Genetic aspects of obesity and metabolic syndrome in people occupationally exposed to arsenic and certain heavy metals

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Abstract

Introduction. Obesity is a common health issue affecting over 650 million people worldwide. It is an integral element of the metabolic syndrome and leads to the development of arterial hypertension and type 2 diabetes, as well as many other different conditions, including numerous tumours. At the same time, exposure to arsenic, cadmium and lead is increasingly attributed to the development of the metabolic syndrome. Environmentally, this influence concerns the entire population, and is particularly visible in the occupationally exposed population.

Objective. The aim of this study is to prevent selected, best-known elements of the genetic variability and its significance in the development of the metabolic syndrome, taking into account exposure to arsenic, lead and cadmium.

State of the art. CNR1, FTO are related to an increased risk of developing metabolic syndrome, while certain variants of genes responsible for arsenic metabolism (As3MT) and lead (ALAD) are related to their higher toxicity. Knowledge about the relationship between exposure to arsenic, lead and cadmium, and the polymorphism of genes responsible for the development of the metabolic syndrome is scarce. This gap does not appear in more recent research based on the micro RNA expression. Expression of certain miRNA allows detection of both exposure to arsenic and increased risk of cardiovascular episodes in the future, as well as the existence of organ damage at present.

Conclusion. Reinforcement and use of the miRNA knowledge appears to be the right direction, but we should not forget current knowledge about polymorphism of individual nucleotides.

Key words metabolic syndrome, obesity, arsenic, cadmium, lead, genetic variability

Streszczenie

Wprowadzenie. Otyłość jest powszechnym problemem zdrowotnym, dotyczy ponad 650 mln osób na świecie. Jest integralnym elementem zespołu metabolicznego, prowadzi do rozwoju nacisknięcia tętniczego i cukrzycy typu 2, ale też wielu różnych innych schorzeń, w tym licznych nowotworów. Jednocześnie coraz większą wagę przywiązuje się do wpływu narażenia na arsen, ołów i kadm na rozwój zespołu metabolicznego. Wpływ ten środowiskowo dotyczy całej populacji, jednak jest szczególnie widoczny u osób narażonych zawodowo na działanie tych pierwiastków.

Cel pracy. Celem tej pracy jest przedstawienie wybranych najlepiej poznanych elementów zmienności genetycznej oraz jej znaczenia w rozwoju zespołu metabolicznego z uwzględnieniem narażenia na arsen, ołów i kadm.

Aktualny stan badań. Polimorfizm pojedynczych nukleotydów jest dość dobrze poznany. Poszczególne warianty genów leptyny, CNR1, FTO związane są z większym ryzykiem rozwoju zespołu metabolicznego, a pewnie warianty genów odpowiedzialnych za metabolism arsenu (As3MT) oraz ołowiu (ALAD) związane są z większą toksycznością tych pierwiastków. Na tconiast wiedza o związkach narażenia na arsen, ołów i kadm w technologicznych genów odpowiedzialnych za rozwój zespołu metabolicznego jest niewielka. Luki pozbywane są nowsze badania, bazujące na ocenie ekspresji mikroRNA. Ekspresja pewnych mikroRNA pozwala wykryć zarówno narażenie na arsen, jak i większe ryzyko epizodów siercowo-naczyniowych w przyszłości, a także istnienie uszkodzeń narządowych już w chwili obecnej.

Wnioski. Ugruntowanie i wykorzystanie wiedzy o mikroRNA wydaje się słuszne kierunkiem, jednak nie powinniśmy zapominać o posiadanej już wiedzy o polimorfizmie pojedynczych nukleotydów.

Słowa kluczowe otyłość, zespół metaboliczny, kadm, arsen, ołów, zmienność genetyczna
INTRODUCTION

Obesity is a common medical and economic problem. According to the World Health Organization (WHO), globally there are over 650 million obese and 1.9 billion overweight people [1]. These data are all the more impressive as they keep growing, although this illness is potentially avoidable. The classic definition of obesity is BMI over 30 [2]. A new definition of obesity was formulated by the American endocrinological associations, which defines stage 1 obesity as BMI over 25 and the occurrence of complications of obesity [3]. This definition attempts to draw attention to the fact that obesity is a prelude to other diseases, particularly from the perspective of the metabolic complex.

