapid tools was performed for 277 PR and 171 RT sequences derived from 277 HIV-1 infected subjects previously determined by phy to be infected by non-B subtypes. At the PR the subtypes were 27A, 2B, 16C, 5D, 4F, 32G, 4H, 1J, 2CRF01_AE, 141CRF02_AG, 2CRF03_AB, 11CRF06_cpx, 2CRF10_CD, 5CRF11_cpx, 9CRF12_BF, 1CRF13_cpx, and 13CRF14_BG. At the RT they were 18A, 7B, 9C, 2D, 3F, 20G, 1H, 81CRF02_AG, 5CRF06_cpx, 3CRF10_CD, 4CRF11_cpx, 7CRF12_BF, 1CRF13_cpx, and 10CRF14_BG (Table 1). Clade B sequences were derived from recombinant viruses showing distinct clade assignment at the PR and RT. In addition, 33 clade B sequences at both genes were tested as controls.

Assignment of subtype B sequences using rapid subtyping tools. All tested subtype B variants, which are the most prevalent in the United States and Western Europe, were correctly assigned by the Stanford and NCBI tools. Both methods detected all PR and RT clade B sequences derived from CRF, as well as all 33 PR and RT clade B sequences used in pure clade B controls (Table 1). In contrast, the REGA tool failed to identify one PR and two RT clade B sequences from CRF carrying sequences from another subtype at *pol* (Table 1). Moreover, 3 of the 33 pure clade B PR/RT control sequences were not assigned by REGA.

Assignment of CRF02_AG sequences using rapid subtyping tools. Among the CRF tested in this study, 76% (141/186) of PR and 73% (81/111) of RT sequences belonged to CRF02_AG variants (Table 1). CRF02_AG sequences were correctly identified by the Stanford and NCBI tools for 92 to 97% and 96 to 99% of the PR-RT sequences, respectively, but only for 18 to 32% of the sequences by the REGA tool (Table 1). Overall, the REGA tool did not provide subtype assignment for 79% of PR and 60.5% of RT sequences derived from CRF02_AG strains, as was shown by phy.

Assignment of pure non-B HIV-1 subtypes. The overall accuracy of the Stanford, NCBI, and REGA tools in the identification of pure non-B clades of HIV-1 and CRF other than CRF02_AG carrying sequences from distinct subtypes at *pol* was relatively low compared to that of phy. As shown in Table 1, rapid subtyping tools showed discrepant results when interpreting sequences from non-B subtypes and CRF defined by phy. The Stanford, NCBI, and REGA tools failed to correctly assign pure non-B clades for 24 to 33%, 35%, and 57 to 64% of PR-RT sequences, respectively. Thus, the Stanford database showed the best concordance with phy for the assignment of pure non-B clades.

Of note, the PR sequence belonging to clade J was detected only by the NCBI tool. The REGA tool showed a similar concordance with other rapid tools for the assignment of clade A at PR, identifying a higher number of subtype A RT sequences (8/18) than the other two methods (Table 1). However, the REGA tool did not assign any subtype for 194 (70%) PR sequences and 95 (55.5%) RT sequences, which were known to belong to CRF in nearly 70% of the PR and RT sequences not assigned a subtype (data not shown).

Assignment of CRF other than CRF02_AG by using rapid subtyping tools. Overall, rapid subtyping tools showed limitations in the assignment of CRF other than CRF02_AG variants for *pol* sequences (45 PR and 30 RT specimens) when phy was taken as the reference. For instance, the Stanford, NCBI, and REGA tools failed to assign CRF correctly in our study for 98 to 100%, 18 to 43%, and 80 to 87% of PR-RT sequences, respectively (Table 1). The NCBI tool demonstrated the best concordance with phy for these CRF variants. In more detail, the concordance between NCBI and phy was complete for CRF03_AB, CRF11_cpx, CRF13_cpx, and CRF14_BG at the PR and for CRF14_BG at the RT, as well as for 75% and 57% of CRF11_cpx and CRF12_BF sequences, respectively, at the RT. Finally, the NCBI tool provided the best concordance with phy for CRF06_cpx, with correct assignment for nearly 80% of PR sequences, while REGA was correct in only 18% and 40% of PR and RT sequences, respectively (Table 1).

