Transmitted HIV drug resistance and subtype patterns among blood donors in Poland

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Surveillance on the HIV molecular variability, risk of drug resistance transmission and evolution of novel viral variants among blood donors remains an understudied aspect of hemovigilance. This nationwide study analyses patterns of HIV diversity and transmitted resistance mutations. Study included 185 samples from the first time and repeat blood donors with HIV infection identified by molecular assay. HIV protease, reverse transcriptase and integrase were sequenced using population methods. Drug resistance mutation (DRM) patterns were analyzed based on the Stanford Interpretation Algorithm and standardized lists of transmitted mutations. Phylogeny was used to investigate subtyping, clustering and recombination patterns. HIV-1 subtype B (89.2%) followed by subtype A6 (7.6%) were predominant, while in three (1.6%) cases, novel recombinant B/A6 variants were identified. Non-B variants were more common among repeat donors (14.5%) compared to the first time ones (1.8%), \( p = 0.011 \), with higher frequency (9.9%) of A6 variant in the repeat donor group, \( p = 0.04 \). Major NRTI DRMs were observed in 3.8%, NNRTI and PI in 0.6% and INSTI 1.1% of cases. Additionally, E157Q polymorphism was observed in 9.8% and L74I in 11.5% of integrase sequences. Transmission of drug resistance among blood donors remains infrequent. Subtype patterns increase in complexity with emergence of novel intersubtype A6B recombinants.

Molecular surveillance allows to model the evolution of the HIV epidemics, while investigation of the drug resistance, subtype variability and identification of the new recombinant variants remains a valuable tool to inform public health strategies1-2. Monitoring of the drug resistance patterns, especially in the evolving landscape of the antiretroviral drugs, rapid antiretroviral treatment introduction and expanding use of the pre-exposure prophylaxis remains key to maintain virologic efficacy of combined antiretroviral treatment (cART) and preserve future treatment options3-4. Studies on the transmission of the HIV drug resistant strains have provided data to prioritize the selection of the first and second-line antiretroviral drugs5. Importantly, molecular characteristics of the HIV strains observed among blood donors may reflect circulation patterns in the untested general population, generally unaware of the transmission risk, which in turn is of the primary importance to the blood safety6. Additionally, amplification efficiency and sensitivity of the molecular assays used for the blood screening may differ for novel variants and resistance mutations7-11.

In Poland, HIV epidemics has evolved to the primarily sexually transmitted, with the highest transmission risk among men-who-have-sex with men and increasing number of new infections despite generally low level prevalence12. Since the beginning of epidemics until the end of 2019 HIV was diagnosed in 25 544 cases with 1429 registered AIDS deaths. Of note, in the recent years increasing number of immigrant population from the regions with the expanding HIV epidemics (Ukraine, Russia and Belarus) have been registered in the country however, records on the nationality among newly diagnosed individuals are commonly missing. So far, the most prevalent HIV-1 strain was subtype B (86.9%) followed subtype A (5.2%), D (3.5%) and C (1.8%) and infrequent identification of the recombinant forms (~ 2.1%)13. Among treatment naïve cases transmitted drug resistance mutations (DRM) were observed in ~ 9% of cases in the national wide studies (5.8% for nucleoside reverse transcriptase, 1.2% for non-nucleoside reverse transcriptase, and 2.0% for protease mutations) and up to ~ 12% in local cohorts with no transmitted major integrase mutations observed so far14-16. Additionally, frequency of transmission clusters ranges from 20–30% which indicates high likelihood of HIV epidemics expansion in the future17,18.

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Among blood donors the frequency of detected HIV infection in Europe remains low, however constant hemovigilance is required to identify the breakthrough transmissions in the pre-seroconversion period\(^{9-11}\). In general, currently implemented blood testing strategy based on nucleic acid amplification testing has proven to be effective\(^9\). The average frequency of HIV detection per 100,000 (confidential limit, 95% CI) among Polish blood donors in period 2005–2018 was 6.26 (5.77–6.85) for seropositive and 0.28 (0.19–0.42) for seronegative infections. In total the frequency of infected donors and donations per 100,000 was 6.56 (6.03–7.14) and 3.3 (3.04–3.59), respectively. In this period a slight increase in infection rate was noticeable and the HIV infection rate was significantly higher among the first time than in repeat blood donors. All seronegative HIV-NAT positive donors were men and all but one were repeat donors, however until now Look-back procedure has not documented any HIV transmission via blood component transfusion. Mathematical models estimate the risk of infectious transmissions at 0.16 to 0.49 per million, depending on screening format (sensitivity) and type of blood component (plasma volume)\(^9\).

