HIV AIDS Review 2017/Volume 16/Number 4

Late presenters among newly diagnosed HIV-infected in Poland in 2006-2008

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Abstract

Introduction: Assessment of incidence and factors associated with late vs. early human immunodeficiency virus (HIV) diagnosis among newly diagnosed patients referred to HIV treatment centres in Poland in the years 2006-2008.

Material and methods: Retrospective analysis of medical records of patients reported from eight regional centres for HIV treatment in Poland in years 2006-2008 was conducted. The study population consisted of 1132 HIV-infected patients newly diagnosed with HIV infection, aged 17 years up. To describe characteristics of this group of patients in terms of clinical and immunological presentation multivariate regression analysis of demographic and epidemiology data including: sex, age, mode of transmission, CD4 T cell count, and viral load at the time of diagnosis was performed.

Results: Among 1132 patients included in the study 564 (49.8%) were late presenters (LP) according to European consensus definition. In multivariate logistic analysis including sex, age and route of infection, as independent factors, age (OR = 1.06), intravenous drug use (OR = 2.17 for IVDU vs. MSM), and heterosexuality (OR = 2.07 for Hx vs. MSM) were risk factors for late testing. The same factors were predictors of advanced HIV disease, as well as unknown route of infection (OR = 1.77; p < 0.05). However, multivariate regression analysis revealed that only advanced age was an independent factor influencing lower CD4 T cell count and late presentation (OR = 1.02 per 1 year of age, p < 0.001).

Conclusions: Nearly 50% of patients were diagnosed after the optimal time for antiretroviral treatment initiation, according to actual European and Polish guidelines. These results strongly point out the necessity of further encouragement toward more frequent and earlier HIV testing.

HIV AIDS Rev 2017; 16, 4: 244-250
DOI: https://doi.org/10.5114/hivar.2017.72025

Key words: late presenters, HIV epidemic in Poland, high-risk group for HIV, HIV surveillance system.

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Article history:
Received: 25.07.2017
Received in revised form: 16.02.2017
Accepted: 09.11.2017
Available online: 30.11.2017
Introduction

Despite more than three decades of the human immunodeficiency virus (HIV) epidemic and enormous progress in the treatment of the infection with common access to the highly effective antiretroviral treatment, almost a third of individuals infected with HIV enter a health care system very late in the course of their HIV infection throughout Central and Eastern Europe [1]. This situation has remained unchanged for the last several years without evidence of improvement [2]. Surveillance to identify the exact extent of the problem remains insufficient across Europe.

Currently available highly active antiretroviral therapy (HAART) has dramatically improved the prognosis and survival of HIV infected patients [3]. However, late HIV diagnosis, especially when acquired immune deficiency syndrome (AIDS) defining disease has occurred, constitutes a relevant problem for treatment and leads to individual suboptimal usage of HAART [4-6].

Furthermore, late presenters represent an ongoing and serious challenge to the control of the AIDS pandemic. In a global perspective, HIV late diagnosis perpetuates HIV pandemics, while about 50% of new sexually transmitted infections are transmitted from persons with early phase of HIV disease [7]. HIV late diagnosis and presentation for care is estimated at about 15-38% of cases across the Europe [8], and this has serious societal and economic consequences, despite access to universal insurance coverage for complex specialist care [9-11].

Moreover, in the past it was difficult to evaluate the extent of this problem, as there were numerous and diverse definitions of late presentation during the evaluation of collected data (Table 1).

Currently we share one universal definition and, every patient with CD4 T cell count < 350 cells/µl and/or AIDS defining illness (ADI) fulfills criteria for being recognised as “late presenter” according to current treatment guidelines for HIV infected adults [12].

Since 1985, when HIV diagnostics was introduced in Poland, the number of newly diagnosed cases of HIV disease and AIDS has kept on increasing, particularly in recent years [13, 14].