Obesity is an integral part of the classically defined metabolic complex, accompanied by hyperglycaemia, arterial hypertension and dyslipidaemia [4]. Subsequently, there were numerous attempts to expand this definition, yet the changes never gained general acceptance.

Numerous studies were conducted in an attempt to explain the causes of the metabolic complex and obesity. They take into account both the genetic diversity and the effect of the outside environment.

Arsenic, as well as certain heavy metals, such as lead and cadmium, contribute to the onset of metabolic syndrome [5, 6, 7]. Although they are common in nature, there is a particular group of occupationally exposed individuals in which their harmful effects can be particularly evident. One of these groups consists of several thousand employees of the copper smelting plants in the Lower Silesian province of south-west Poland. Apart from the exposed population, heavy metals are also present in tap water. Numerous studies conducted in different sites in Poland have confirmed the presence of metals (especially lead), and in almost every study concentration above acceptable drinking standards were observed. However, over the years tendency of metals concentrations seems positive.

On the other hand, one study showed that the standard monitoring of heavy metals concentrations in Poland is not reliable. Arsenic is a known carcinogen, but its harmful effects, particularly during chronic exposure, are much broader – there were observations of increased cardiovascular risk, peripheral vessel diseases, respiratory system diseases, and neutropaenia [8]. The effect of the arsenic on the development of insulin resistance and diabetes [9] is particularly noticeable. The effects of cadmium exposure mostly include not only thickening of the arterial intima-media, but also renal damage and bone demineralization [9]. The first symptom of chronic lead poisoning is usually anaemia and iron deficiency. These are followed by damage to the kidneys, immune system, or reproductive system [10].

The effect of arsenic, lead and cadmium on the development of the metabolic syndrome is well known. Nevertheless, numerous observations indicate a high individual variability – individuals with the same exposure and similar environmental conditions were observed to have various concentrations of arsenic or lead [11]. This is the effect of a considerable genetic variability of the genes responsible for the broadly-defined metabolism of arsenic, lead and cadmium. Hence, it is difficult to assess the effect of metals on the development of the metabolic syndrome in isolation from individual genetic variability – both responsible for the metabolism of arsenic or lead, and the genes responsible for the development of metabolic syndrome.

The most common well-documented genetic mutations are changes to the genetic polymorphism and micro-RNA variability.

OBJECTIVE

The aim of this study is to present selected, best-known elements of the genetic variability and its significance in the development of the metabolic syndrome, taking into account exposure to arsenic, lead and cadmium.

GENETIC VARIABILITY OF SELECTED GENES RELATED TO THE DEVELOPMENT OF THE METABOLIC SYNDROME – STATE OF THE ART

Leptin. Leptin is one of the more important and best-known compounds participating in regulating the sense of hunger. It is a hormone produced mostly by adipocytes. It triggers a sense of satiety, stops the sense of hunger, and stimulates the sympathetic nervous system through the interaction with leptin receptors expressed mostly in the hypothalamus. The gene responsible for its production is located on the 7th chromosome. Both leptin gene polymorphism rs7799039 or leptin receptor gene polymorphism rs137101 have clinically relevant single nucleotide polymorphisms. Despite numerous studies, there is no explicit proof of any relationship between the leptine gene polymorphism and obesity [12]. Various genetic models have been studied, including models with dominant and co-dominant expression. On the other hand, new studies have shown a relationship between the leptine gene polymorphism and arterial hypertension, coronary heart disease, or LDL cholesterol concentration [13, 14].

Cadmium exposure has the most evident effect on leptine concentration – cadmium causes the emergence of abnormal, small adipocytes and reduces secretion of leptine, which leads to an increase in insulin resistance, obesity and the development of metabolic syndrome [15]. However, the role of genetic variability in the context of cadmium exposure remains unknown and the authors were unsuccessful in finding studies which consider both genetic variability and cadmium exposure.