In this considerably sized nationwide study, HIV diversity, patterns of transmitted drug resistance and clustering among Polish blood donors were analysed. This is the first study informing on the molecular variability of HIV in this group, identifying risk of DRM transmission and evolution of novel viral variants, which is of the utmost importance for the blood services.

Methods

Study group. For this study 235 samples with HIV infection confirmed by the molecular test, diagnosed during blood donation, were collected, of these 185 (78.7%) samples were successfully sequenced and included in the analyzed dataset. Sampling period spanned from 2009 to 2017 and included samples collected from the entire Poland, with positive samples obtained from the following blood collection centers: Białystok, Bydgoszcz, Gdańsk, Kalisz, Katowice, Kielce, Kraków, Lublin, Łódź, Olsztyn, Opole, Poznań, Racibórz, Radom, Rzeszów, Słupsk, Szczecin, Wałbrzych, Warszawa, Wrocław, Zielona Góra and military blood transfusion service. In this period in total 387 blood donors with HIV positive nucleic acid amplification test (NAAT) were identified (370 donors with detectable both HIV-RNA and anti-HIV antibody and 17 in the pre-seroconverted period with negative HIV serology, supplemental Table 1); therefore the study represents 60.7% of the total number of HIV NAAT positive blood donors. For HIV serological markers screening in blood donors 3rd or 4th generation CE-marked assays were used. All blood donations were tested in individual donations (IDT) with TMA based assays: initially with Procleix Ultrio Plus and later with Procleix Ultrio Elite (Gen-probe, USA) or with real-time PCR in minipools of 6 donations (MP6) using MPX system (Roche, USA). Donations reactive in screening were shipped to the reference laboratory at the Institute of Haematology and Transfusion Medicine in Warsaw for confirmatory tests that included Western/immuno blot and RNA HIV testing. Western/immuno blot analyses were performed using commercial assays that were changed periodically: HIV BLOT, Genelabs’ Diagnostics (Singapore); INNO-LIA™ HIV I/II Score, Innogenetics (Belgium). HIV BLOT MP Diagnostics (Singapore). NAAT test used were changed over the years to reflect technological progress and increasing sensitivity: Firstly, Cobas Ampliscreen HIV-1 v 1.5 Roche and Procleix Ultrio Plus were used, then Gen-probe USA/ and later Procleix Ultrio Elite, Gen-probe USA/ Confirmatory PCR Kit HIV-1 v 1.2 GFE Blut Germany. Newer assays were able to detect 2–3 HIV genome regions; each sample was confirmed using at least two methodologies, as noted above.

The study was approved by the bioethical committee of Pomeranian Medical University, Szczecin, Poland (approval number BN-001/34/04). The research was conducted in accordance with the Declaration of Helsinki. All data were anonymized. At the time of consent for the blood collection procedure the participant provides an informed consent related to the molecular analyses of the presence of the viral pathogens and subsequent analyses including molecular epidemiology of the viruses. Such an informed consent was obtained from all subjects included in the study. HIV-1 viral loads were not available, as the screening in blood collection centers is performed using qualitative HIV assay.

Data collected included type of donation (first time vs. repeated), gender, age, nationality, time since last donation (if repeated donor) and HIV infection (Fiebig) stage\(^{20,21}\). Fiebig stage was assessed based on the HIV molecular markers (HIV-RNA), p24 antigen, enzyme immunoassay reactivity and Western-blot patterns.

Before donation, prospective donors were screened for high-risk activities through a predonation, paper-based questionnaire. A brief physical examination was also performed. This procedure allows to determine whether the candidate is suitable for donating blood. Based on this evaluation, prospective donors could be temporarily or permanently deferred from donating blood. Donors also were deferred based reported certain risky behaviors including high-risk sexual activity (sex with multiple partners or with unknown partner (-s), having a sexual partner who injects drugs, commercial sex work, reported contact with person infected with HIV, HB, HCV or and Treponema pallidium), a history of criminal arrests or detention, intravenous drug use, exposure to blood from another person, selected medical (surgery, transplantations, gastroscopy etc.) or cosmetical (piercing, tattoo etc.) procedures. There were no HCV or HBV coinfected individuals in the analyzed group.

