Cardiovascular risk and metabolic profile of Polish citizens from Lower Silesia. First signs of metabolic crisis?

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Abstract

Introduction: Population biobanks are essential for the development of public health screening and improvement of personalized medicine. Since 2012, Biobank of Łukasiewicz Research Network – PORT Polish Center for Technology Development (PORT Biobank) has collected more than 120 000 biological samples from nearly 5000 inhabitants of Lower Silesia, together with a variety of demographic, anthropometric, life style and health information.

Material and methods: The analyzed group consisted of 2274 participants (1398 women, 876 men). Both women and men were further subdivided into five age decades (20+, 30+, 40+, 50+, 60+). For this study, the level of lipids (total cholesterol, high-density lipoprotein [HDL], low-density lipoprotein [LDL], triglycerides) was estimated and correlated with the level of high-sensitivity C-reactive protein (hs-CRP) and biometric parameters.

Results: We have demonstrated for the first time that biochemical changes that may lead to cardiovascular diseases (CVD) occurred already in the group of people aged 30+. Our observation is based on measurements of lipids, glucose, inflammatory (hs-CRP) and biometric markers such as body mass index (BMI) and waist-to-hip ratio (WHR).

Conclusions: Positive correlations with age for these variables suggest the ongoing progress of metabolic changes, which in the end may lead to a fatal outcome such as myocardial infarction or stroke. It suggests that CVD screening programs should be dedicated to a wider group, especially younger citizens, in order to prevent fatal outcomes related to CVD.

Key words: cardiovascular diseases, metabolic crisis, population-based screening, biobanking.

Introduction

Population-based biobanks are essential for the development and monitoring of public health [1–5]. Poland became a member of the biggest biobanking infrastructure in Europe (BBMRI-ERIC) in 2016 and a large proportion of public funds has been dedicated towards development of biobanking infrastructures [6]. Indeed there is an ongoing effort towards building regional biobanks and longitudinal cohorts which can be used for the development of precision medicine and profiling of disease screening programs [7, 8]. The population-based study conducted on citizens of Lower Silesia (2012–2016) is one of the first projects performed by the newly established PORT Biobank, created due to lack
of population-based cohorts in this area. The research profile is defined by high incidence of cardiovascular diseases (CVD) already observed in other Polish studies (WOBASZ I, WOBASZ II and NATPOL – National Multicenter Health Survey and Arterial Hypertension and other CVD risk factors in Poland, TERCET, REGARDS) [9–13]. The PORT Biobank population consisted initially of almost 4000 participants aged 20–70 years with a main scope to characterize the risk of CVD and diabetes on the basis of biomarker and biometric measurements. The additional aspect that convinced us to create such a program was related to the fact that the younger generation (younger than forty years old) is not eligible for some screening programs, despite the fact that during the last few decades eating habits as well as level of physical activity have markedly changed. As described previously [14], a reduced fat and cholesterol diet was reported only by every 5th patient and a low calorie diet by every 100th participant in the WOBASZ study. Furthermore, the quality of nutrition and health knowledge in subjects with diagnosed CVDs in the Polish population turned out to be insufficient. Another study also reported that intake of saturated fat, sugar and complex carbohydrates is a cause for concern especially in Eastern European countries [15]. Impaired glucose (GLUC) level among European societies has a direct effect on total CVD incidence, and together with an elevated level of low-density lipoprotein cholesterol (LDL-C) potentially accelerates atherosclerosis [13, 16, 17]. Moreover, physical activity including active commuting (i.e. walking, cycling) of Polish citizens remains unsatisfactory, despite promotion of a healthy lifestyle [18]. For example about 66–68% of Finnish or Danish citizens report practicing vigorous sport once a week in contrast to 28% of Poles [19]. Taking into consideration risk factors mentioned above, we may assume that intervention actions such as broad screening for CVD and diabetes is required and should also

Table I. Sex distribution and quantitative parameters of the studied cohort. The study group consisted of 2274 participants divided into five age decades (20+, 30+, 40+, 50+, 60+). Distributions of sex and quantitative information in relation to decades are shown in the table.

