



HIV subtype diversity worldwide

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Purpose of review

To provide a summary of the current data on the global HIV subtype diversity and distribution by region. HIV is one of the most genetically diverse pathogens due to its high-mutation and recombination rates, large population size and rapid replication rate. This rapid evolutionary process has resulted in several HIV subtypes that are heterogeneously globally distributed.

Recent findings

Subtype A remains the most prevalent strain in parts of East Africa, Russia and former Soviet Union countries; subtype B in Europe, Americas and Oceania; subtype C in Southern Africa and India; CRF01_AE in Asia and CRF02_AG in Western Africa. Recent studies based on near full-length genome sequencing highlighted the growing importance of recombinant variants and subtype C viruses.

Summary

The dynamic change in HIV subtype distribution presents future challenges for diagnosis, treatment and vaccine design and development. An increase in recombinant viruses suggests that coinfection and superinfection by divergent HIV strains has become more common necessitating continuous surveillance to keep track of the viral diversity. Cheaper near full-length genome sequencing approaches are critical in improving HIV subtype estimations. However, missing subtype data and low sequence sampling levels are still a challenge in some geographical regions.

Video abstract

<http://links.lww.com/COHA/A14>.

Keywords

diversity, HIV, subtype

INTRODUCTION

As of 2017, there were 36.9 million people living with HIV worldwide with a majority of these infections in sub-Saharan Africa [1]. AIDS is a global pandemic caused by two genetically diverse lentiviruses, HIV-1 and HIV-2 [2], that were introduced through multiple cross-species transmissions of simian immunodeficiency viruses from nonhuman primates to humans. These distinct zoonotic viral transmissions gave rise to separate HIV-1 groups M (Major), O (Outlier), N (non-M, non-O) and the most recent group P [2]. The origin of HIV-1 has been documented around Kinshasa in the present day Democratic Republic of Congo around the 1920s from where it spread along a transport network to other areas in sub-Saharan Africa, West Africa, Europe and the rest of the world [3]. This global spread was marked by a geographically defined distribution of several genetically distinct viruses. For instance, subtype B became prevalent in almost all parts of Europe and the Americas while a variety of subtypes and intersubtype recombinants are found in Africa with the highest diversity reported in West Central Africa. Whereas group M viruses have dominated the

global HIV pandemic since its inception, other group N, O and P viruses have not been disseminated widely. Group M viruses are further subdivided into nine subtypes (A-D, F-H, J, K) [2]. The genetic distance within a subtype can be between 15 and 20%, whereas genetic distances across subtypes are usually 25 and 35% [4]. HIV-2 remains largely restricted to the

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KEY POINTS

- Subtypes A, C and CRF02-AG are the most prevalent on the African continent, whereas subtype B predominates in Europe/the Americas and CRF01_AE dominate in Asia.
- There is an increase in the prevalence of recombinant viruses particularly in geographical areas where multiple subtypes cocirculate such as Asia and Africa.
- The emerging and dynamic HIV genetic diversity needs constant monitoring due to the challenges it presents towards diagnosis, phylogenetic reconstruction, treatment and vaccine development.
- The classification of HIV variants based on near full-length genomes is still limited but needs to be scaled up in a cost-effective way.
- The problem of missing HIV subtype data and low sequence sampling levels needs to be addressed.

western part of Africa although viral introductions have been reported elsewhere in Europe (Portugal and France), India and the United States of America [5]. HIV-2 has also been documented to be less infectious than HIV-1 [6,7]; it is comprised of at least nine groups (formerly referred to as subtypes; A to I) of which groups A and D are presently circulating [5]. HIV-2 subtype data is still limited and only a few recombinants have been described [5,8].

CIRCULATING RECOMBINANT FORMS

Recombination is an important phenomenon for viral diversification that allows the virus to evade the host immune system and antiretroviral treatment [9]. A recombinant is a viral sequence that has regions from two or more distinct parental strains [10]. A circulating recombinant form or circulating recombinant form (CRF) is a combination of viral genomes of different subtypes in dually infected people that results in a mosaic genome composed of regions from each of the different subtypes. When these recombinants are transmitted and spread within a population, they are recognized as circulating strains in the HIV epidemic and are classified as CRFs [11]. The virus must be isolated from at least two unlinked individuals and fully sequenced [12]. The designation 'cpx' (complex) is used when more than three subtypes are present [13]. To date, 98 CRFs have been identified and updated in the Los Alamos National Laboratory HIV sequence database [14]. These are designated with sequential numbering as they are discovered and reported.