The problem of late presentation was not widely addressed in Poland. The first Polish data that we present were collected from significant number of Polish patients from different HIV clinics and hospitals in the period 2006-2008.

Therefore, the aim of the present study was to determine the frequency of late presenters among newly diagnosed individuals in the years 2006-2008 using current “late presentation” definition, to compare frequencies of late presentation according to previous definitions and to describe late presenters clinical and immunological characteristics.

Material and methods

The study population consisted of 1132 HIV-infected, Caucasian patients newly diagnosed with HIV infection, aged 17 years up, reported from eight regional centres for HIV treatment in Poland.

Out of 11 centres, operating during the study period in Poland and invited for the survey, four centres provided data for 2006 and 2007 (n = 563) and eight centres for 2008 (n = 591). Foreigners and immigrants as well as patients diagnosed elsewhere were not included. Duplicate records of patients registered in two centres during the analysis period were excluded, with only the earlier record being taken into account.

To evaluate whether the study group was representative of the HIV patient population nationwide, the study data were compared with those registered by the National Health Institute for the same period, i.e. 2006 through 2008.

Data on demographic and epidemiology, including age, sex, mode of HIV transmission (injection drug user – IDUs, heterosexual contact – Hx, male having sex with male – MSM, unknown/other – UNK), CD4 T cell count, serum viral load at the time of diagnosis, and the patient’s medical history were gathered and analysed.

In terms of the numerous definitions of “late presentation” available at the time of the study the data were initially analysed applying a different set of parameters for late presentation (Fig. 1).

Clinical categories A, B, and C, as data were collected and used in the period 2006-2008, were applied according to revised classification system for HIV infection among adolescents and adults from the year 1993 [15]. In addition, a group of persons with very late presentation and CD4 T cell count under 50 cells/µl were analysed.

The appropriate statistical tests were applied for parametric and non-parametric data using Statistica 7.0 software, with p value under 0.05 considered as clinically significant. Tests used for demographic data included Fisher test, Kruskal-Wallis test, Mann-Whitney U-test. Odds ratios were assessed for late presenting risk using as independent factors age, gender, route of infection (IDU, Hx, MSM, and unknown), and univariate and multivariate analyses were performed.

Results

In total 1132 medical records of newly diagnosed patients were identified. The general characteristic of this group is presented in Table 1. These patients represented 46% of all patients diagnosed with HIV in Poland during the analysis period. Two hundred and sixty-four persons were recorded in 2006 and 290 persons in 2007, and 578 in 2008.

Male gender strongly dominated in the analysed group – 911 out of 1132 patients (80.5%). Median age at HIV diagnosis was 31.3 (IQD 26.8-38.6) years; female patients were younger than male: 30 (IQD 24.6-36.3) and 32 (IQD 27.3-39.1) years, respectively, and the difference was statistically significant (p < 0.0001). With respect to the mode of HIV transmission, there were 262 (23.1%) current or previous IDUs, 498 (44%) MSM (including bisexual men) and 275 (24.3%) Hx persons. The route of HIV infection remained unknown for 97 (8.6%)
Table 1. Comparison of medium age, CD4 T cell count, and viral load between particular gender, mode of transmission, and late vs. non-late testers. Statistical significance assessed using Mann-Whitney U test