CB1. Endogenous cannabinoids, which affect the cannabinoid receptors type 1 (CB1), have similar activity compared to leptine in the context of metabolic syndrome. They are located mostly in different parts of the brain and spine, with little expression in other organs. Their expression prevents development of undue neurological activity. As a result, they are responsible for the feelings of satiety and hunger. Genetic variability of several locations of the gene responsible for the CB1 creation have been described. The best known example is the transformation of adenosine into locus rs 1049353, which causes higher BMI and bigger waist circumference [16, 17]. There is no obesity without excessive calories in the food. Hence, it is likely that individuals with a specific genetic configuration who eat more food also consume more arsenic and are exposed to its toxic effects. As mentioned before, arsenic directly leads to the development of insulin resistance, and the main route of exposure in the general population...
is food. As commonly known, exogenous cannabinoids stimulate appetite and, as a result, arsenic intake.

Another problem is the accumulation of arsenic and other heavy metals in cannabis plants. These plants accumulate significant amounts of heavy metals, and using its products, either legally or illegally, is connected with heavy metal exposure. In cannabis sativa this accumulation is so great that it is used in the process of phytoremediation – the process of purifying soil from heavy metals by small herbs and plants. An adverse synergy is very likely – a disadvantageous genetic constellation – a genetic predisposition to excessive calorie intake and increased exposure to arsenic, which jointly lead to the development of insulin resistance. However, there are no studies to prove this explicitly.

**As3MT.** Upon entering the organism, arsenic undergoes methylation and partial reduction, and is then expelled with urine. This process also creates oxygen free radicals and more toxic forms of arsenic, such as monomethylarsonic acid. The main participating enzymes are arsenite methyltransferase (As3MT), purine nucleoside phosphorylase (PNP), or glutathione S-transferase omega-1 (GSTO1) [18]. The genetic variability of the genes responsible for these enzymes leads to various concentration of arsenic in the exposed population, as well as changes in the ratios of individual arsenic forms excreted with urine. It has been demonstrated that certain polymorphic forms of the As3MT gene are related to higher concentrations of dimethylarsenates [19]. In the Mexican population, As3MT gene polymorphisms – in locus rs12767543, rs3740393 and rs11191453 of chromosomes 10, were all associated with a high uDMA/uMMA ratio. Similar dependences are observed in the Polish population. In turn, higher concentration of the dimethylarsenates increases the insulin resistance by blocking the insulin-dependent transported GLUT4 [20].

Increased insulin resistance is the onset of metabolic syndrome which, after several years, leads to complications and vascular disease, particularly coronary heart disease. As demonstrated by Gong, even a minor, chronic exposure to arsenic, such as drinking ground water, increases the risk of coronary heart disease, the same as the adverse variant to arsenic, such as drinking ground water, increases the risk of coronary heart disease. Designation of the expression this relationship shows how arsenic directly contributes to an increase in cardiovascular risk [28]. The discovery of the aforesaid miRNAs. At the same time, there is a reported dependence between polymorphisms – in this case, original number of new reports increases every month, unlike the dependence between polymorphisms – in this case, original and contributing studies are scarce. For this reason, a detailed description of all the miRNAs which number several dozen, exceeds the scope of this study. Explicit examples have been selected which demonstrate the significance of miRNA in the context of environmental and occupational exposure. In potable water there is a clearly visible effect of arsenic exposure on the expression of miRNA 155 and 126 – the higher the arsenic concentration, the greater the expression of the aforesaid miRNAs. At the same time, there is a reported relationship between the expression of the aforesaid miRNAs and the increase in cardiovascular risk [28]. The discovery of this relationship shows how arsenic directly contributes to an increase in cardiovascular risk. Designation of the expression of these miRNAs allows identification of individuals who are both potentially exposed to arsenic and bear an increased cardiovascular risk.

