OBESITY IN CHILDHOOD AND ADOLESCENCE,
GENETIC FACTORS

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ABSTRACT

Obesity and excess weight are a pandemic phenomenon in the modern world. Childhood and adolescent obesity often end up in obesity in adults. The costs of obesity and its consequences are staggering for any society, crippling for countries in development. Childhood obesity is also widespread in Macedonia. Metabolic syndrome, dyslipidemia and carbohydrate intolerance are found in significant numbers. Parents and grandparents are often obese. Some of the children are either dysmorphic, or slightly retarded. We have already described patients with Prader-Willi syndrome, Bardet-Biedl syndrome or WAGR syndrome. A genetic screening for mutations in monogenic obesity in children with early, rapid-onset or severe obesity, severe hyperphagia, hypogonadism, intestinal dysfunction, hypopigmentation of hair and skin, postprandial hypoglycaemia, diabetes insipidus, abnormal leptin level and coexistence of lean and obese siblings in the family discovers many genetic forms of obesity. There are about 30 monogenic forms of obesity. In addition, obesity is different in ethnic groups, and the types of monogenic obesity differ. In brief, an increasing number of genes and genetic mechanisms in children continue to be discovered. This sheds new light on the molecular mechanisms of obesity and potentially gives a target for new forms of treatment.

Keywords: obesity, children, adolescents, genetic causes

INTRODUCTION

Worldwide prevalence of obesity and childhood obesity is in constant and steep increase. Obesity has become pandemic. It affects at least 250 million people (7% of the estimated current world population), while at least 2-3 times more people are overweight [1].

Children are affected, too. 21-24% children and adolescents in USA are overweight and another 16-18% is obese. The prevalence of overweight children and adolescents in the United States has increased by 50-60% in a single generation, while the prevalence of obesity has doubled. Australia, Canada, France, Germany, Brazil, Chile, Finland, France, Germany, Greece, Japan, the UK, and the USA saw the prevalence doubled or tripled between the early 1970s and late 1990s [1, 2].

There is an ethnic and racial difference in the prevalence of obesity. In American Indians,
Hispanics, Hawaiians, Hispanics, and blacks obesity is 10-40% higher than in whites [3]. Moreover, the majority of adults in some societies are overweight. In the United States, 61 percent of all adults are overweight. In Russia, the figure is 54 percent; in the United Kingdom 51 percent; and in Germany 50 percent. For Europe as a whole, more than half of those between 35 and 65 years of age are overweight [1, 2].

It is of note that the adolescent obesity is predictive of adult obesity: 80% of teenagers who are obese continue on to be obese as adults. The prevalence of obesity is high in Macedonia, too [4]. There is also a sign of hope: the 2006 review suggests that the increase in childhood obesity in the USA, the UK, and Sweden might be abating [5, 6, 7].

This explosion of obesity is probably a result of historical convergence. Early on humans with parsimonious caloric intake had a biological advantage as food was scarce and starvation common. Later one, this "thrifty gene" was massively challenged by the abundance of food in most of the developed world.

Obesity is costly for any society: some estimates suggest that the management of obesity in the USA costs approximately $100 billion yearly.

Criteria
In children, effects of age, sex, puberty, and race or ethnicity on growth make classification difficult. Methods based on weight, weight-height, skinfold thickness have advantages and disadvantages. The World Health Organization (WHO) bases its criteria for obesity in childhood and adolescence on BMI. BMI that is greater than the 85th (overweight) or the 95th (obesity) percentile, for age-matched and sex-matched control subjects. Overweight, obese, and morbidly obese refer to children and adolescents whose weights exceed those expected for heights by 20%, 50%, and 80-100%, respectively.

The International Obesity Taskforce (IOTF) international standard growth chart enables global comparison of prevalence [8]. Many countries continue to use their own country-specific charts [9]. The dominantly used thresholds for being overweight or obese in childhood are: 110% or 120% of ideal weight for height; weight-for-height Z scores of higher than 1 or higher than 2, and BMI at the 85th, 90th, 95th, and 97th percentiles (on the basis of international or country specific reference populations). It is of note that, the IOTF classification has high specificity, but low sensitivity [10].