<table>
<thead>
<tr>
<th>All groups</th>
<th>N (%)</th>
<th>Age (years) Median (IQD: Q25-Q75)</th>
<th>CD4 (cells/μl) Median (IQD: Q25-Q75)</th>
<th>VL (copies/ml) Median (IQD: Q25-Q75)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All groups</td>
<td>1132</td>
<td>31.3 (26.8–38.6)</td>
<td>355 (171–530)</td>
<td>30300 (5260–110580)</td>
</tr>
<tr>
<td>Female</td>
<td>221 (19.5%)</td>
<td>30.5 (24.6–36.3)</td>
<td>305 (125–503)</td>
<td>1950 (2640–75250)</td>
</tr>
<tr>
<td>Male</td>
<td>911 (80.5%)</td>
<td>32 (27.3–39.1)</td>
<td>368 (185–533)</td>
<td>35000 (6020–117000)</td>
</tr>
<tr>
<td>p</td>
<td>0.001</td>
<td>p = 0.02</td>
<td>p = 0.006</td>
<td></td>
</tr>
<tr>
<td>IDUs</td>
<td>262 (23.1%)</td>
<td>31.7 (27.7–38)</td>
<td>289 (90–504)</td>
<td>31713 (6017–106000)</td>
</tr>
<tr>
<td>MSM</td>
<td>498 (44%)</td>
<td>30.3 (26.1–36.4)</td>
<td>418 (265–576)</td>
<td>28555 (6278–103000)</td>
</tr>
<tr>
<td>Hx</td>
<td>275 (24.3%)</td>
<td>33 (26.8–41.9)</td>
<td>282 (125–453)</td>
<td>29650 (3890–115000)</td>
</tr>
<tr>
<td>UNK</td>
<td>97 (8.6%)</td>
<td>36 (24.6–36.3)</td>
<td>364 (120–472)</td>
<td>41494 (1456–142500)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>MSM vs. Hx: p = 0.002</td>
<td>MSM vs. IDUs: p = 0.002</td>
<td>NS</td>
</tr>
<tr>
<td>Clinical category A</td>
<td>760</td>
<td>29.8 (25.4–36.5)</td>
<td>461 (348–615)</td>
<td>17950U (3840–77500)</td>
</tr>
<tr>
<td>Clinical category B</td>
<td>85</td>
<td>34.3 (29.0–39.9)</td>
<td>230 (134–310)</td>
<td>37100U (7240–124000)</td>
</tr>
<tr>
<td>Clinical category C</td>
<td>287</td>
<td>35.2 (30.0–43.9)</td>
<td>65 (22–158)</td>
<td>100000U (1456–142500)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>C vs. A and B vs. A: p &lt; 0.001; B vs. C: NS</td>
<td>C vs. B and C vs. A: p &lt; 0.001</td>
<td>NS</td>
</tr>
<tr>
<td>CD4 &lt; 50 cells/μl</td>
<td>137</td>
<td>34.6 (30.4–42.5)</td>
<td>20 (10–31)</td>
<td>120000U (85000–520000)</td>
</tr>
<tr>
<td>CD4 &gt; 50 cells/μl</td>
<td>995</td>
<td>30.9 (28.2–37.9)</td>
<td>398 (243–562)</td>
<td>23450U (4290–98000)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>p &lt; 0.001</td>
<td>p &lt; 0.001</td>
<td></td>
</tr>
<tr>
<td>CD4 &lt; 200 cells/μl</td>
<td>323</td>
<td>34.8 (30.3–43.9)</td>
<td>64 (23–135)</td>
<td>100000U (29800–323593)</td>
</tr>
<tr>
<td>CD4 &gt; 200 cells/μl</td>
<td>809</td>
<td>30.0 (25.5–36.7)</td>
<td>452 (337–604)</td>
<td>17800U (4170–80800)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>p &lt; 0.001</td>
<td>p &lt; 0.001</td>
<td></td>
</tr>
<tr>
<td>CD4 &lt; 200 cells/μl + clinical category C</td>
<td>363</td>
<td>34.8 (30.0–43.6)</td>
<td>76 (28–160)</td>
<td>100000U (27093–279000)</td>
</tr>
<tr>
<td>Other (all without CD4 &lt; 200 + clinical category C)</td>
<td>769</td>
<td>29.8 (25.3–36.5)</td>
<td>462 (348–615)</td>
<td>17000U (3865–74300)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>p &lt; 0.001</td>
<td>p &lt; 0.001</td>
<td></td>
</tr>
<tr>
<td>CD4 &lt; 350 cells/μl</td>
<td>556</td>
<td>34.0 (29.4–41.6)</td>
<td>166 (51–266)</td>
<td>71750U (15609–191000)</td>
</tr>
<tr>
<td>CD4 &gt; 350 cells/μl</td>
<td>576</td>
<td>29.8 (25.3–36.5)</td>
<td>526 (434–670)</td>
<td>13100U (3122–61200)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>p &lt; 0.001</td>
<td>p &lt; 0.001</td>
<td></td>
</tr>
<tr>
<td>CD4 &lt; 350 cells/μl + clinical category C</td>
<td>564</td>
<td>34.0 (29.4–41.5)</td>
<td>170 (52–170)</td>
<td>72200U (15609–191000)</td>
</tr>
<tr>
<td>Other (all without CD4 &lt; 350 + clinical category C)</td>
<td>568</td>
<td>29.0 (24.7–35.6)</td>
<td>529 (435–672)</td>
<td>13056U (3100–58400)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td>p &lt; 0.001</td>
<td>p &lt; 0.001</td>
<td></td>
</tr>
</tbody>
</table>