UNIQUE RECOMBINANT FORMS

Unique recombinant forms (URFs) consist of a mixture of subtypes, but unlike the CRFs, they were sampled only once from a single multiply-infected individual [13]. Hence, all sequences corresponding to a CRF have the same recombination break points in the genome while each URF shows unique break-points. Intrasubtype recombination does occur but as it does not lead to subtype recombination, these viruses are not described as URFs. Dual infections with HIV-1 and HIV-2 have frequently been reported in regions where both viruses circulate [15] although no recombinants between them have been described. Several AD and AC intersubtype recombinants have been described in Eastern Africa [16,17], where these three subtypes cocirculate. BF intersubtype recombinants have been found in Brazil and Argentina, where subtypes B and F are both common [18,19]. Subtypes A and C cocirculate in India with AC recombinants [20].

LITERATURE SEARCH

We performed a structured literature search in PubMed using MeSH terms. A combination of keywords related to the subject of 'HIV genetic diversity worldwide' were used with Boolean operators (OR and AND) in PubMed's advanced search builder. The results generated in the search builder at each level were combined to output all articles that overlapped across all the search words for the specified review period (Fig. 1). This article focused mainly on HIV-1 as there was limited data on HIV-2 during the review period.

GLOBAL DISTRIBUTION OF HIV-1 BY REGION

Africa

The greatest genetic diversity of HIV-1 has been found in Africa especially in West Central Africa [21]; however, other parts of the continent show an assortment of diverse viral strains. In Southern Africa, subtype C collectively predominates [22,23], whereas in Eastern Africa subtype A is the most dominant [24²²] although other subtype D and C viruses cocirculate as well [25]. An increase in subtype C and a decrease in subtype D have been reported in Kenya together with extensive intersubtype recombination [24²²]. A recent study in Uganda reported an increased prevalence of HIV-1 intersubtype recombinants of about 46% based on near full length HIV sequences [26²²]. A separate study by the Phylogenetic And Networks for Generalized HIV Epidemics in Africa (PANGEA)-HIV consortium [27²²] showed a high proportion (about 50%) of