There is a modification of those criteria for adults: grade 1 overweight (overweight) is a BMI of 25-29.9 kg/m2, grade 2 overweight (obesity) is a BMI of 30-39.9 kg/m2 and grade 3 overweight (severe or morbid obesity) is a BMI greater than or equal to 40 kg/m2. In addition, there are surgical definitions which describe BMI greater than 40 kg/m2 as severe obesity, a BMI of 40-50 kg/m2 is termed morbid obesity, and a BMI greater than 50 kg/m2 is termed super obese.

Pathophysiology
This is an energy imbalance between excessive energy intake and/or reduced energy expenditure. Sedentary lifestyle with excessive television viewing and/or excessive computer use coupled with insufficient physical activity results in obesity in children and adolescents. In infancy, excess fat deposition occurs when excess energy is provided, especially when the protein-to-energy ratio is altered. Excess weight in children depends on both genetic and environmental factors.

Etiological factors are many. As much as >90% of cases are idiopathic, <10% are associated with hormonal or genetic causes [1]. Hormonal disorders: growth hormone deficiency and growth hormone resistance, hypothryoidism, leptin deficiency or resistance to leptin action [11, 12, 13], glucocorticoid excess (Cushing syndrome), prolactin-secreting tumors, precocious puberty, polycystic ovary syndrome (PCOS) all can be manifested with obesity. Some medications cause excess weight/obesity: glucocorticoids, oral contraceptives, insulin, sulfonylureas, risperidone, thiazolidinediones, clozapine, tricyclic antidepressants, monoamine oxidase inhibitors (MAOIs: e.g phenelzine).

Other causes
Inflammatory and infective etiology may exist for obesity: adenovirus 36 infection is associated with obesity in chickens and mice. In humans, those who are not obese have a 5% prevalence of adenovirus 36 infection, while those who are obese have a prevalence of 20-30%.
MENDELIAN OBESITY

Mendelian pattern of inheritance is observed in some types of obesity [14].

Syndromic monogenic obesity

Syndromic monogenic obesity is rare and featuring obesity and mental retardation, dysmorphic features and organ-specific abnormalities [14]. Prader–Willi syndrome (PWS), has an incidence of 1 in 15000–30000 [15]. Patients have hypotonia, feeding difficulties, poor growth and delayed development in the first year of life. Later, patients develop hyperphagia, childhood obesity, short stature and cognitive disability [15]. In PWS only the paternal gene copies are expressed [16]. Paternal deletions are found in 65–75% of the patients [17], while maternal uniparental disomy is found in 20–30% [17]. In 1-3% of the patients there are imprinting defects, caused by epimutations or incomplete processing of the imprint from the father or from microdeletions in the DNA imprinting centre [17]. Several genes are implicated with PWS: makorin ring finger protein 3 (MKRN3), Necdin (NDN), nuclear pore associated protein 1 (NPAP1), SNRPN upstream reading frame (SNURF-SNRPN), MAGE family member L2 (MAGE-L2), and 5 small nucleolar RNA [18, 19, 20, 21, 22, 23].

Bardet–Biedl syndrome (BBS) is an autosomal recessive ciliopathy with retinal degeneration, polydactyly, cognitive disability, and genital and renal anomalies [24]. Obesity occurs mostly in the early years of life and sometimes results in T2D [25]. There are at least 19 BBS genes [18, 19, 20, 26]. Interestingly the BBS proteins mediate leptin receptor (LEPR) signaling [27]. It has been suggested that children with BBS at the initial work up should undertake imaging studies of the kidney and urinary tract. Also, in order to prevent end-stage renal disease (ESRD), close renal follow up from an early age of life is proposed [28].

