Non-Alcoholic Fatty Liver and Metabolic Syndrome in Children: A Vicious Circle

Arianna Alterio    Anna Alisi    Daniela Liccardo    Valerio Nobili
Hepato-Metabolic Disease Unit and Liver Research Unit, ‘Bambino Gesù’ Children’s Hospital, IRCCS, Rome, Italy

Introduction

Non-alcoholic fatty liver disease (NAFLD) is the most common cause of chronic liver disease in Western countries and its worldwide prevalence in adults and children is increasing and becoming an obesity epidemic [1, 2]. Particularly, there is an alarming increase of number of children affected by NAFLD supported by high prevalence data ranging from 3 to 12% in the general paediatric population up to 70–90% in young obese [3].

Pediatric NAFLD is associated with several factors of metabolic syndrome (MS), like abdominal (central) obesity, dyslipidaemia (hypertriglyceridaemia and/or hypercholesterolaemia) and insulin resistance. Therefore, NAFLD can be considered as the hepatic manifestation of MS, even if there is no agreement that NAFLD contributes to MS and vice versa [4]. In fact, the aetiology and pathogenesis of both diseases are multifaceted and closely related to genetic predisposing factors, intrauterine environment and unhealthy lifestyle [5]. Bad eating habits and sedentarity may favour the onset of childhood obesity strongly contributing to make a child more prone to develop NAFLD and MS during adolescence [6]. Furthermore, as a large part of children with NAFLD and traits of MS remain undiagnosed, it is plausible that, in the coming years, adolescents with NAFLD characteristics may present a rapid course of disease towards more severe forms of liver damage (i.e. cirrhosis), MS and associated cardiovascular disease (CVD). In fact, during the
last decade, several epidemiological data have revealed a dramatic increase of the MS incidence in urban adolescents mainly due to overnutrition and a sedentary lifestyle [7].

In this scenario, the role of unhealthy lifestyle prevention is critical, particularly in subjects genetically predisposed to both diseases. Here, we provide an overview of current genetic, pathogenetic and clinical evidence of the vicious circle created by NAFLD and MS in children.

Methods

A PubMed/MEDLINE search of the literature in the last 5 years was performed, using the following search terms: NAFLD and MS and children. This search yielded 121 publications of which 35 were relevant recent clinical studies and 6 were review articles. To these we added 12 relevant articles on NAFLD and MS published between 2009 and 2013.

Characteristics That Define NAFLD and MS in Children

Non-Alcoholic Fatty Liver Disease

NAFLD is defined as an intrahepatic fat accumulation in people who drink little or no alcohol (>20 g/day). In general, it is considered a multifactorial liver disease that ranges from benign steatosis to the more severe non-alcoholic steatohepatitis (NASH), which is characterized in children by lobular inflammation and hepatocellular ballooning, and frequently coupled to fibrosis [2, 8].

During NAFLD pathogenesis, according to the ’multiple hits’ hypothesis, two primary hits including insulin resistance and free fatty acid (FFA) accumulation induce steatosis (>5% of hepatocytes, histologically) making the liver more susceptible to secondary hits, such as the action of adipocytokines, oxidative stressors and immune response that subsequently lead to NASH and fibrogenesis [9]. It is well recognized that diet, exercise, central obesity, insulin resistance, and hyperlipidaemia strongly influence the epigenetic dependent development and progression of NAFLD as well as MS. However, not all patients with this pedigree of risk factors for MS develop NAFLD. Therefore, it is not surprising that a possible genetically established predisposition, including single nucleotide polymorphisms (SNPs), may explain both the ethnic differences in the prevalence of disease and an increased risk for advanced NASH and fibrosis [2].

Metabolic Syndrome

MS in adults is defined by the International Diabetes Federation (IDF) as the presence of abdominal obesity (by measuring waist circumference) and two or more of the following factors: hypertriglyceridaemia, low HDL cholesterol, fasting glucose and hypertension (http://www.idf.org/idf-worldwide-definition-metabolic-syndrome). IDF proposed similar criteria for definition of MS in the paediatric population according to three age groups: from 6 to 10 years, from 10 to 16 years, and ≥16 years was considered as adults (http://www.idf.org/node/1405?unode=4A7F23CB-FA35-4471-BB06-0294AD33F2FC) [9].