*NS – not statistically significant
IDUs – injection drug user, MSM – male having sex with male, Hx – heterosexual contact, UNK – unknown/other

persons. MSMs were significantly younger than IDUs and Hxs (p = 0.002 for both).

Median CD4 T cell count at diagnosis was 355 cells/μl (IQD 171–330) in the whole group. Female patients had significantly lower CD4 T cell count than male; 305 cells/μl (IQD 125–503) and 368 (IQD 185–533), respectively (p = 0.02). Median CD4 T cell counts did not differ statistically between IDUs and Hxs; 289 (125–453) and 282 (125–453) cells/μl, respectively, but they were significantly higher in MSM – 418 (265–576) cells/μl than in both other groups (p < 0.001 for both). Among IDUs, male patients had significantly lower CD4 T cell count (276 cells/μl) than female ones (349 cells/μl) from this group (p < 0.05).

Depending on the definition of “late presentation”, the incidences for the whole group varied from about 25% when the category C was defined to nearly 50% when the most
Late presenters among HIV-infected in Poland in 2006-2008

103 (20.7)
122 (55)
556 (49.1)
101 (38.5)
86 (21.6)
50.2 μl

192 (38.6)
84 (32)
276 (30.3)
45 (17.2)
196 (39.4)
43 (15.6)
87 (39.4)
103 (11.3)
434 (47.6)
110 (42)
113 (41.1)
32.1
100 (36.4)

DU group the frequency of "very late
287 (25.4)
165 (60)
50.9
91 (33)
155 (59.2)
49.8
28.5
225 (24.7)
363 (32.1)
243 (26.7)
564 (49.8)
167 (60.7)
122 (55)
43 (15.6)
87 (39.4)
103 (11.3)
434 (47.6)
110 (42)
113 (41.1)
32.1
100 (36.4)

Table 2. Frequencies of late testing in groups of different routes of infection and gender

Definition | All, n = 1132 (%) | IDUs, n = 262 (%) | MSM, n = 498 (%) | Hx, n = 275 (%) | Female, n = 221 (%) | Male, n = 911 (%)
--- | --- | --- | --- | --- | --- | ---
≤ 50 | 137 (12.4) | 45 (17.2) | 37 (7.4) | 43 (15.6) | 34 (15.3) | 103 (11.3)
Clinical category C | 287 (25.4) | 84 (32) | 76 (15.3) | 91 (33) | 62 (28) | 225 (24.7)
≤ 200 | 323 (28.5) | 101 (38.5) | 86 (21.6) | 100 (36.4) | 80 (36.2) | 243 (26.7)
< 200 and/or category C | 363 (32.1) | 110 (42) | 103 (20.7) | 113 (41.1) | 87 (39.4) | 276 (30.3)
≤ 350 | 556 (49.1) | 153 (58.3) | 192 (38.6) | 165 (60) | 122 (55) | 434 (47.6)
< 350 and/or category C | 564 (49.8) | 155 (59.2) | 196 (39.4) | 167 (60.7) | 122 (55) | 437 (48)