Albright hereditary osteodystrophy (AHO), or pseudohypoparathyroidism Ia, is an autosomal dominant disorder with clinical features of hyperphagia, obesity, mental retardation, short stature, round facies and skeletal anomalies [32]. AHO is caused by the mutations in the guanine nucleotide binding the alpha-sub-unit of the stimulatory G protein (Gsα). It mediates the actions of hormones, neurotransmitters and paracrine/autocrine factors [33], resulting in resistance to the parathyroid hormone, thyroid stimulating hormone and gonadotropins [34].

Cohen syndrome, or obesity-hypotonia syndrome has characteristic facial features, microcephaly, hypotonia, non-progressive psychomotor retardation, motor clumsiness, progressive myopia and truncal obesity [35]. This is an autosomal recessive disorder caused by mutations in the Cohen 1 (COH1/VPS13B) gene [36].

Kabuki syndrome is characterized by specific face, mental retardation, visceral and skeletal malformations, growth deficiency, obesity and endocrinological anomalies [37]. Heterozygous mutations in the gene lysine (K)-specific methyltransferase 2D (KMT2D/MLL2) as the cause of Kabuki syndrome in 56–76% of the cases [38, 39, 40]. Infrequently, the syndrome has been linked to a mutation in the lysine (K)-specific demethylase 6A (KDM6A) gene [39].

Borjeson–Forssman–Lehmann (BFL) is characterized with severe mental disability, microcephaly, epilepsy, hypogonadism, obesity and gynecomastia [41]. This is an X linked disorder, caused by the mutations in the PHD finger protein 6 (PHF6) gene [42]. Not all patients carry mutations in PHF6.

Carpenter syndrome, or acrocephalopolysyndactyly type II, is characterized with acrocephaly, preaxial polydactyly, soft tissue syndactyly, brachy- or agenesis mesophalangy of the hands and feet, congenital heart disease, hypogonadism, obesity, umbilical hernia and mental retardation [43]. This is an autosomal recessive disorder, caused by the mutations in the RAB23 gene [44, 45].

Smith–Magenis syndrome (SMS) is a neuro-behavioral disorder characterized with obesity, sleep disturbance and multiple developmental anomalies [46]. This is an autosomal dominant disorder caused by heterozygous mutations in retinoic acid induced 1 (RAI1) gene.

Wilms tumour, aniridia, genitourinary abnormalities and mental retardation (WAGR) syn-
drome is caused by 11p13 deletions [47]. Obesity has been observed in approximately 30% of WAGR patients [48]. Most of obese WAGR patients have altered brain-derived neurotrophic factor (BDNF) gene [47]. Fluorescence in situ hybridization (FISH) study of a 8.5 year-old girl revealed a deletion of the WT1 and PAX6 gene in the 11p13 WAGR region [49].

NON-SYNDROMIC MONOGENIC OBESITY

Non-syndromic monogenic obesity refers to a single gene disorder.

Leptin. A mutation in the leptin (LEP) gene with truncated transcription of leptin was described in two severely obese cousins within a highly consanguineous family of Pakistani origin [50]. Other reports followed describing patients with no detectable leptin, in Pakistan, Turkey and Egypt [51, 52]. The patients with a homozygous LEP mutation had detectable circulating leptin levels, indicating the existence of a bio-inactive leptin protein [53].

LEPR deficiencies were found in severely obese siblings [54], in patients with severe early-onset obesity with mutations in LEPR [55], having high serum levels of leptin which indicated receptor insensitivity [55]. Further reports found Pakistani patients [56], patients in French population from Reunion Island [57].

Patients with mutations in LEP or LEPR had severe hyperphagia, rapid weight gain within the first year of life, and intolerant behavior when food restrictions were demanded [58]. Hypo-gonadotrophic hypogonadism is frequent [55]. Those children have defective T-cell mediated immunity, with high rates of infection and mortality [55]. Strikingly, loss-of-function mutations in LEP and LEPR have low blood pressure, despite obesity [59].

Leptin treatment in a girl with leptin deficiency resulted in weight reduction, reduced energy intake and increase in gonadotropin concentrations [60].

Loss-of-function mutations in SH2B1 an regulator of leptin, led to severe early-onset obesity, hyperphagia, IR, reduced height and behavioral abnormalities [61].