The pathogenesis of MS is not completely understood but two factors are certainly crucial in its genesis: central obesity and insulin resistance. Central obesity is associated with hypertension, hypertriglyceridaemia and hyperglycaemia. Children with excess fat have a higher risk of CVD in adulthood [10]. In fact, when white adipose tissue (WAT) excess is localized in the upper parts of the body, defining central obesity, it represents a strong risk factor for metabolic and pro-inflammatory complications [11]. Therefore, even in the case of MS a genetic susceptibility to WAT accumulation concomitantly to lifestyle factors (i.e. diet and exercise) may lead to a specific lipid partitioning and consequent adipose tissue inflammation and increased FFA flux, which drive the various elements of disease including insulin resistance and affect endothelial function [12].

Role of Genetic Predisposition

A large part of the translational research is today devoted to the pursuit of genetic variants that can be used to identify subjects with a genetic susceptibility or resistance to multifactorial diseases, including NAFLD and MS. These genetic variants in the DNA code (known as alleles), in hetero- or homozygosity, may be associated with specific traits of these diseases acquiring a high predictive value as markers. Variants at a single DNA base-pair or SNPs have received a great attention as potential genetic markers. They have the advantage of a high frequency in the human genome (on average, 1 occurs in every 1,000 nucleotides) and are relatively easy to genotype using next-generation sequencing technologies. Although genetic susceptibility to NAFLD and MS are not completely known, it was suggested that their development may be associated with SNPs in genes involved in the control of insulin resistance, lipid metabolism, inflammation and oxidative stress [5].
During last 5 years, several potential genetic SNPs have been suggested as potential candidates for either paediatric NAFLD susceptibility or disease progression (i.e. NASH and fibrosis), including the rs738409 variant of adiponutrin/patatin-like phospholipase domain-containing 3 (PNPLA3) coding gene, the rs1801278 variant of insulin receptor substrate-1 (IRS-1) coding gene, the rs3750861 variant of Kruppel-like factor 6 (KLF6) gene and the rs4880 variant of manganese-dependent superoxide dismutase (SOD2) gene. Genome-wide association studies have identified many SNPs associated with a large number of conditions related to MS [13]. However, only recently it has been reported that the rs1800849 variant of uncoupling protein 3 (UCP3) gene associates with MS components and increased risk of Chinese children with NAFLD [14]. Moreover, as the severity of obstructive sleep apnoea (OSAS) also correlates with the increased risk of paediatric NAFLD and MS, the presence of selective SNPs in the gene encoding for fatty acid-binding protein 4 (FABP4) might explain the susceptibility to OSAS in both diseases [15, 16].

**Role of Central Obesity-Dependent Inflammation and Insulin Resistance**

Central obesity correlates with an accumulation of visceral fat, which in turn depends on adipose tissue dysfunctions. In fact, adipose tissue is now considered like an ‘endocrine’ organ, that may produce and release circulating factors, called adipocytokines, that are involved in the alterations of MS. In fact, in adipocytes, bad dietary intake (i.e. elevated consumption of sweet high-fat foods) causes a specific pattern of lipid storage and metabolic stress, which in turn activate signalling cascades that induce oxidative stress and trigger an inflammatory response. Therefore, metabolic stressors and lipid partitioning may result in the release of several circulating adipocytokines with local action (i.e. on macrophages), and systemic effects such as muscle and liver insulin resistance. Furthermore, these circulating factors including tumour necrosis factor α (TNF-α), interleukin 6 (IL-6), leptin, adiponectin, retinol-binding protein-4 (RBP4) and resistin play a critical role in inflammation that often characterized subjects with insulin resistance, NAFLD and MS predisposition [5, 17, 18]. This nexus between NAFLD and MS, summarized in figure 1, is supported by clinical evidence reported in children.

**Evidence of NAFLD and MS Nexus in Children**

Over the last decade there has been increased interest in the determination of possible pathogenetic and clinical relationships between NAFLD and MS in obese subjects during childhood [5, 19]. A retrospective study including 43 American children with biopsy-proven NAFLD has provided the first evidence of NAFLD/MS association at paediatric age reporting that approximately 95% of patients with liver disease were obese and were insulin-re-
sistant [20]. Accordingly, Manco et al. [21] have demonstrated that 92 and 84% of 197 Caucasian children with NAFLD displayed a body mass index (BMI) >85th percentile and waist circumference ≥90th percentile, respectively. The strong correlation between central obesity and elevated waist circumferences has also been confirmed by a case-control study conducted on 150 overweight/obese children with biopsy-proven NAFLD and 150 without [22]. Moreover, the same authors found that among the subjects studied, MS showed 5.0 times the odds of having NAFLD compared to without the metabolic disease.

