Takayasu’s arteritis: a rare disease in Poland

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\begin{abstract}

Introduction. Takayasu’s arteritis (TA) is a rare and potentially life-threatening granulomatous large-vessel vasculitis that involves mostly in the aorta and its proximal branches, and occurs most commonly in young females. This study measures the incidence and prevalence of TA, and assesses the gender distribution and territorial differences in the occurrences of this disease in Poland over a five-year period. To the best of our knowledge, this is the first evaluation of this rare disease in Poland based on a hospital morbidity database.

Materials and method. Analyses were performed with population-based administrative data obtained from a national hospital morbidity study carried out between January 2011 – December 2015 by the Polish National Institute of Public Health. Yearly incidence rates and prevalence of TA were calculated using the number of TA patients and corresponding census data for the overall Polish population.

Results. Data included 660 hospitalization records. The final study sample comprised 177 patients: 154 female (87%) and 23 male (13%) with first-time hospitalization for TA. The mean age was 45.4 years (95% CI: 42.9 – 47.8; SD 16.8; range 4 –81 years), median 47. The incidence rate of TA was estimated at 0.92 per million per year (95% CI: 0.68 – 1.16). Five-year TA prevalence was estimated to be 4.6 per million. Incidence rates of TA did not vary significantly between more urban and more rural regions.

Conclusions. The incidence of TA in Poland was similar or lower to data reported by other European countries. The study provides epidemiological data on TA in Poland that may be useful while comparing it with other geographical regions.

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Key words
vasculitis, epidemiology, aortoarteritis, pulseless disease, granulomatous arteritis, rural regions, urban regions

INTRODUCTION

Takayasu’s arteritis (TA), a potentially life-threatening, granulomatous large vessel vasculitis involves mostly the aorta and its proximal branches, and occurs most commonly in young females. The data on the epidemiology of TA is limited, probably due to the rarity of the disease. Although the disease has a worldwide distribution, it is generally thought to be much more common among Asian populations. The true incidence and prevalence of TA is underestimated, and many patients remain undiagnosed or wait for a long time before the correct diagnosis is made [1–2]. In a recent study from Japan it was reported that the time from onset to diagnosis was significantly shortened in the past decade, which may be related to the development of noninvasive diagnostic imaging tools [3]. Pregnancy seems to cause serious risks for both maternal and foetal health. Epidemiological studies on TA are necessary as it was reported that there was a high prevalence of pregnancy-related concerns in TA patients, although the maternal and foetal outcomes were favourable [4]. TA negatively affects pregnancy outcomes and disease activity increases the risk of obstetric and maternal complications [5].

The etiology of TA remains poorly understood, but genetic contribution to the disease pathogenesis is supported by the genetic association with HLA-B*52 [6–7]. Genetic predisposition may be responsible for an increased occurrence of TA; however, there are differences in incidence and prevalence of this disease in different parts of the world. According to a nationwide Japanese registry, there were at least 5,881 patients with TA in Japan in 2012, with the prevalence believed to be over 40 per million [8]. In another study in Japan, the incidences of TA were estimated to be 1–2 per million [9]. In Turkey the situation was different region-dependent. In a hospital-based study conducted in the north-western part of Turkey (Eastern Thrace) the incidence was estimated to be 3.4 per million per year [10]. In other Turkish study the annual prevalence was estimated as 12.8 (95% CI 12.0 – 13.6) per million; 23.5/million (95% CI 21.9 – 25.0) in females and 1.9/million (95% CI 1.5 – 2.4) in males. The prevalence was higher at 8.8/million (95% CI 7.7–10.0) in the population >40 years of age. During the study period, the mean annual incidence of TA was estimated as 1.11/million (95% CI 0.54–1.67) [11]. In Israel, the incidence of Takayasu’s arteritis was reported to be 2.1 per million (95% CI 1.2–2.9) [12]. In Western Australia, per million inhabitants the annual incidence and prevalence was 0.3 and 3.2 for Caucasians and 1.1 and 15 for Asians [13]. In Arabs, the demographical and clinical findings of TA were reported to be similar to that reported from different parts of the world [14].
TA is regarded as an endemic disease among Asian populations. However, recently studies from Europe show an increasing trend of incidence and prevalence of this rare disease in the Old Continent although the studies on the incidence of TA are rather limited. A study conducted in Germany (Federal State of Schleswig-Holstein, north Germany) that covered the period 1998–2002 indicates the incidence of Takayasu’s arteritis to be 0.4–1.0 per million per year [15], based on data from the Regional Registry that covered mainly hospital records. Similar incidence can be observed in the United Kingdom (Norfolk county, eastern England) in a study that covered the period 2000–2005. It indicates the incidence of Takayasu’s arteritis to be 0.8 per million per year overall, and about 0.3 per million per year when only patients below 40 years of age were take into consideration [16]. A Swedish (Skåne) hospital-based study for the period 1997–2011 estimated the incidence of TA to be about 0.7 per million per year [17]. Meanwhile, a Danish study that covered the period 1990–2009 in the eastern part of the country, based on the Central Registry (1990–2006) and hospital-based data (2007–2009), estimated the incidence of TA to be about 0.4 per million per year overall, and about 0.6 per million per year when the patients under 40 years were taken into consideration [18]. Contrary to these incidences, three other European countries–south-east Norway [19], Spain [20] and Lithuania [21], are situated where the incidences of TA is estimated at between 1–2 per million. Among Scandinavian countries, the prevalence of Takayasu’s arteritis is highest in Norway and calculated to be between 22–25 per million [19]. In contrast, in Sweden the prevalence is twice as low and estimated to be 13.2 per million [17]. The prevalence in Spain in hospital-based study is estimated to be 10.5 per million [20]. The lowest prevalence in Europe is estimated in the United Kingdom – 4.7 per million [16]. In Denmark, in a study based on Central Registry and hospital data, the prevalence of Takayasu’s arteritis is calculated to be 8 per million overall, and about 12 per million when patients <40 years are considered [18].