Proopiomelanocortin (POMC) deficiency in humans has the following characteristics: obesity, adrenal insufficiency, red hair, skin hypopigmentation, neonatal hypoglycemia, seizures, cholestasis and voracious appetite [62, 63, 64]. Severe motor and mental retardation was also reported [65].

Deficiency in prohormone convertase 1 (PC1/3), results in early-onset obesity, hyperphagia, postprandial hypoglycemia and other endocrine dysfunction. The main reason is that its role in the cleavage of proinsulin into insulin and POMC into alpha-melanocyte-stimulating hormone (α-MSH) is lacking [66, 67, 68]. Null mutations have diarrhea and diabetes insipidus [66, 68, 69, 70, 71], while a nonsense loss-of-function mutation at the heterozygous state causes familial obesity associated with glucose intolerance/diabetes [72].

Melanocortin 4 receptor (MC4R) mutations cause autosomal dominant obesity. It is of note that not all heterozygous carriers of MC4R become obese, but homozygous all have early-onset obesity [73]. MC4R deficient patients display hyperinsulinemia, increased linear growth, and an increase in bone mass in both children and adults [74, 75, 76]. Additionally, patients experience an increase in both fat and lean mass, which is not observed in other forms of monogenic obesity [77].

The neurotrophic tyrosine kinase receptor type 2 (NTRK2) missense mutation was found in a boy with early-onset obesity, hyperphagia, developmental delay, impairment in short-term memory and impaired nociception [78]. There was an alteration of the BDNF stimulated protein kinase phosphorylation, too [78]. In addition, loss of one functional copy of BDNF manifested hyperphagia, severe obesity, cognitive impairment and hyperactivity [79].

Single-minded homologue 1 (SIM1) has an essential role in formation of the paraventricular nucleus (PVN) of the hypothalamus [80, 81]. SIM1 haploinsufficiency leads to hyperphagia, obesity and reduction in the PVN [82]. SIM1 haploinsufficiency led to excessive growth, severe early-onset obesity [83]. Heterozygous deleterious mutations in SIM1 were observed in obese children with additional Prader–Willi-like neurobehavioral features [84, 85, 86].

KSR2 loss-of-function mutations were found in patients with hyperphagia, early-onset obesity, low heart rate, reduced basal metabolic rate and severe IR [87]. Mutations in Tubby bipartite transcription factor (TUB) were observed in patients with deteriorating vision, obesity and normal lucose/cholesterol/triacylglycerol levels [88].
OLIGOGENIC OBESITY

In contrast to monogenic obesity which is rare, obesity in humans carrying heterozygous deleterious coding mutations in these genes is significantly more frequent but not fully penetrant [89]. Based on loss-of-function mutation frequency of MC4R in the population of United States (US) (0.07 %), 426701 heterozygous MC4R carriers compared with 149 homozygous carriers (N =305000000) are probable. Partial MC4R deficiency may explain obesity in 256021 individuals, whereas complete MC4R deficiency may be the cause of obesity for only 149 subjects in the US population [90]. Those humans have also interaction with “obesogenic” environment [91, 92]. In different ethnic communities there is a prevalence of 0.2–5.6% of MC4R heterozygous, heterozygous compound and homozygous loss-of-function mutation carriers [73].

The same is true for the heterozygous loss-of-function mutations in POMC which result in a non-fully penetrant form of obesity 93, 94, 95]. Partial deficiency of LEP and LEPR has been found in humans with a higher percentage of body fat mass [55, 96].

POLYGENIC OBESITY

In polygenic obesity multiple gene defects interact with the environment [97] e.g. A three SNP haplotype in ectonucleotide pyrophosphatase/phosphodiesterase 1 (ENPP1) was found to contribute to childhood and adult obesity in a recessive model in European populations [98, 99]. K121Q has also been associated with adult obesity [100].