**ALAD.** Lead is an inhibitor of several enzymes which participate in the haem synthesis. Its inhibiting effect is the strongest for the delta-aminolevulinic acid dehydratase, displacing zinc from the active enzyme location. Therefore, polymorphism of the gene responsible for production of the delta-aminolevulinic acid is a factor determining the concentration of lead in the serum and its potential toxic effect [22]. Higher lead concentration is also related to the development of metabolic syndrome [23]. It was also found that mixed exposure to heavy metals, including lead, is also related to the development of obesity, arterial hypertension and type II diabetes [7]. The authors, however, failed to find any research investigating a direct connection between the ALAD gene polymorphism and the metabolic syndrome, although this connection appears to be likely.

**FTO.** The FTO gene (fat mass and obesity-associated gene), commonly known as the obesity gene, is probably the most examined genetic mutation which contributes to the development of obesity. It codes the enzyme alpha-ketoglutarate dioxygenase responsible for the demethylation of numerous nucleic acid molecules. The latest meta-analyses confirm that the adverse variants of the FTO gene polymorphism are significantly related to obesity [24]. On the other hand, dietary interventions has the same effect, irrespective of one’s polymorphic variants, which suggests that environmental interventions and lifestyle prevail over the innate genetic variability [25]. At the same time, there is no known connection between the FTO gene polymorphism and exposure to metals, although, again, one may expect an adverse synergy in the context of metabolic syndrome development.

**MICRO-RNA.** Apart from polymorphism of individual nucleotides, the literature also broadly discusses the participation of the micro RNA (miRNA) in pathogenesis and development of numerous diseases, e.g. tumours, cardiac and vascular diseases, diabetes and insulin resistance, or immunologic disorders.

MicroRNA is a group of small, non-coding RNA which, in mature form, regulate the gene expression at the post-transcription level. They play an important role in the processes crucial for the proper functioning of cells, and may affect the signal-transferring process and be a part of the mechanism defending the organism against viruses [26].

Numerous miRNA have been identified, both with pro-adipogenic and anti-adipogenic effect. Currently, it is believed that distortion of the miRNA balance leads to the development of obesity and increased insulin resistance, and individual miRNAs are a potential therapeutic target in obesity treatment [27].

The effect of arsenic, lead and cadmium exposure on the miRNA variability is currently widely examined. The number of new reports increases every month, unlike the dependence between polymorphisms – in this case, original and contributing studies are scarce. For this reason, a detailed description of all the miRNAs which number several dozen, exceeds the scope of this study. Explicit examples have been selected which demonstrate the significance of miRNA in the context of environmental and occupational exposure.

In potable water there is a clearly visible effect of arsenic exposure on the expression of miRNA 155 and 126 – the higher the arsenic concentration, the greater the expression of the aforesaid miRNAs. At the same time, there is a reported relationship between the expression of the aforesaid miRNAs and the increase in cardiovascular risk [28]. The discovery of this relationship shows how arsenic directly contributes to an increase in cardiovascular risk. Designation of the expression of these miRNAs allows identification of individuals who are both potentially exposed to arsenic and bear an increased cardiovascular risk.
to have a potential impact – by an increased expression of individual miRNAs – on the development of large intestine cancer, type 2 diabetes, and damage of the signal pathways in the immune system [29]. Studies on cellular cultures confirmed that it is arsenic, particularly in mixed exposure to arsenic, lead and cadmium, that induces an increased miRNA expression, not any other factors [30].

CONCLUSIONS

The designation of micro-RNA expression seems to be a promising direction not only in the diagnostics of exposure to arsenic, but also to other metals, such as lead and cadmium. It not only permits the isolation of risk groups, but also an almost direct diagnosing of organ damages caused, for instance, by exposure to arsenic. Further studies of miRNA diversity are still needed which would confirm their significance and isolate specific changes with the highest potential for clinical use. At the same time, one should not forget about the polymorphism of individual nucleotides of the genes responsible both for the development of the metabolic syndrome, and for arsenic metabolism. Although these changes have been known for many years, to-date there has been little success in connecting the relationship between polymorphism of the genes responsible for the development of the metabolic syndrome, and exposure to arsenic and certain heavy metals.

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