Sequencing. HIV-1 protease (PR) and reverse transcriptase (RT) genotyping and sequence assembly was performed using Virosol 2.9 genotyping kit (Abbott Molecular, Abbott Park, IL) according to manufacturer’s protocol providing a sequence of 1302 base pair (b.p.) long with inclusion of 1–99 codons in the PR and 1–335 in the RT. Additionally, HIV-1 integrase (IN) region (866 b.p., codons 1–288) was amplified and sequenced with reagents and conditions specified by Laethem et al.\(^{22}\). Amplicons obtained by the nested PCR method were used for sequencing by standard techniques with BigDye technology on an ABI 3500 platform (Applied Biosystems, Foster City, CA). Integrate sequence assembly was performed with the Recall online tool\(^{22}\). PR/RT sequencing
was successful for 159 (85.9%) cases, integrate for 174 (94.1%) samples. Final dataset included 148 (78.9%) patients with both PR/RT and IN sequence, 11 (5.9%) only with PR/RT and 26 (14.1%) only with IN sequence.

**Subtyping and drug resistance interpretation.** Initial subtyping was performed using automated genotyping software (REGA genotyping 3.46 tool34, based on the obtained PR, RT and IN sequences. Subtyping, including identification of subgroups for the subtype A, was verified using phylogenetic methods with reference sequences with known subtype from the 2018 version of the HIV sequence compendium (Los Alamos National Laboratory) (https://hivdb.lanl.gov/content/sequence/HIV/COMPRENDIUM/compendium.html), supplemented with local sequences from the HIV sequence database (http://www.hiv.lanl.gov/components/sequence/HIV/search/search.html). Phylogenetic trees of A-clade were made with a representative number of 178 sequences, comprising 101 references from the LANL-HIV database, 58 unique regional sequences with homology over 95% (based on BLAST analysis), and finally subtype O—as an out-group sequence. Method of maximum likelihood (ML) with approximate likelihood ratio test (aLRT) and Shimodaira–Hasegawa (SH) algorithm was performed among the support of PHYMLv3.0 web server.

For identification of drug resistance mutations Stanford Genotypic Resistance Interpretation Algorithm (https://hivdb.stanford.edu/hivdb/sequences/) was used, with classification of drug resistance mutations into major and accessory for PR and IN, as well as nucleoside and non-nucleoside inhibitor drug resistance (NRTI and NNRTI) for RT. Mutations with the scoring ≥ 10 for at least one active drug were included in the analyses. Additionally, PR/RT mutations were assessed according to WHO surveillance list35, while for integrase strand transfer inhibitor mutations standardized list of INSTI-resistance mutations was used36. In the final analyses we have also included the L74M integrase polymorphism as included in the IAS 2019 drug resistance update37.

**Phylogenetic analyses.** For the phylogenetic relationships, the PR/RT and IN sequences were concatenated (2168 bp length) and aligned with Clustal Omega28 software separately for subtype A and B. Subtype C and URFs were excluded from the phylogenetic analyses due to small sample size. Methodology to use concatenated sequences spanning different locations in HIV genome was used previously in numerous studies29–31 and HIV subtyping program (https://hivdb.stanford.edu/page/hiv-subtyper). Following the alignment, the optimal tree model was estimated using jModelTest 2.1.10 software for subtype A and subtype B sequences32. In both cases, the best fitting model was the GTR with four gamma categories. Rate parameters were as follows, for subtype A and subtype B, respectively: freqA = (0.3975/0.4221), freqC = (0.1581/0.1577), freqG = (0.2240/0.2077), freqT = (0.2205/0.2125), gamma shape parameter 0.4960/0.8810) and proportion of invariant sites of 0.3180/0.4850. Under these parameters three separate Bayesian Monte Carlo Markov Chain analyses were run in triplicates for 50 million of generations in Beast v.2.033 using Yuole Model with a strict molecular clock using freqT = (0.2205/0.2125), gamma shape parameter 0.4960/0.8810) and proportion of invariant sites of 0.3180/0.4850. All trees were visualized in Figtree v.1.4.4.

**Statistics.** Statistical comparisons were performed using Fisher’s exact and Chi² tests for nominal variables as appropriate. Continuous variables were analysed using the Mann–Whitney U-test for nonparametric statistics. Confidence intervals (CI) and interquartile ranges (IQR) were indicated where appropriate. Commercial software (Statistica 11.0PL, Statasoft, Warsaw, Poland) was used for these statistical calculations.