Since the discovery of FTO, many other loci that contribute to BMI, adult obesity, childhood obesity have been identified [101, 102, 103, 104]. GWAS have identified 135 variants associated with BMI level and/or obesity status. SNPs in most Mendelian non-syndromic genes (BDNF, NTRK2, LEPR, MC4R,PCSK1, POMC, SH2B1, TUB) and some Mendelian syndromic genes (SD-CCAG8, BBS4) have been proven to play role in polygenic obesity.

FROM GENOMICS TO CLINICAL PRACTICE

Strikingly, the US Supreme Court considers gene patenting as illegal [105]. However, some patents are still active, so identifying mutations must be done within a legal framework, using expensive patented genetic tests instead of whole genome/exome sequencing experiments [106, 107].

In children, no evidence was found for effects of 12 GWAS-based obesity marker alleles on weight regain [108], and only the FTO common variants [109] were associated with weight regain. However, these findings await further confirmation, and highlight the challenges of replicating gene–diet interactions in randomized clinical trials [110].

Case reports on carriers of homozygous LEPR and MC4R mutations showed lower weight loss and poorer outcomes after bariatric surgery [111, 112]. A more complex relationship has been reported for heterozygous MC4R mutations, showing no significant effects on bariatric surgery outcomes [112, 113]. In a study matching carriers of functional MC4R mutations or MC4R variants and two randomly paired controls without mutations, no difference in weight loss was observed [114], however, the design of functional characterization of mutations and variants was questionable [115]. Carriers of rare variants of MC4R matched with the MC4R reference allele carriers also demonstrated comparable weight loss [116]. In the Swedish Obesity Study, FTO was associated with maximum weight loss in gastric banding surgery subjects but not in gastric bypass subjects [117]. GWAS of gastric bypass subjects found that the 15q26.1 locus was significantly associated with weight loss [118]. However, larger studies, longitudinal analyses, and subsequent meta-analyses comprising of not only the genome, but also the epigenome and metagenome, are required to definitively establish whether treatment outcomes can be improved through assignment of patients to personalized surgical techniques.

The multifactorial origin of obesity gives rise to a variable response to anti-obesity medication, suggesting that efficacy of all new centrally active anti-obesity drugs [119] should be carefully assessed.
by using genomic information to ensure proper prescription and dispensing, in order to avoid unnecessary and potentially life-threatening side effects [120]. Identifying biomarkers for the development of diagnostics to guide prescriptions carries the potential of reducing adverse drug reactions and improving outcomes, while saving the healthcare system and patients from ineffective prescriptions.

CO-MORBIDITIES AND COMPLICATIONS

Obesity has a considerable impact on life quality, and some reduce life expectancy. Being overweight or obese between ages 14 and 19 years was associated with increased adult mortality (from age 30 years) from various systemic diseases [121]. Co-morbidities and complications are severe:


2. Respiratory: obstructive sleep apnea, Pickwickian syndrome), increased predisposition to respiratory infections, increased incidence of bronchial asthma.

3. GI: cholecystitis, cholelithiasis, steatohepatitis, fatty liver infiltration, reflux esophagitis.


5. Orthopedic: osteoarthritis, coxa vera, slipped capital femoral epiphyses, Blount disease and Legg-Calvé-Perthes disease, lumbago. Tibia vara (Blount’s disease) [122], slipped capital femoral epiphyses [123]. Alterations in the FTO gene, suggested that high-fat mass in children was associated with increased [124].

6. Metabolic: insulin resistance, hyperinsulinemia, type 2 diabetes mellitus, dyslipidemia. The rate of increase in BMI during adolescence may be a significant risk factor for diabetes [125, 126]. About half of the children with BMI higher than the 97th percentile have one or more of the disorders that make up the metabolic syndrome [127].

7. Reproductive: anovulation, early puberty, infertility, hyperandrogenism and polycystic ovaries in women, hypogonadotropic hypogonadism in men. There is an acceleration in timing of the larche and menarche in girls [128, 129], pubertal advancement in boys [130] and adverse effects on maturation [131] and alignment [132] of developing bones in both sexes. The advanced skeletal maturation has been attributed to increased adipose tissue aromatization. Pubertal timing might be altered by nutrition-related signals (eg, insulin and leptin) [133].