IDUs – injection drug user, MSM – male having sex with male, Hx – heterosexual contact

Discussion

Primarily, it is worth underlying the fact that in the presented study almost half of the patients were considered late presenters according to the European Late Presenter Consensus Working Group. This recent universal European definition considers a person presenting with CD4 T cell count less than 350 cells/µl and/or AIDS defining event, regardless of CD4 T cell count as being a late presenter, and an individual with an advanced HIV disease as having CD4 T cell count below 350 cells/µl and/or category C (AIDS-defining disease diagnosed) (Table 2).

According to the current definition, the groups with the highest rate of late presentation were IDUs (59.2%) and Hxs (60.7%). Late presentation was also more frequent among female patients (55%) than among males (48%).

In addition, 12% of “very late testers” were identified. These were persons with CD4 T cell count under 50 cell/µl at HIV diagnosis. In the IDU group the frequency of “very late testers” was 17% while that among persons infected via the heterosexual route was 15.3% and for MSM it was 7.4%. The frequency of very late presenting females was 15.3% while for males it was 11.3%. Statistically significant differences in the frequencies were found for all late presenting definitions between groups of IVDUs and heterosexually infected persons vs. MSM (p < 0.05 for all comparisons).

Patients diagnosed with clinical category A were significantly younger (median 29.8; IQD 25-36.5 years) than those with categories B and C (median 34, IQD 29-39.9 years and 35.2, IQD 30-43.9 years), respectively – differences A vs. B and A vs. C were statistically significant, p < 0.001. The same tendency was observed for all “late presenter” definitions, and patients diagnosed late were significantly older than those who presented earlier for diagnosis Table 1.

Data on HIV viral load was available for 792 out of 1132 patients (70%). As was suspected, medium viral load was significantly higher in late presenting group of patients regardless of definition (p < 0.05 for all) (Table 1).

Age, higher viral load, intravenous drug use, and heterosexuality were all predictors of late presentation according to current definition: IDUs vs. MSM – OR (odds ratio) 2.17 (95% CI: 1.6-3.0), Hx vs. MSM – OR 2.1 (95% CI: 1.5-2.0), age OR = 1.07 (95% CI: 1.05-1.08) for 1 year of age (p < 0.001 for all). For advanced stage of HIV disease (CD4 < 200 cells/µl and C category of disease) female sex was also a predictor (OR = 1.49, p = 0.01). Multivariate logistic analysis showed that age, heterosexuality and intravenous drug use were independent predictors of late presenting regardless of definition (Table 3). Additionally, for advanced HIV-disease and AIDS diagnosis (category C of HIV disease) unknown route of infection was also a predictor. However multivariate regression analysis revealed that elder age was an independent factor influencing lower CD4 T cell count and late presentation. Multivariate logistic analysis showed that age, heterosexuality, and intravenous drug use are independent predictors of late presenting regardless of definition (Table 3).

In univariate analysis female sex, increased age, higher viral load, and intravenous drug use were risk factors for lower CD4 T cell counts (p = 0.02, p < 0.001, p < 0.001, p = 0.005). However, the multivariate regression analysis revealed that increased age was independently related to the lower CD4 T cell count (p < 0.001) in males with a history of IDU (p = 0.003).

Table 2. Frequencies of late testing in groups of different routes of infection and gender

Figure 1. Different characteristics of analyzed population according to accepted definition

The frequency of very late presenting females was 15.3% while that among persons infected via the heterosexual route was 15.3% and for MSM it was 7.4%.

Patients with CD4 T cell count under 350 cells/µl and/or AIDS diagnosis (category C of HIV disease) unknown route of infection was also a predictor. However multivariate regression analysis revealed that elder age was an independent factor influencing lower CD4 T cell count and late presentation.