**Sequence data.** Sequences from this study have been submitted to GenBank and may be accessed with the following IDs: MZ218761 - MZ218932 and MZ218933 - MZ219089.

**Results**

**Overall group characteristics and HIV-1 subtypes.** The studied group included predominantly male individuals (95.7%) with the median age of 29 (IQR: 24–34) years. Of these, 54 (29.2%) individuals were the first time, while 131 (70.8%) were the repeat donors. The most prevalent HIV-1 variant was subtype B (n = 165, 89.2%) followed by subtype A (n = 15, 8.2%) and subtype C (n = 2, 0.1%) (Table 1). Of note, when utilizing the REGA 3.46 on-line automated subtyping tool all subtype A sequences were assigned as A1 variant, while phylogeny with reference sequences confirmed that in fact only one sequence belongs to the A1 subgroup, while the remaining sequences (n = 14, 7.6%) are in fact A6 subtype (Fig. 1). In three cases, novel recombinant variants with breakpoints between the reverse transcriptase and integrase coding region, were found and confirmed phylogenetically (2 sequences with B/A6 and one with A6/B) (Fig. 2).

In the group of the recurrent donors median time from the last donation was 226 (IQR: 121–654) days (median 169 (IQR: 68–315) days for the Fiebig I stage, median 93 (IQR:66–121) for II, median 81 (IQR: 42–111) for IV, median: 152 (88–333) for V and median 420 (165–1025) for VI. Median time from the last donation was notably longer for individuals with HIV diagnosed in Fiebig stages I-IV [median time 126 (IQR:66–169) days compared to stages V and VI [median time 241 (IQR: 136–730) days] (p = 0.003, and Fiebig stages I-V [median time 139 (IQR:79–304)] days vs VI (p < 0.0001).

There was a notable difference in the distribution of the Fiebig stages between the first time and recurrent donors (p = 0.03, Table 1), with Fiebig stage I-IV being more common among repeat donors (n = 13, 9.9%) vs. first time donors (n = 1, 1.9%), p = 0.047. Additionally, stages I-V were observed among 40.5% (n = 53) repeat donors vs. 16.7% (n = 9) first time donors (p = 0.002).
Interestingly, HIV-1 non-B variants were notably more common among repeat donors (n = 19, 14.5%) compared to the first time ones (n = 1, 1.8%), \( p = 0.011 \), which was associated with higher frequency of A6 variant in the repeat donor group (n = 13, 9.9%) \( p = 0.04 \).

**HIV-1 drug mutation patterns.** Based on the Stanford Genotypic Resistance Interpretation Algorithm major or non-accessory NRTI drug resistance was observed in 6 (3.8%) sequences, PI or NNRTI resistance in one (0.6%) case each, and INSTI in two (1.1%) sequences. Additionally, accessory mutations were observed in 5 (3.1%) PI, 17 (10.7%) NNRTI and 20 (11.5%) IN sequences (Fig. 3a–d).

Transmitted drug resistance mutations (tDRM), based on WHO mutation list, were observed in one (0.6%) protease (L54M mutation), 6 (3.8%) NRTI (mutations in the codon positions: M41L, D67N, T215S/V , K219Q) and two (1.1%) (both E138K mutations) integrase sequences (supplemental table 2). No NNRTI tDRM were observed. E157Q polymorphism was the most prevalent (9.8%) resistance associated variant in the analysed dataset and was notably more common among female individuals (n = 4, 50.0%) compared to 8.4% (n = 14) among males, \( p = 0.004 \). Additionally, we have analyzed the distribution of the L74I polymorphism which was present in virtually all A6 sequences it was also observed within the identified clusters. In subtype B one cluster with NNRTI K101H/E138A mutation was observed, in four sequence pairs there was also evidence of the shared resistance patterns (NRTI: D67N/K291Q and T215V , NNRTI: V106I, integrase: E157Q).

**Phylogenetic analyses.** Clustering was assessed using Bayesian inference, with genetic distance of 1.5%, separately for subtype A and B with 9 (50%) and 44 (26.2% ) sequences, respectively contained within transmission clusters. There were 16 sequence pairs and 3 (two containing 3 sequences, one with 6 sequences) clusters identified for subtype B and three (all with 3 A6 sequences) for subtype A (Fig. 4, supplemental Fig. 1). It should be noted that the two identified sequences with B/A6 recombinants, proved to be a sequence pair with high similarity both for the protease/reverse transcriptase (red branches in the Fig. 4) and integrate coding regions (red branches in the supplemental Fig. 1). As L74I variant was present in virtually all A6 sequences it was also observed within the identified clusters. In subtype B one cluster with NNRTI K101H/E138A mutation was observed, in four sequence pairs there was also evidence of the shared resistance patterns (NRTI: D67N/K291Q and T215V, NNRTI: V106I, integrase: E157Q).