**DISCUSSION**

HIV-1 subtype characterization is becoming an important aspect of the clinical management of infected persons. Thus, updated information about circulating HIV-1 variants in distinct countries is helpful for epidemiological purposes as well as for confronting potential problems with the use of diagnostic assays (3, 18), viral load measurements (19), drug resistance testing, and the prescription of antiretroviral drugs (10). Since phylogenetic analyses are not widely used in clinical settings, most clinicians base subtype characterization on the interpretation of *pol* sequences obtained following drug resistance testing by using online tools. Here we have performed the first comparison of the reliabilities of different rapid HIV-1 subtyping tools (the Stanford, NCBI, and REGA tools), taking as a reference the results obtained using phy by testing a relatively large group of clinical specimens obtained from patients infected with HIV-1 non-B subtypes and CRF.

The number of HIV-1 strains that have been fully sequenced has increased within the past few years, revealing a high prevalence of recombinant variants among strains previously considered to represent pure non-B subtypes, based on information derived from single viral coding regions. For instance, viruses carrying subtype G sequences at *pol* could be pure G subtypes or recombinants, and interestingly, 18 of the CRF described (http://www.hiv.lanl.gov/content/sequence/HIV/CRFs/CRFs.html) harbor clade G sequences in partial or complete *pol* coding regions. Therefore, it is not surprising that CRF may constitute at least 10 to 20% of new HIV-1 infections worldwide and that recombination involving different non-B subtypes is rising where diverse variants cocirculate, particularly in areas where distinct subtypes are highly prevalent (8, 12, 21, 24).

The concordance between rapid subtyping tools and phy for clade B viruses was excellent in this study. These results reinforce the good performance of rapid subtyping tools for this subset of patients. Our results also revealed that the Stanford database, the tool most widely used by clinicians for routine clinical characterization of HIV-1 subtypes, showed the best concordance with phy for the recognition of pure non-clade-B subtypes. However, the Stanford database failed to identify nearly 25% of PR/RT sequences belonging to non-B clades, as well as 98% of PR and all RT sequences belonging to CRF variants other than CRF02_AG. The NCBI tool showed the best concordance with phy for CRF identification, although it
failed to assign 35% of pure non-B subtypes correctly in our study. The REGA tool failed to assign either pure non-B subtypes or CRF correctly for an unexpectedly large number of samples. However, it was the only tool that identified CRF06_cpx variants, and it showed the best performance for identifying clade A sequences at the RT gene. Taken together, our results point out that misclassification of HIV-1 non-B subtypes and CRF strains other than CRF02_AG may occur frequently when rapid subtyping tools are used, a limitation that may have epidemiological and clinical consequences. However, the concordance between the Stanford tool, the NCBI tool, and phy was relatively good, with complete agreement for nearly two-thirds of pol sequences.

It is reassuring that CRF02_AG strains, which are spreading rapidly in West and Central Africa (16, 17), as well as in European countries with strong links to, and immigrants from, these regions, are well represented in subtyping databases. Correspondingly, rapid subtyping tools performed quite well with these sequences. However, CRF variants other than CRF02_AG were less well represented in these databases and often were misclassified by rapid subtyping tools. Since these strains are similarly spreading worldwide, it may be worth highlighting the importance of reporting these non-B sequences and stressing the need for improved accuracy of rapid subtyping tools for these variants. Other potential bioinformatics tools have been developed, or are in development, to deal with the unreliability of subtyping tools for correctly identifying HIV-1 subtypes, subsubtypes, CRF, and other HIV variants using viral sequences (20, 23).

In summary, this study revealed that HIV-1 rapid subtyping tools can be useful for identifying clade B sequences. However, phy of pol sequences continues to be the only reliable method for correctly assigning HIV-1 non-B subtypes and CRF, which are growing in number and complexity. Clearly, the number of sequences from non-B and CRF strains in reference databases must increase, and rapid tools should improve their prediction of subtyping for the distinct non-B variants, including CRF, before being used in routine clinical settings.

ACKNOWLEDGMENTS
This work was supported in part by grants from the Red de Investigación en SIDA (RIS, project ISCIII-RETIC RD06/006) and the Fondo de Investigaciones Sanitarias (FIS PI06/0925). A.H. is supported by FIS and the Agencia Lain Entralgo. The authors do not have any conflict of interest.

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