OBJECTIVES

Apart from case reports, there is little information regarding TA in Poland [22–28]. The presented study measured the incidence and prevalence of TA, and assessed its gender distribution and territorial differences in Poland over a five-year period. To the best of our knowledge, this is the first evaluation of this rare disease in Poland based on a hospital morbidity database.

MATERIALS AND METHOD

This is a population-based and retrospective study using hospital discharge records with Takayasu's arteritis diagnosis obtained from a Polish National Institute of Public Health survey. The data originates from the period 2011–2015. The study evaluated records on all first-time hospitalized patients with Takayasu's disease diagnosis. This data are obligatory sent to the National Institute of Public Health from all hospitals, with the exceptions of psychiatric and military hospitals. These anonymous records consist of information on ICD10 diagnosis, date of admittance, date of discharge, date of birth, gender, and place of residence.

In a Turkish study, a case search was performed electronically in the information systems of these hospitals using The International Classification of Diseases Tenth Revision (ICD-10) code for Takayasu’s arteritis (M31.4) [11]. In the current study, two analytic samples were taken into consideration – all hospitalizations for Takayasu’s arteritis and first-time hospitalizations for Takayasu’s arteritis. Beside data from the hospital morbidity study, demographic data for the general Polish population were obtained from the Statistic Poland [29]. Based on that data, the incidence and prevalence rates were estimated using the number of Takayasu’s arteritis patients and corresponding census data. The study was conducted in accordance with generally applicable law.

TA is a primary vasculitis that causes stenosis or occlusion, rarely aneurysm and distal ischemia [30]. This type of arteritis may require treatment and advanced diagnostic procedures that can be performed in inpatient settings. Analysis of hospitalizations of TA patients from the National Register in multi-year time of observation may be a good way to estimate the epidemiology of this rare, but still life-threatening disease which may negatively affects pregnancy outcomes.

Statistical analysis. All statistical analyses were performed using the licensed version of Statistica Software [31] and WINPEPI programmes [32]. The results of descriptive analyses were expressed as means, medians and ranges for continuous variables, and counts and percentages for categorical data. 95% CIs were calculated assuming Poisson distributions of the observed cases. The incidence rate was calculated using the number of disease cases divided as the population figure. Linear regression was used for trend analysis. Nonparametric tests were used as appropriate when normality assumptions were not met. A two-sided P value less than 0.05 was considered to be statistically significant.