Multivariate logistic analysis showed that age, heterosexuality, and intravenous drug use are independent predictors of late presenting regardless of definition (Table 3).
Table 3. Multivariate analysis of factors influencing late presentation for different definitions of late presenters

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>Age* (95% CI, p)</th>
<th>Female vs. male (95% CI, p)</th>
<th>IDUs vs. MSM (95% CI, p)</th>
<th>Hx vs. MSM (95% CI, p)</th>
<th>UNK vs. MSM (95% CI, p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>1132</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OR for CD4 &lt; 50 cells/µl</td>
<td>137</td>
<td>1.04 (1.02-1.06, p = 0.000005)</td>
<td>1.15 (0.71-1.86, p = 0.56)</td>
<td>2.44 (1.5-3.97, p = 0.0003)</td>
<td>1.87 (1.1-3.19, p = 0.02)</td>
<td>1.37 (0.67-2.8, p = 0.38)</td>
</tr>
<tr>
<td>OR for C category of HIV infection</td>
<td>287</td>
<td>1.06 (1.04-1.07, p = 0.000000)</td>
<td>0.89 (0.6-1.3, p = 0.54)</td>
<td>2.72 (1.87-3.97, p = 0.000000)</td>
<td>2.55 (1.7-3.83, p = 0.000007)</td>
<td>2.69 (1.62-4.45, p = 0.0001)</td>
</tr>
<tr>
<td>OR for CD4 &lt; 200 cells/µl</td>
<td>323</td>
<td>1.06 (1.05-1.08, p = 0.000000)</td>
<td>1.26 (0.87-1.83, p = 0.23)</td>
<td>2.83 (1.97-4.06, p = 0.000000)</td>
<td>2.16 (1.42-3.15, p = 0.0002)</td>
<td>2.13 (1.3-3.52, p = 0.003)</td>
</tr>
<tr>
<td>OR for CD4 &lt; 200 cells/µl + C category</td>
<td>363</td>
<td>1.06 (1.05-1.08, p = 0.000000)</td>
<td>1.19 (0.82-1.72, p = 0.35)</td>
<td>2.64 (1.86-3.75, p = 0.000000)</td>
<td>2.13 (1.45-3.13, p = 0.0001)</td>
<td>1.77 (1.08-2.9, p = 0.02)</td>
</tr>
<tr>
<td>OR for CD4 &lt; 350 cells/µl</td>
<td>556</td>
<td>1.06 (1.05-1.08, p = 0.000000)</td>
<td>1.04 (0.73-1.5, p = 0.82)</td>
<td>2.15 (1.55-2.99, p = 0.000005)</td>
<td>2.07 (1.44-2.98, p = 0.000096)</td>
<td>1.08 (0.68-1.72, p = 0.75)</td>
</tr>
<tr>
<td>OR for CD4 &lt; 350 cells/µl + C category</td>
<td>564</td>
<td>1.07 (1.05-1.08, p = 0.000000)</td>
<td>1.004 (0.7-1.45, p = 0.99)</td>
<td>2.17 (1.56-3.02, p = 0.000004)</td>
<td>2.1 (1.45-2.03, p = 0.000078)</td>
<td>1.03 (0.7-1.45, p = 0.9)</td>
</tr>
</tbody>
</table>

*OR – per year
IDUs – injection drug user, MSM – male having sex with male, Hx – heterosexual contact, UNK – unknown/other

A group of the highest risk of infection was the group of IDUs. Nowadays the incidence of HIV acquisition by using drugs intravenously has decreased significantly. This change has taken place due to the implementation of harm-reduction programs such as needle and syringe exchange or free needle and syringe distribution and opiate substitution programs, e.g. methadone programs [25]. It seems worth noting that types of drugs were also changed during the past few years, from domination of “homemade” heroin to usually oral drugs.