<table>
<thead>
<tr>
<th>Age decade</th>
<th>Women (n = 1398)</th>
<th>Men (n = 876)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20+</td>
<td>460 (68.15%)</td>
<td>215 (31.85%)</td>
</tr>
<tr>
<td>30+</td>
<td>261 (54.26%)</td>
<td>220 (45.74%)</td>
</tr>
<tr>
<td>40+</td>
<td>274 (58.55%)</td>
<td>194 (41.45%)</td>
</tr>
<tr>
<td>50+</td>
<td>263 (64.30%)</td>
<td>146 (35.70%)</td>
</tr>
<tr>
<td>60+</td>
<td>140 (58.09%)</td>
<td>101 (41.91%)</td>
</tr>
</tbody>
</table>

be addressed to the younger generation. Monitoring of diet and physical activity, as well as regular measurement of blood pressure, lipids and other CVD markers, should be performed even at the age of thirty, in order to determine the critical time point when the human body stops compensating side effects of the modern lifestyle [20, 21]. Attenuated atherosclerosis is strongly related to inflammatory status. Hence the relationship between level of lipids and high-sensitivity C-reactive protein (hs-CRP) has been associated with adverse cardiovascular events due to vascular inflammation and formation of atherosclerotic plaque [22, 23]. The aim of the study was to characterize metabolic changes and determine the age when the first symptoms of metabolic crisis occur in the population of inhabitants from the Lower Silesia region. Our population-based study employs a novel approach such as a broad spectrum of metabolic data and incorporation of the youngest participants from the age of twenty and is one of the first to be organized in Poland.

Material and methods

Study group

The initial cohort of volunteers who decided to participate in the PORT Biobank study consisted of 4098 individuals, aged 20–70, which is about 1% of the local population. All participants who declared any chronic disease (for example high blood pressure, CVD, diabetes, thyroid problems), regular use of medicines or those individuals who had an elevated level of CRP (> 10 mg/dl) were excluded from the data analysis. We did not exclude patients on the basis of smoking status or physical activity as the idea of the study was to present a typical cross-sectional population. The final group consisted of 2274 participants (1398 women, 876 men). Both women and men were further subdivided into five age decades (20+, 30+, 40+, 50+, 60+). Detailed distribution of sex in relation to decades of age and quantitative information about the cohort are shown in Table I. The study protocol was examined and consent to proceed was granted by the supervising Bioethical Committee (decision numbers: 73/2012, 482/2015, 477/2016 and 607/2017). Informed consent was obtained from all participants prior to sample collection.

Sample collection

Fasting blood samples were collected using the BD Vacutainer collection system (plastic tubes with spray-coated silica for serum extraction) according to the manufacturer’s protocol. Serum for biobanking and routine in vitro diagnostic (IVD) laboratory testing was spun at 2500 g for 15 min at
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Body metrics

Body metrics and a medical questionnaire (including familial history of CVD and diabetes) were recorded for each study participant. Weight and height were used to calculate body mass index (BMI) according to the following formula: BMI = weight [kg] divided by the square of height [m] [24]. Waist and hip measurements [cm] were used to calculate waist-to-hip ratio (WTH).

Lipid profile

Level of total cholesterol (tCHOL), triglycerides (TG) and high-density lipoprotein cholesterol (HDL-C) were determined by enzymatic assay using colorimetric methods performed on the fully automated biochemical analyzer Cobas Integra 400 plus System from Roche Diagnostics GmbH. All measurement steps were performed according to the manufacturer’s protocols (cholesterol 2nd generation test ID 0-586; triglycerides test ID 0-010; HDL 3rd generation test ID 0-331). Low-density lipoprotein cholesterol level was calculated with the Friedewald formula: LDL-C [mmol/l] = total cholesterol [mmol/l] – (HDL-C [mmol/l] + triglycerides [mmol/l]/2.2), when TG level is lower than 4.5 mmol/l.

High-sensitivity C-reactive protein level

Level of hs-CRP was measured by the high-sensitivity method on the fully automated biochemical analyzer Cobas Integra® 400 plus System from Roche Diagnostics GmbH. All measurement steps were performed according to the manufacturer’s protocols (CRPHS test ID 0-033). In brief, the method was based on particle-enhanced (latex) immunological agglutination and turbidimetric precipitate determination.

Glucose

Level of GLUC was measured photometrically on the Cobas c System from Roche Diagnostics GmbH. The enzymatic reference method with hexokinase was performed according to the manufacturer’s protocol (Glucose HK 3rd generation, test ID 05 6831 6).

Statistical analysis

The R package for Windows (version 3.5) was used for statistical calculations [25]. The Shapiro-Wilk test and Q-Q plot were used to assess normality of data. Due to an unacceptable deviation from normality we used nonparametric tests in our study. In order to analyze obtained data, the Kruskal-Wallis test was performed. To verify differences between age groups, pairwise comparison using the post-hoc Nemenyi test with $\chi^2$ approximation for independent samples was adopted. Spearman’s rank correlation between age groups and variation level was estimated. A $p$-value of $\leq 0.05$ was acknowledged as significant.