Further cross-sectional studies conducted on different ethnic groups have supported the connection between NAFLD and MS in the paediatric population. Kelishadi et al. [23] have in fact reported that in 1,107 Iranian subjects (aged 6–18 years) central obesity may be used as a predictor of NAFLD assessed by ultrasound and surrogate markers, such as alanine aminotransferase (ALT). Accordingly, Wei et al. [24] have demonstrated that approximately 16% of a cohort of obese British children display an increase of serum ALT levels, elevated BMI and altered glucose homeostasis. It has also been reported that in Chinese obese children the overlapping between NAFLD and MS may be found in 84.61% of cases [25]. Interestingly, Deboer et al. [26] have found that ALT elevation varied by race/ethnicity demonstrating that non-Hispanic Black adolescents exhibit a relationship between insulin resistance and ALT lower than in non-Hispanic White and Hispanic age-matched subjects, supporting the weight of genetic predisposition in the link between NAFLD and MS. More recently, two other prospective studies have confirmed a positive association between elevated ALT levels and MS prevalence in overweight and obese Hispanic children observed in tertiary care centres [27, 28]. Finally, very recently a cross-sectional study on 182 obese sedentary children and adolescents (6–16 years) has documented the significant correlation between intra-abdominal fat and NAFLD (p = 0.005), MS (p = 0.013), dyslipidaemia (p = 0.001) and HOMA-IR (p = 0.007) [29].

Besides central obesity, adipocytokine circulating levels have also been demonstrated as a common feature between NAFLD and MS in children. Interestingly, among these adipocytokines, leptin serum levels increase in concert to steatosis grade, ballooning, inflammation and fibrosis severity, indicating that hyperleptinaemia in NAFLD children could also be a prerequisite for insulin resistance and other MS features in these subjects [29]. On the contrary, adiponectin and resistin act as anti-inflammatory molecules, because their circulating levels display a negative correlation with liver steatosis assessed by abdominal ultrasound [30]. Moreover, Leibnitz et al. [31] have clarified that only hypoadiponectinaemia is significantly associated with a decreased risk of NAFLD and insulin resistance. As adiponectin and leptin may control the expression and secretion of pro-inflammatory cytokines such as TNF-α and IL-6, it is expected that these adipose tissue-dependent circulating molecules also associate with NAFLD and insulin resistance [32]. The real role of these two adipocytokines in the development and progression of NAFLD in children remains to be defined. However, Mager et al. [33] reported that meals high in saturated fat may lead to postprandial dyslipidaemia and hyperinsulinaemia concomitantly to a change in pro-inflammatory and lipoprotein pattern in obese children with and without NAFLD.

In addition to these commonly studied adipocytokines, the circulating levels of RBP4, which are strongly associated with insulin resistance, also display an inverse association with liver damage progression in paediatric NAFLD [34]. On the contrary, Boyraz et al. [35] have recently demonstrated that RBP4 levels are increased in 63 obese children with ultrasound and ALT elevation evidence of NAFLD.

Although several differences exist in the power of correlation of these adipocytokine levels with the severity of NAFLD and MS traits, recent clinical evidence reinforces the role of these circulating molecules as biomarkers of both diseases [36–38]. Noteworthy, Walker et al. [39] have recently suggested the connection between WAT inflammation, adipocytokine levels and NAFLD severity in children, demonstrating that the increasing obesity is, on the one hand, strongly related to increased intrahepatic lipid deposition and adipocytokine-dependent activation of Kupffer cells explaining the progression from NAFLD to NASH, and, on the other hand, it is also significantly associated with markers of adipose tissue damage via crown-like structure resembling the process of liver fibrogenesis.

Role of Intrauterine Environment in NAFLD and MS Cross-Talk

Intrauterine growth retardation (IUGR) is the most important cause of perinatal mortality and morbidity, and is often used as a synonym of small for gestational age (SGA). Interestingly, several studies have highlighted the relationship between IUGR and the increased risk of MS in children [5]. The pathogenetic machinery that pro-
motes MS in IUGR remains unclear, even though the increase of insulin resistance appears to play a pivotal role by two potential mechanisms: (1) the establishment of an insulin-resistant genotype independently of intrauterine environment, and (2) the foetal reprogramming due to a thrifty phenotype hypothesis [40]. Furthermore, experimental studies have demonstrated that IUGR is associated with not only obesity and lipid abnormalities, but also with liver steatosis and inflammation [40]. Interestingly, Nobili et al. [41] have found in 90 Italian children an association between paediatric NAFLD with IUGR, independently of insulin resistance. Accordingly, Faienza et al. [42] more recently have reported 34.8% of NAFLD out of 23 SGA children demonstrating that the insulin resistance index significantly correlates with liver steatosis at ultrasound.