**Discussion**

This study presents the novel data on the HIV-1 subtyping and patterns of drug resistance variants among blood donors from Poland collected in the years 2009–2017. No similar study was performed in the country; moreover the added value of this dataset is the analysis of not only HIV-1 PR/RT but also integrase coding regions. Sampling obtained for sequencing included majority of available samples from the donors with positive HIV molecular test in the country for the above timeframe, therefore may be considered representative for entire population of Polish blood donors.
Subtyping patterns remain in line with the previous data published for the region, with the highest prevalence of the subtype B followed by the subtype A1,11,13,14,35. Of note, in this study we identified 14 sub-subtype A6 sequences and three unique recombinants containing these variants, confirming its import from Russia and Ukraine, most likely by immigration36. Interestingly, non-subtype B frequency, especially A6 was associated with repeat donors. This is indicating the circulation of this variant in Polish population adding to the subtype complexity. Additionally, we have identified three recombinants between A6 and B subtype, with a pair of B/ A6 sequences showing high similarity despite diagnosis in the distant centers, which may indicate formation of the novel circulating recombinant form. No circulating recombinant forms between A6 and B variants have been described so far. Furthermore, L74I polymorphism was almost invariably (94.1%) present in A6 sequences.

Figure 1. Maximum likelihood tree showing the subtype A identification with the reference sequences from HIV sequence compendium supplemented with A1 and A6 sequences from GenBank. A1 variant is colored in blue, A6 in green.
Figure 2. Three unique recombinant form genome maps and the separate phylogenetic ML trees with the corresponding partial protease/reverse transcriptase and integrase sequences confirming the subtype assignment. For the tree reconstruction a dataset obtained from HIV sequence compendium 2018 was used. Multiple branches for the same subtype were collapsed. A. sequence 2886 acquired in the city of Warsaw, B. sequence 2495 acquired in the city of Krakow, C. sequence 5474 acquired in the city of Warsaw.
This polymorphism, albeit not included in the drug resistance interpretation algorithms, was associated with increased risk of the virologic failure among patients infected with A6/A1 variants treated with long acting cabotegravir/rilpivirine in the ATLAS 2 M study. Further increase in the frequency of A6 sub-subtype in Poland may negatively affect the future virologic response rates to these injectable agents, and underscores the necessity for subtyping and resistance testing prior to introduction of this combination. We have also observed high frequency of transmission clusters calculated with the genetic distance of 1.5%, however DRM were infrequent among closely related sequences. Clustering is a common phenomenon among HIV sequences, also frequently observed among subtype A infected individuals in Europe.

In general, data on transmitted drug resistance and HIV subtyping patterns are not collected systematically, especially in the region of the central and eastern Europe and as such this dataset provides an important insight on this issue. This is also the first study reporting on the integrase resistance patterns among European blood donors. In the recert reports, frequency of protease/reverse transcriptase DRM among blood donors ranged from 14% in Catalonia, 12.1–13.2% across Chinese provinces, 11% in Brazil, however transmission of major DRMs remains infrequent. This is in line with presented data, with frequency of major, non-accessory drug resistance variants for protease, reverse transcriptase or integrase being low, not exceeding 5% for each drug class. As this is a first study on HIV drug resistance among blood donors in the country, no previous patterns in this group may be compared. However, in the largest Polish study published so far on the 833 antiretroviral naïve cases transmitted drug resistance to PR and RT was observed in 9% of sequences, being the most common for NRTI (5.8%), followed by PI 2.0% (2%) and NNRTI (1.2%) mutations, with the highest frequency among heterosexually infected individuals (13.4%) and MSM (8.3%). These frequencies are slightly higher than the frequencies observed in the current study. Moreover, there is an emerging signal for the transmission of the non-polymorphic integrase mutation (E138K), previously not observed in the country. As expected the most