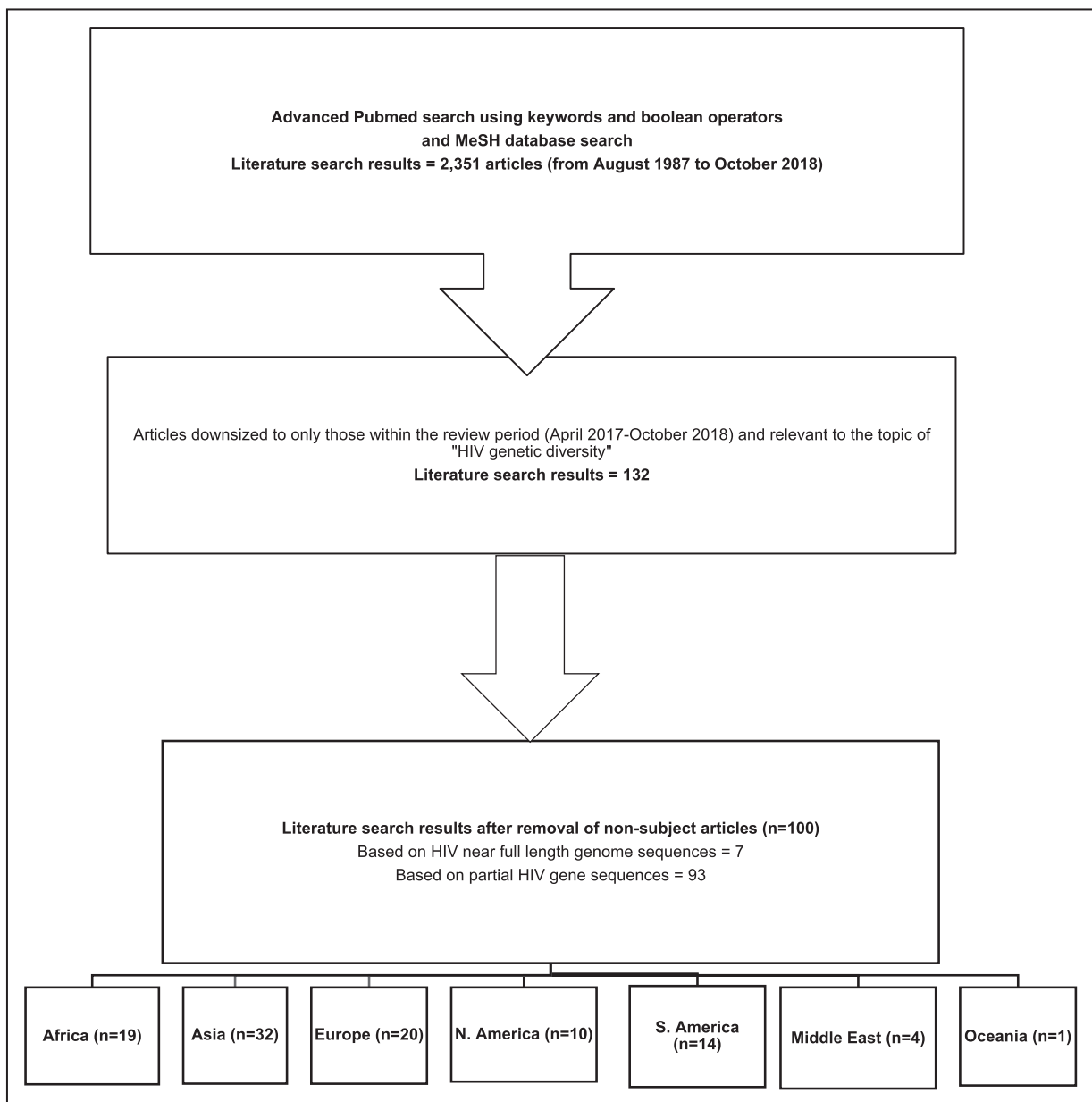


FIGURE 1. Literature search. Diagrammatic representation of the literature search and selection of articles.

HIV-1 intersubtype recombinants in Uganda based on deep sequencing of near full-length viral genomes (unpublished data). In West and West Central Africa, the majority of viruses are CRF02_AG [21,28]. Subtypes A and G cocirculate in some countries such as Nigeria [21]. Mosaic viruses involving CRF02_AG have been reported in several African countries [29] and an increase of subtype F2 and other recombinant forms in Cameroon [30] (Fig. 2).

Europe and North America

In North America [31], Europe [32–36] and Australia [37], subtype B is still the most widespread viral

strain. Other subtypes have also been introduced. For example, in Spain, subtype F1 was identified in a cohort of MSM [38] or CRF02_AG among immigrant populations [39].

South America

Subtype B predominates in South America [40,41]. Other intersubtype BF recombinants have also been recently reported [42]. An increase in pure subtype C compared with subtype B has also been reported among HIV/viral hepatitis coinfecting patients in Southern Brazil [43].