RESULTS

Data included 660 hospitalization records. The final study sample consisted of 177 patients with first-time hospitalization for TA between 2011–2015. The number of first-time hospitalization per year varied from 47 in 2011 to 31 in 2015. The mean age was 45.4 years (95% CI: 42.9–47.8; SD 16.8; range 4–81 years), median 47. Among the TA patients, 37% were aged ≤40 years and 59% were aged ≤50 years at diagnosis of TA. The mean age of female patients was 47.2 years (95%CI: 44.6–49.7; SD 16.0; range 4–81 years) and the mean age of male patients was 33.0 years (95% CI: 25.7–40.4; SD 16.9; range 14–66 years). The mean age of male patients was significantly lower compared to female patients (p < 0.001). Based on hospitalization records and census data, the annual incidence of TA was estimated to be 0.92 per million (95%CI: 0.677–1.161). Five-year TA prevalence was estimated to be 4.6 per million. TA was more frequently observed among women than men (87% vs 13%; P<0.001). The number of newly-diagnosed TA cases in Poland by gender per year across the study period is presented in Figure 1. The number of newly-diagnosed TA cases in Poland by gender per year across the study period is presented in Figure 1. Analysis of hospitalizations of TA patients from the National Register in multi-year time of observation may be a good way to estimate the epidemiology of this rare, but still life-threatening disease which may negatively affects pregnancy outcomes.

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an Italian study among 104 patients, 91 were women (87.5%)—a quite high female to male ratio of 9:1. Data from the French Takayasu network reported female to male ratio at 6.7:1. Other studies from northern Europe [19] reported female to male ratio at 6.7:1. Data from the French Takayasu network reported that median age at TA diagnosis was 36 years [33]; in Greece, the median age at TA onset was 31 years [38]; in southern Sweden the median age at TA diagnosis was 23 years [17] and in eastern Denmark the median patient age at the time of the TA diagnosis was 36.3 years [18].

In a few European countries median age was higher than that reported from Poland in the current study, e.g. in the UK, the median age was 51 years [16]. In a recent study, northern Europeans had a mean age of 36.3 years at diagnosis [19]. In an Italian study, the mean age at disease onset (determined by the occurrence of the earliest specific symptom or a finding attributable to the disease) was 29.4 years in men and 29.1 years in women, a difference that was not statistically significant [1]. It is worth mentioning that in studies from China the reported mean age at onset was 28.9 ± 12.0 years [36], and in another study the median age at onset of the disease was 23 years old [37]. In a retrospective, one-centre study from Poland the mean age at disease onset was 26 years, but it was highlighted that the mean diagnostic delay was 19 months [28].

These variations among the presented studies may derive from geographical and genetic differences between the populations, but also may be due to methodological differences [39]. The results of the current study suggest the participation of many factors that may increase the risk of disease. The genetic factors and gender may probably play a more important role in the development of TA in Poland than territorial factors.

In this study we observed 4 deaths during five years observation. In a US cohort of 126 patients, the overall survival was 97% at 10 years and 86% at 15 years [40]. In a retrospective multi-centre study from the French Takayasu network after a median follow-up of 6.1 years, death was observed in 5% [33]. Among the Chinese, the median survival time was 102.5 months [41]. Mortality rate estimation was reported to be 2.8% (2000–2010) among the Japanese [3]. In other study from Japan, the overall survival rates for patients with TA were 97.7%, 97.7% and 95.3% at one, five and 10 years [42]. In a study from South Korea, survival rates were 92.9% at the fifth year and 87.2% at the tenth year [43].

Strengths and limitations. The authors believe that the current study has major strengths. Since it was performed in a large population of Poland, it produced a large, unselected TA cohort. However, the presented study also has some limitations, one of which is that it was based on a retrospective review, and relied on the discharge records from inpatient hospitalizations. This excludes patients who were seen only as outpatients or were in a stable period of this disease and had not needed hospitalization for more than five years. The first
appearance of diagnosis of Takayasu’s arteritis in the inpatient discharge database does not necessarily have to be the date of first diagnosis. This may lead to overestimation of the count of incident cases. Nevertheless, the 5-year period time of observation in this study may minimize the overestimation.

CONCLUSIONS
Takayasu’s arteritis incidence in Poland is lower or comparable to that in other European countries. Incidence rates of TA did not vary significantly between more urban and more rural regions. The disease was observed more often in females than males. The occurrence of Takayasu’s arteritis may be related to many factors, but genetic and personal factors seem to be more relevant than those connected with the environment. The number of people suffering from Takayasu’s arteritis may be underestimated, but a better described epidemiology may be achieved using data from national registers in a longer follow-up.

REFERENCES