Results

In order to evaluate the prevalence of metabolic crisis and its components (such as lipids and GLUC impaired level, abdominal obesity or hs-CRP expression level) as CVD risk factors in the Lower Silesia population, we collected and examined serum samples for lipid and GLUC profile, as well as hs-CRP level. For each donor, biometric parameters were also measured. The group characteristics of PORT Biobank participants are shown in Table I. It is important to underline that we excluded from the statistical analysis those who might suffer from or who reported any chronic disease that leads to increased CVD risk. Certainly there is a higher number of women amongst all participants (61.48% vs. 38.52%), who are more willing to participate in the study (Table I).

Lipid profile in different groups of donors showed major changes that are correlated with age (Figure 1). Total cholesterol, TG and LDL-C concentration increased along with maturity. Taking into account upper reference limits for these metabolites (URL) recommended by ESC/EAS guidelines [26], donors from the 30+ group already reached values higher than the URL for total cholesterol. Moreover, total cholesterol level exceeded reference values with an increasing tendency regarding age (Figure 1). An increasing concentration of TG was noted in all studied age groups, but it did not exceed the URL in any of them.

LDL-C level, the major driver for the development of atherosclerotic plaque, demonstrated higher than URL values in the group of donors aged 40+, with an increasing tendency correlated with age. Level of HDL-C did not decrease below reference values in any age group studied (Figure 1).

The next step was to evaluate changes in body metrics that may be correlated with lipid level and may lead to the observed increasing obesity in the Lower Silesia population and increased risk of CVD. Taking into account the URL for this measurement we observed elevated BMI already in the 30+ group and an increasing tendency correlated with age (Figure 2). A similar tendency was observed in the WHR measurements (exceeding reference values evaluated for women from the groups aged 30+ up to 60+). Additionally, hs-CRP and GLUC levels were measured, and an increasing tendency regarding all age groups was noted, and all values could indicate increased risk for future CVDs (Figure 2).
To further confirm obtained results indicating increased risk of CVD in Lower Silesia subjects, correlation analysis between age groups and lipids, hs-CRP, GLUC levels and biometric parameters was performed (Table II). A noteworthy correlation was observed between age groups and tCHOL, TG, LDL-C, hs-CRP, GLUC, BMI, and WHR, demonstrating that the level of metabolites responsible for CVD development changes and is correlated with alteration in body composition.

**Discussion**

The leading cause of mortality in the 21\textsuperscript{st} century is coronary heart disease [27, 28]. More importantly, premature ischemic heart disease is an increasing cause of hospitalization among populations living in developed countries, including Poland [29, 30]. The socio-economic impact triggered by both hospitalization and recovery after myocardial infarction is vast, especially among patients during their period of professional activity [31, 32]. The problem we face today is the lack of efficient screening programs, capturing individuals with the initial stages of CVD, as well as inefficient monitoring of success rate related to the change of lifestyle and sometimes pharmacotherapy. Studying the literature related to the development of cardiovascular biomarkers in relation to the prediction of clinical end-points and the final outcome, still lipids, hs-CRP, GLUC, body metric

**Figure 1.** Changes in lipid profile relative to age groups in Lower Silesia inhabitants. Serum lipid profile, A – total cholesterol (tCHOL), B – triglycerides (TG) and D – high-density lipoprotein cholesterol (HDL-C) levels were measured by enzymatic assay, as described in Materials and Methods section. C – Low-density lipoprotein cholesterol (LDL-C) level was calculated with Friedewald formula. Results exceeding reference values are indicated with a star. Values that increased significantly between particular age decades are indicated with a linker (Pearson’s \( \chi^2 \) test; \( p < 0.0001 \)). Reference values for: tCHOL 2.97–4.91 mM, TG 0.56–1.69 mM, LDL-C < 2.97 mM, HDL-C > 1.03 mM.
measurements, blood pressure, lifestyle and familial history remain crucial [33–35]. That brings us to the conclusion that appropriate tools which are relatively low-cost are available. In fact, it is important to create an entire algorithm and early detection national program that could guarantee a better outcome seen as a reduction of CVD cases including myocardial infarction, especially in the younger age group.