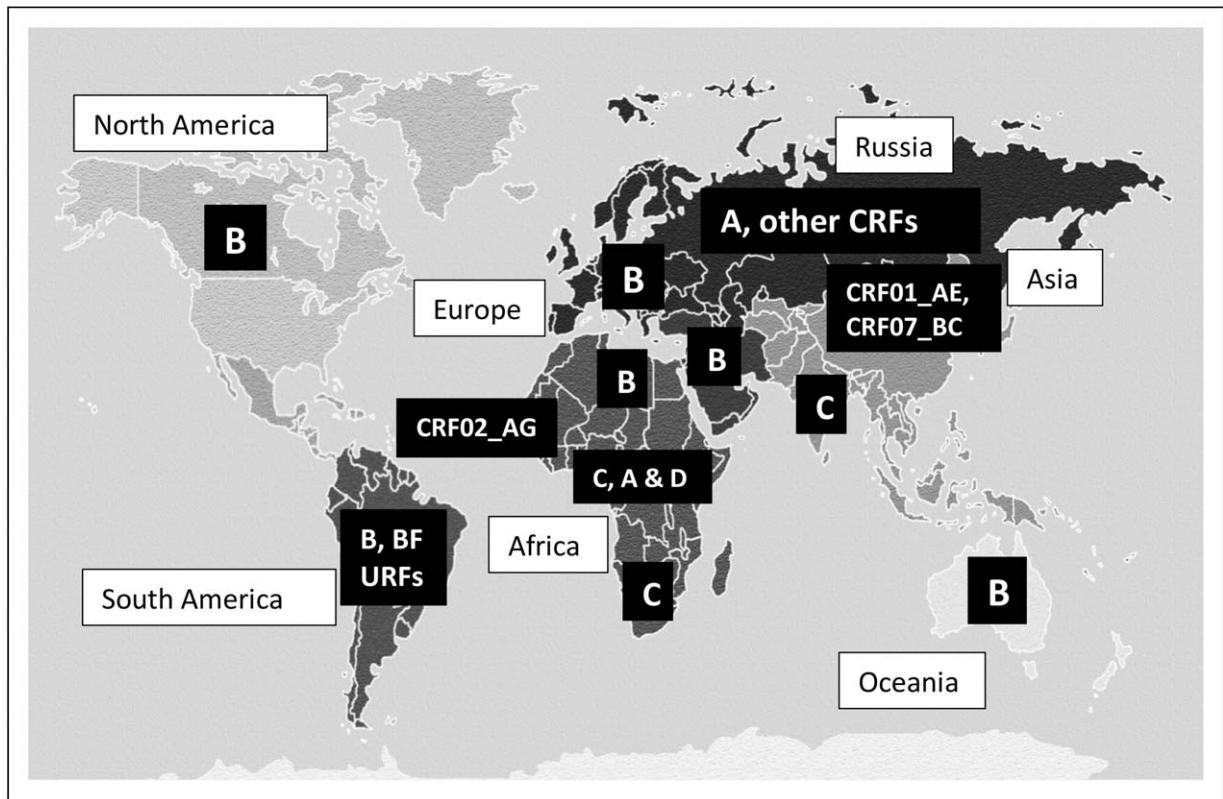


FIGURE 2. Global distribution of major HIV subtypes. Map showing the global distribution of the major HIV subtypes and circulating recombinant forms from the review. Source: Map of the world modified to show HIV-1 subtype diversity worldwide. Map-menu.com.

Asia

Several subtypes circulate in Asia and it has been described as the ‘hotbed’ of recombinant viruses with several CRFs [44]. Subtype A predominates in Russia and other former Soviet Union countries although CRF02_AG is common in Kyrgyzstan [45]. CRF01_AE and CRF07_BC predominate in China [46–48]; new CRFs have been found in China more than in any other country [49–52]. In south-east Asia, CRF01-AE is the most prevalent [53,54] while subtype C predominates in India with reports of increased URFs in the north eastern part of the country [55].

Middle East and North Africa

Middle East and North Africa countries are grouped together because subtype B is the predominant subtype in the region and these countries show relatively low prevalence and limited sampling of infections [56^a,57].

CONCLUSION

Previous reviews have shown that subtype C accounts for more than 50% of all infections worldwide with a concentration in East and Southern

Africa and India, CRF01_AE as the major strain throughout Southeast Asia, CRF02_AG as the dominant virus in West Africa and about 30% of strains in East Africa to consist of URFs [13,21]. However, there is increased and extensive HIV-1 genetic diversity in Asia [47,49–52,58] and Eastern Africa [24^{aa},26^{aa}]. By 2010, there were about 45 reported CRFs [59], this figure has more than doubled within a 7-year period with at least 98 CRFs that have currently been described [14]. This fast rate of increase in the prevalence of CRFs and the fact that several subtypes cocirculate globally suggests that these numbers will likely continue to grow in the future. The largest source of HIV sequence data (subtype B) is from Europe and the Americas with less data coming from the Middle East (Table 1). As a result, HIV sequence databases like Los Alamos [60] tend to represent a biased sampling and therefore do not necessarily reflect the distribution of subtypes worldwide. For instance, subtype B accounted for 55% of HIV sequences in the Los Alamos database [60] which is about five times its frequency in the world [61]. Information on subtype diversity based on near full-length genomes is still relatively scarce with majority of the available data still based on partial HIV fragments. In East Africa, classification of HIV based on near full-length sequences showed a prevalence

Table 1. Global prevalence and geographical distribution of HIV-1 subtypes and recombinant forms by studies reviewed