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Frequency of major and transmitted drug resistance mutations (red) and variants associated with decreased drug susceptibility (Stanford HIV DB, score ≥ 10, green) for nucleoside reverse transcriptase (a), non-nucleoside reverse transcriptase (b), protease (c) and integrase (d) inhibitors. As non-nucleoside reverse transcriptase E138A mutation is not included in the tDRM list, but is associated with significant reduction of susceptibility to rilpivirine, it was marked in violet.
common NRTI DRMs were thymidine analog mutations, namely M41L, D67N, T215S/V, K219Q, associated with high levels of resistance to zidovudine, but also affect the abacavir and tenofovir sensitivity. Both these agents remain the cornerstone of the first-line treatments according to the recent national and European guidelines. In one sequence the NNRTI non-polymorphic E138K mutation, associated with reduced RPV susceptibility, was found. The remaining observed NNRTI DRMs were accessory, potentially reducing susceptibility to etravirine or rilpivirine (E138A). We have previously observed the similar frequency (5.3%) of the rilpivirine associated DRMs with E138A and E138G being the most common DRM. On the other hand, frequency of the V106I variant, which may be affecting doravirine susceptibility was higher (4.4%) than in reference data (0.8%) from Italy and France published by Soulie et al.

In the integrase region, the most common (9.8%) polymorphism was E157Q, which is usually selected in patients receiving raltegravir or elvitegravir, but not associated with significant effect on the integrase treatment efficacy. This variant may reduce integrase susceptibility if present in combination with other DRMs within this region, especially R263K. It was previously observed that in Poland polymorphism was frequent (21%) especially among females, people with history of injection drug use and hepatitis C coinfection. Blood donor regulations exclude patients with history of drug use or HCV coinfection, however in the current study association with female gender was confirmed.

This study also adds valuable information on the recency of HIV infection among blood donors in Poland, reflected by the Fiebig stages at HIV diagnosis. For this purpose, referral to the infection stages labelled as ‘acute’ (Fiebig I), ‘recent’ (Fiebig II–IV) or ‘established’ (Fiebig V–VI) is commonly used. We have noted, that stages associated with recent infection were observed among 9.9% repeat donors, increasing to 40.5% if the stage V was added to the calculations. This is in line with the previous reports from Poland for the years 2001–2007, however in our study calculation of HIV recency was based solely on the HIV-RNA, p24 and Western-blot patterns, with no implementation of the Recent Infection Testing Algorithm (RITA) assays. Donor testing is obviously intended to ensure blood safety, but it should not be overlooked that early HIV diagnosis in the cohort of blood donors in the setting of the low populational testing prevalence allows for the rapid antiretroviral treatment initiation, reduction of infectivity and risk of onward transmissions.

![Figure 3. (continued)](image-url)
Figure 4. MCMC tree showing the relationship between clusters and mutations and city of diagnosis for subtype B with the inclusion of the protease/reverse transcriptase region of the of the subtype B/A6 recombinant (tree branches marked in red). Transmitted drug resistance substitutions were color-coded and included at the external taxonomical units: brown—resistance against nucleoside reverse transcriptase inhibitors, red—resistance against non-nucleoside reverse transcriptase inhibitors, blue – resistance against protease inhibitors, green—resistance against integrase inhibitors. Clusters have been indicated on the tree with magenta highlight using < 1.5% genetic distance and > 90% branch support.

Limitations of the study include lack of more detailed data on the transmission routes or the risk among identified blood donors. Also, calculation on the HIV infection based on Fiebig scale might have underestimated early infection frequency. For this purpose testing with RITA algorithm would add valuable data on the duration of the infection; testing of blood donors with this algorithm should be considered for the future.

To conclude, this study provides a valuable insight on the HIV molecular epidemiology among blood donors in Poland. Transmission of drug resistance in this group was infrequent, however possible emergence of integrase resistance was noted. This emphasises the necessity to continue surveillance on the HIV mutation patterns. Moreover, high frequency of A6 subtype was found indicating migration associated introduction of this subtype to Poland with subsequent local spread and emergence of the new recombinants with the dominant subtype B. This increase in the HIV diversity may potentially affect the antiretroviral susceptibility, even in the context of the novel integrase inhibitors such as cabotegravir.

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Author contributions

M.P. performed the drug resistance interpretations, interpreted phylogenetic data, analysed the datasets, calculated statistics, written and reviewed the manuscript. E.S. collected the data and samples, interpreted the clinical data, reviewed the manuscript. A.U. performed the sequencing, interpreted the resistance data, reviewed the manuscript. K. Scheibe: performed the sequencing, interpreted the resistance data, reviewed the manuscript. K. Serwin: performed phylogenetic analyses, reviewed the manuscript. P.G. collected the data and samples, interpreted the clinical data, reviewed the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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