9. Surgical: increased surgical risk and postoperative complications.

10. Psychologic: social isolation, peer problems, depression.

11. Miscellaneous: reduced mobility, difficulty maintaining personal hygiene, Stress incontinence, Intertrigo (bacterial and/or fungal), acanthosis nigricans, hirsutism, increased risk for cellulitis and carbuncles, venous varicosities, lower extremity venous and/or lymphatic edema.

TREATMENT

The team approach to therapy (nurse educators, nutritionists, exercise physiologists, and counselors) is essential [134]. Family-based behavioral weight control is effective [135]. Physical activity should be encouraged and time spent watching television and playing computer games reduced. The quantity and the content of daily meals should be adjusted: less sugars, more vegetables and fruit. Improving fitness in children correlates with a lower incidence of obesity in adolescence [136]. The severe controlled–energy diets is not fit for children and adolescents.

Medication. Sibutramine may be classified as an anorectic drug, whereas orlistat’s [18, 19] mechanism of action involves induction of lipid malabsorption. A randomized placebo-controlled trial of sibutramine in adolescents resulted in a significant reduction in body mass index (BMI), without [20]. Benzphetamine (Didrex), phendimetrazine (Bontril), diethylpropion, and phentermine (Ionamin) have also been used.

Rhythm Pharmaceuticals (ESPE 2016) presented preliminary results from a phase 1b clin-
ical trial assessing the safety and efficiency of setmelanotide (RM-493), an MC4R agonist, in obese patients with a heterozygous genetic defect in MC4R. Patients lost weight and the treatment was well tolerated. The same company is now extending the clinical trial to patients with mono- genetic obesity mutations in POMC (ESPE 2016).

Surgical Care. Various bariatric surgical procedures have been used in some adolescents [137]. Usual criteria include patients >15 y, with a BMI of more than 40 or weight exceeding 100% of ideal body weight (IBW).

In the vertical-banded gastropasty (VBG), a pouch of 15-mL to 30-mL capacity is constructed, greatly reducing the amount of food that can be eaten. In gastric bypass, a larger pouch that empties into the jejunum is created [138]. Laparoscopic placement of an adjustable gastric band (LAGB) has supplanted the vertical-banded gastropasty as safer and reversible. LAGB positions a collar with an internal, saline-filled balloon around the upper stomach [139].

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Резиме

ОБЕЗНОСТА ВО ДЕТСТВОТО И АДОЛЕСЦЕНЦИЈАТА, ГЕНЕТСКИ ФАКТОРИ

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Обезноста и прекумерната тежина се пандемски феномени во модерниот свет. Адултната обезност често е резултат на обезноста во детството и во адолесценцијата. Трошоците поврзани со обезноста и компликациите што произлегуваат од неа се запрепастувачки, особено за земјите во развој. Обезноста во детството е, исто така, широко распространен и во Македонија. Метаболниот синдром, дислипидемијата и јаглехидратната интолеранција се присутни во значителен број. Родителите, бабите и дедовците се често со обезност. Некои од децата се или дизморфични или со благо заостанување во менталниот развој, како што ги имаме веќе опишано кај синдромите на Prader-Willi, Bardet-Biedl или WAGR.

Генетскиот скрининг за мутации за моногенетска обезност кај децата со ран и брз почеток на обезноста, односно екстремна обезност, тешка хиперфагија, хипогонадизам, интестинална дисфункција, хипопигментација на косата или кожата, постпрандијална хипогликемија, инсипиден дијабет, абнормални нивоа на лептин и постојење браќа и сестри со обезност или посно тело во семејството, открива многу форми на обезност.

Дополнително, како што обезноста покажува разлики во различни етнички групи така и типовите на моногенетска обезност се разликуваат. Накратко, сè поголем број гени и генетски механизми сè уште се откриваат кај децата. Ова отвора нови порти во молекуларните механизми на обезноста и дава нови потенцијални форми на третман.

Ключни зборови: обезитет, деца, адолесценти, генетски причини