Major subtype	Global Prevalence based on the level of geographic spread and available data in the HIV sequence database	Description	Countries	Geographical location	Study (year)	HIV genomic region/available sequences
A	High	Includes intrasubtypes like A1 and A6	Kenya, Pakistan, Armenia, Azerbaijan, Belarus, Estonia, Georgia, Kazakhstan, Kyrgyzstan, Latvia, Lithuania, Moldova, Russia, Ukraine and Uzbekistan	East Africa, Asia, EuroAsia	Gounder <i>et al.</i> (2017), Khan <i>et al.</i> (2018), Aibekova <i>et al.</i> (2018), Iapovok <i>et al.</i> (2017), Kazennova <i>et al.</i> (2017)	Gag and env (n = 386); gag, pol and env (n = 755); pol (n = 3022), integrase (n = 506)
B	High	Includes reports of other multiple nonsubtype B virus importations	United states of America, Canada, France, United Kingdom, Belgium, Bulgaria, Cyprus, Spain, Iceland, Mexico, Brazil, Guatemala, Honduras, Panama, Nicaragua, Belize, El Salvador, Amazonas, Australia, Middle East and North Africa that include Algeria, Bahrain, Djibouti, Egypt, Iran, Iraq, Jordan, Kuwait, Lebanon, Libya, Morocco, Palestinian Territories, Oman, Qatar, Saudi Arabia, Somalia, Sudan, Syria, Tunisia, United Arab Emirates, and Yemen	North America, South America, Australia, Europe	Dapp <i>et al.</i> (2017), Pérez-Losada <i>et al.</i> (2017), Vrancken <i>et al.</i> (2017), Dennis <i>et al.</i> (2017), Oster <i>et al.</i> (2017), Tumiotto <i>et al.</i> (2018), Volz <i>et al.</i> (2018), Hebberecht <i>et al.</i> (2018), Alexiev <i>et al.</i> (2018), Yebra <i>et al.</i> (2018), Pineda-Peña <i>et al.</i> (2018), Rojas Sánchez <i>et al.</i> (2017), Sallam <i>et al.</i> (2017), Hernandez-Sanchez <i>et al.</i> (2018), Alves <i>et al.</i> (2017), Martin <i>et al.</i> (2017), Chaillon <i>et al.</i> (2017), Villanova <i>et al.</i> (2017), Delatorre <i>et al.</i> (2017), Andrade <i>et al.</i> (2017), Barral <i>et al.</i> (2017), Machado <i>et al.</i> (2017), Lima <i>et al.</i> (2017), Casiley <i>et al.</i> (2017), Sallam <i>et al.</i> (2017), Daw <i>et al.</i> (2017)	Gag (n = 1790); env (n = 1755); pol and integrase (n = 63 917); pol (n = 23 196), Near full length (n = 450), gag & pol (n = 66), vif (n = 317)
C	High	Consists of half of all HIV infections worldwide. Higher prevalence has been found among coinfecting patients in Brazil	South Africa, Tanzania, India, Southern Brazil	Southern Africa, East Africa, Asia	Sivay <i>et al.</i> (2018), Ngcapu <i>et al.</i> (2017), Ngandu <i>et al.</i> (2017), Bauer <i>et al.</i> (2017), Naidoo <i>et al.</i> (2017), Billings <i>et al.</i> (2017), Sharma <i>et al.</i> (2018), Sharma <i>et al.</i> (2017), Avanzi <i>et al.</i> (2017)	pol (n = 200); RNase H (n = 140); gag (n = 91); gag-protease (n = 28); Near full length (n = 170), vpu (n = 80); nef (n = 57); pol/tat-vpu-env (n = 110); pol and env (n = 38)
A/D	Moderate	Intersubtype recombinants of subtypes A1 and D	Uganda	East Africa	Lee <i>et al.</i> (2017)	Near full length (n = 200)

Table 1 (Continued)