Although significant gaps still exist in the mechanisms inducing metabolic profile abnormalities and NAFLD in children, both diseases should be recognized as a coupled emerging health problem in IUGR/SGA prepubertal individuals.

**Emerging Role of Hepatokines and Thyroid Hormones**

Liver responds to adipose tissue inflammation not only with the histological damage that characterizes NAFLD but also with production and release of some circulating molecules called hepatokines, which may directly promote metabolic dysfunctions that lead to MS [43]. In particular, three hepatokines, including insulin-like growth factors (IGF)-I and -II, fibroblast growth factor 21 (FGF21) and fetuin-A, have been reported as novel prom-}

... support a pathogenetically plausible explanation given by the association between the common features of MS observed in both NAFLD and hypothyroidism [47]. Pacifico et al. [48] provide the first evidence between NAFLD, thyroid function and MS in childhood. These authors have demonstrated that thyroid function tests, particularly the thyroid-stimulating hormone (TSH) test, positively associate with NAFLD and metabolic variables in 402 consecutive overweight/obese children, suggesting TSH as a potential predictor of liver disease and lipid and glucose dysmetabolism, independently of visceral obesity. Accordingly, Torun et al. [49] have reported that free thyroxine and free triiodothyronine levels are comparable in obese children and age- and gender-matched subjects with normal weight, whereas TSH levels significantly increase correspondingly to the steatosis degree assessed by ultrasound, and correlate with ALT and BMI.

**NAFLD and MS Collaboration in Early CVD Development**

The concept that NAFLD and MS may represent overlapping diseases is also suggested by the fact that intrahepatic fat accumulation seems to predict more strongly than central obesity the likelihood that metabolic abnormalities are associated with early CVD in youth [10]. In fact, as demonstrated by several authors, children with NAFLD may present certain endothelial dysfunctions and greater carotid intima-media thickness, the latter considered a surrogate marker of early atherogenesis in childhood [50–53]. In fact, Nobili et al. [50] have shown for the first time that in children with NAFLD, the severity of liver injury is strongly associated with the presence of a more atherogenic lipid profile measured as triglyceride/high-density lipoprotein cholesterol (HDL), total cholesterol/HDL, and low-density lipoprotein cholesterol (LDL)/HDL ratios. Accordingly, Pacifico et al. [51] have demonstrated in a population study including 548 children that subjects with a high triglycerides/HDL-C ratio have an increased risk of insulin resistance and display an association between this ratio and NAFLD, independently of obesity.

Left ventricular (LV) dysfunction is a further abnormality in cardiac function that has recently been found to be associated with NAFLD and MS in children. In particular, Fintini et al. [52] have demonstrated that LV hypertrophy can be detectable early in NAFLD children even if it does not appear to be linked to other cardiovascular and metabolic alterations. Moreover, Pacifico et al.
[53] have found that asymptomatic obese children with NAFLD exhibit early LV diastolic and systolic dysfunctions that become more severe in NASH subjects.

Conclusions and Future Perspectives

The rapid prevalence of NAFLD and MS among children in recent years has made research urgent to gain the scientific knowledge required of genetic and molecular mechanisms in order to perform possibly early prevention and design efficient therapeutic approaches. Several lines of clinical evidence have demonstrated that a vicious circle exists between NAFLD and MS in children. It is plausible that a common genetic predisposition may provide fertile soil for the complex molecular overlap of multiorgan response to different intrauterine and postnatal environmental factors, even though much remains to be explored in this field. Furthermore, as NAFLD and MS phenotypes are associated with a high risk of CVD, such information may be critical for the development of public health promotion strategies that approach children as future adults. In the meantime, as the adipose tissue-dependent inflammation and its cross-talk with the liver appears to be in some cases central for the NAFLD/MS connection, weight loss and weight control, coupled with a multidisciplinary approach to metabolic dysfunction and liver damage, should be considered by paediatricians the first-line treatment in the management of both diseases.

References

Non-Alcoholic Fatty Liver and Metabolic Syndrome in Children

Horn Res Paediatr 2014;82:283–289
DOI: 10.1159/000365192