Thus, we are facing a situation where half of the newly diagnosed patients in the years 2006-2008 were not able to fully profit from HAART, as recommended by European and Polish guidelines for treatment [16, 17].

Furthermore, these data are similar to those obtained from other European countries, most often applying the 200 cells/µl threshold as a definition of later presentation. For example, a large Swiss cohort study revealed that about 30% of patients diagnosed with CD4 T cell count below 200 cells/µl and even 10% with below 50 cells/µl [18]. The factors associated with late diagnosis in the Swiss study were increased age, heterosexual contact, and Danish concur with our observation that a significantly higher proportion of heterosexual patients than homosexuals are late presenters [23, 24].

At the start of the HIV epidemic in Poland, it was clear that injecting-drug-users were driving the spread of illness. However, unlike “safe sex” practices, frequent testing is not a HIV preventive measure.
Late presenters among HIV-infected in Poland in 2006-2008

Late presenters are more likely than early presenters to be diagnosed with opportunistic infections, their risk of death is higher, and their rate of immunological improvement is slower [26, 27]. Initiation of HAART at higher CD4 T cell counts is associated with a lower frequency of antiretroviral drug resistance mutations at virological failure [28]. Late presentation (CD4 < 350 cells/μl) was found to be related with significantly higher costs, especially inpatient costs during the first year of medical care, and two-fold higher costs were observed during further treatment compared to patients presenting with CD4 T cell counts above 350 cells/μl [11].

Moreover, ignorance of the serological status means that late presenters are likely to transmit the virus [29]. However, to present for testing requires the perception of being at risk of those potentially infected and a higher level of suspicion among clinicians of all specialties [30].

So far, several barriers impairing HIV testing have been identified, both among patients and health care providers [31-33]. Among the most important of these barriers in Poland are perception of low or no risk, fear of disclosure among patients (HIV stigma), and lack of knowledge and self-confidence among health care providers when proposing HIV testing. Therefore, it is very difficult to establish the "dark figure", the actual number of currently undiagnosed HIV positive individuals in our country.

Late presentation of HIV infected persons for care seems to be a universal problem in Europe [4], despite significant and constant progress in health care. Therefore, HIV testing policies and practices should be thoroughly analysed, adapted to local conditions, and subsequently implemented. To realise these goals the "HIV in Europe Initiative" was started. In view of the public health implications of early HIV diagnosis for reducing HIV transmission and preventing late presentation, HIV testing should be more frequently offered in all health care settings.

The most important issue is to find an answer to the question: Who is the person with the highest risk of presenting late?

Therefore, nowadays it is believed that the key element in the control of the HIV epidemic is the optimisation of testing for HIV. It is worth emphasising that carried-out studies like the European project HIDES 1 and 2 (HIV Indicator Disease Across Europe Study) have shown that the testing model based on the routine indicator condition-guided HIV testing, the prevalence of which in the population amounts to > 0.1%, is very effective [34, 35].

Since May 2015 at the Emergency Department of the Hospital for Infectious Diseases of Medical University of Warsaw and Out-Patient Clinic of the Hospital for Infectious Diseases in Warsaw, the realisation of a subsequent scientific project, aimed at the evaluation of a percentage of late presenters among patients with the HIV admitted to this unit in recent years, was commenced.

It should be noted that the characteristics of this group of persons will most likely enable the introduction of effective methods aimed at the said group of persons in the future. It seems to be clear that the awareness of transmission trends lets us find and understand an effective way to reach the groups at the highest risk of late presentation.

Conclusions

1. Almost 50% of patients were diagnosed late, after the optimal time to start HAART, in the years 2006-2008 in Poland.
2. Increased age and being a male intravenous drug user were risk factors for lower CD4 T cell count and late presentation.
3. MSMs were diagnosed much earlier in the course of HIV infection, but heterosexuals are at risk of late diagnosis and AIDS-defining disease occurrence.

Conflict of interest

The authors declare no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

References