Major subtype	Global Prevalence based on the level of geographic spread and available data in the HIV sequence database	Description	Countries	Geographical location	Study (year)	HIV genomic region/available sequences
F1 and B/F recombinants	Low	Increasing prevalence among MSM in Spain and Argentina	Spain, Argentina	Europe, South America	Patiño-Galindo <i>et al.</i> (2017), Cevallos <i>et al.</i> (2017)	Near full length (n=49), Near full length (n=25), pol (n=312)
CRF01_AE	High	Intersubtype recombinant of subtypes A and E. Previously classified as subtype E	Indonesia, China, Vietnam, Myanmar	Asia	Ueda <i>et al.</i> (2018), Li <i>et al.</i> (2018), Xu <i>et al.</i> (2018), Chen <i>et al.</i> (2018), Chikata <i>et al.</i> (2018), Yang <i>et al.</i> (2018), Zhang <i>et al.</i> (2017), Sun <i>et al.</i> (2017), Li <i>et al.</i> (2017)	pol (n=837); Near full length (n=335); partial gag, pol and env (n=638); gag, pol and nef (n=388), env and gag (n=70); gag+RT (n=187)
CRF02_AG	High	Intersubtype recombinants of subtype A and G although may show more extensive diversity	Ghana, Cameroon	West Africa	Nii-Trebi <i>et al.</i> (2017), Nanfack <i>et al.</i> (2017), Rodgers <i>et al.</i> (2017)	pol (n=99); pol & env (n=555); NFL (n=1)
CRF07_BC	Moderate	Intersubtype recombinants of subtypes B and C	China	Asia	Yang <i>et al.</i> (2017), Zhang <i>et al.</i> (2017), Wang <i>et al.</i> (2017), Li <i>et al.</i> (2017)	pol (n=276); gag (n=610); Near full length (n=1); gp160 (n=749)
CRF08_BC	Low	Recombinant between subtypes B and C and commonly cocirculate with CRF07_BC and CRF01_AE in parts of Yunnan, China	China	Asia	Chen <i>et al.</i> (2018)	gag, pol and env (n=291)
CRF35_AD	Low	Recombinant between subtypes A and D	Iran (Tehran)	Middle East	Vahabpour <i>et al.</i> (2017)	pol (n=42)
Mixed subtypes and new CRFs or URFs	Low	CRF01_AE, CRF02_BC, subtypes A-H, J and K. Also include newly-recommended strains like CRF55_01B, CRF67_01B and CRF69_01B, CRF96_cpx, CRF01/CRF07_BC, CRF07_BC/CRF55_01B, CRF87_cpx/CRF88_BC	Several countries within congo Basin, China, African countries, Canada, South Africa, Uganda and Kenya	Central and West Africa, N. America, South and East Africa	Tongo <i>et al.</i> (2018) and Tongo <i>et al.</i> (2018), Fan <i>et al.</i> (2018), Miao <i>et al.</i> (2018), Lu <i>et al.</i> (2018), Kong <i>et al.</i> (2017), Li <i>et al.</i> (2017), Wu <i>et al.</i> (2017), Hu <i>et al.</i> (2017), Kiguoya <i>et al.</i> (2017)	Near full length (n=305); pol (n=18); env (n=48); gag/pol/env (n=183), Gag-protease (n=1342)

CRF, circulating recombinant form; NFL, near full-length; RT, reverse transcriptase; URF, unique recombinant form.

of 46% of recombinant viruses in Uganda [26^{***}]. The classification of HIV variants based on near full-length genomes is still relatively low (Table 1) although some consortia have generated data that spans entire HIV genomes [62]. One such consortium is the PANGEA HIV [27^{**}] which is a multinational project funded by the Bill and Melinda Gates Foundation to study the transmission dynamics of HIV within generalized epidemics in Africa. PANGEA aimed at generating close to 20 000 HIV near full-length sequences mainly from sub-Saharan Africa which have been valuable in characterizing HIV variants and understanding the subtype diversity in several African countries. Preliminary results have reported close to 50% of HIV-1 intersubtype recombinants in Uganda based on near full-length genomes (unpublished data from PANGEA-HIV, [27^{**}]).

Recent data suggest a significant increase in the prevalence of recombinants in some countries [26^{***},55,63^{***}] and several newly discovered CRFs/URFs in parts of Asia and Africa [49–52,58,59]. This suggests that coinfection or superinfection by divergent HIV-1 strains has become more common in regions where multiple subtypes cocirculate [26^{***},42,49,52,64]. Viral sequences from recombinant forms have been shown to affect the accuracy of phylogenetic reconstructions [65,66]. This is due to the different regions of the aligned recombinant genome having distinct evolutionary relationships that affect tree topology and branch lengths [67]. Furthermore, such sequences introduce errors in phylodynamic inferences [68] by biasing ancestral state reconstruction and estimations for the most recent common ancestor [65]. As the prevalence of recombinant forms increases, these challenges are expected to become more important for molecular epidemiological-based investigations. The scale up of cheaper near full-length genome sequencing approaches is critical in improving the accuracy of viral classification. It is important to consider how the fast-increasing emergence of HIV recombinant forms will impact accurate diagnosis, phylogenetic reconstruction, antiretroviral treatment and future vaccine development.

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Conflicts of interest

There are no conflicts of interest.

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- of outstanding interest

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