Our data showed that inhabitants of Lower Silesia who were included in the study group (about 1% of the local population) are willing to participate in preventive programs, and a greater proportion are women than men, which probably results from various psychological barriers. For the first time we documented the so-called metabolic crisis seen in study participants aged 30+ onward when compared to younger individuals. Previous studies on CVD incidence in Poland did not include such a young population and division of the cohort into such narrow age groups. Our observation was based on measurements of lipids, GLUC, inflammatory and biometric markers such as hs-CRP BMI and WHR. The tendency to exceed reference values of tCHOL seen in the 30+ group and LDL-C seen in the 40+ group correlated well with changes in the body composition and obesity observed in these groups. These may be very valu-

Table II. Correlation between age and lipid level, hs-CRP and biometric parameters. Spearman’s rank correlation between age groups (n = 2274) together with correlation coefficient (ρ) was calculated. P-values < 0.0001 are indicated in bold.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Age groups (n = 2274)</th>
<th>p-value</th>
<th>ρ</th>
</tr>
</thead>
<tbody>
<tr>
<td>tCHOL</td>
<td>&lt; 0.0001</td>
<td>0.471</td>
<td></td>
</tr>
<tr>
<td>TG</td>
<td>&lt; 0.0001</td>
<td>0.318</td>
<td></td>
</tr>
<tr>
<td>LDL-C</td>
<td>&lt; 0.0001</td>
<td>0.45</td>
<td></td>
</tr>
<tr>
<td>HDL-C</td>
<td>0.05</td>
<td>−0.0414</td>
<td></td>
</tr>
<tr>
<td>hs-CRP</td>
<td>&lt; 0.0001</td>
<td>0.277</td>
<td></td>
</tr>
<tr>
<td>BMI</td>
<td>&lt; 0.0001</td>
<td>0.428</td>
<td></td>
</tr>
<tr>
<td>WHR</td>
<td>&lt; 0.0001</td>
<td>0.280</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. A – High-sensitivity C-reactive protein (hs-CRP), B – glucose (GLUC), C – body mass index (BMI) and D – waist-to-hip ratio (WHR) profiles in relation to age decades. Serum hs-CRP level was measured by immunological agglutination and turbidimetric precipitate determination; GLUC level was measured by enzymatic reaction with hexokinase; BMI and WHR were calculated as described in Materials and Methods section. Results exceeding reference values are indicated with a dot. Values that increased significantly between particular age decades are indicated with a linker (Pearson’s χ² test; p < 0.0001). Reference values for: hs-CRP 0–3.0 mg/l, GLUC: 3.6–5.8 mM, BMI 18.5–24.99, WHR for men ≤ 0.9; WHR for women ≤ 0.85. 
able data that address important issues and may have an impact on the clinical practice in Poland. Increasing BMI, WHR and the level of TG (although still within the URL) affect the liver function and attenuate lipogenesis, leading to the overload of adipocytes. That is the first stage of metabolic syndrome, which accelerates other signaling cascades. Consequently we observed an increase in the hs-CRP level with age, since the fat tissue acts as an endocrine organ promoting synthesis of inflammatory cytokines such as IL-6, which mediates synthesis of CRP in the liver [36]. Dysfunction of lipid homeostasis together with expression of inflammatory molecules is closely linked to severe metabolic disorders, metabolic syndrome and CVD, such as myocardial infarction or stroke [37]. Observations performed on different populations showed that this is a mechanism that cannot be ceased completely, but should be identified at the earliest stage, as it can be reduced by making lifestyle changes by increasing physical activity and changing dietary habits. [19]. Our data in some aspects are very consistent with other studies performed in Poland such as WOBSZ I, WOBSZ II and NATPOL, but we characterized each age group more deeply to accurately determine the age decade at which the metabolic crisis may occur [9, 38]. In fact until now there is not a prevention program to control CVD risk factors among the younger generation, such as our study group. The idea of screening programs often fails to keep up with dynamic environmental changes, and too little attention is paid to those groups of patients who could actually benefit from them. Biobanks due to their broad approach to population-based research may influence the public health policy and may improve early CVD diagnosis and effective prevention.

The study set up by PORT Biobank continues. The current research protocol for the cohort also involves monitoring of blood pressure and measurement of insulin resistance (using the HOMA calculation) that begin to confirm our initial findings based on body metric measurements and an increasing level of TG (data not shown). The analyzed metabolic profile indicated that more specific testing (i.e. ultrasound imaging) could be performed during current recruitment such as carotid-artery intima media thickness. We are considering monitoring of dietary habits and physical activity in a longitudinal project, which may perhaps answer more precisely the question when the metabolic crisis occurs and will clearly indicate the age of metabolic crisis shift towards the younger generation in the future. CVDs must be targeted at a primary health promotion level and traditional and relatively inexpensive tools play a substantial role in population-based health monitoring.

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**Conflicts of interest**

The authors declare no conflict of interest.

**References**