

Upshot of Epigenetics and Famine on Metabolic Syndrome

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Abstract

Chronic diseases are stimulated in the womb through adaptations acquired by the fetus in response to malnutrition. Maternal malnutrition during early pregnancy, in relation to low birth weight as well as uterine growth restriction, may adversely influence offspring metabolism and health. Parental nutritional imbalance, either through global nutritional manipulation or deficiencies in specific nutrients, predisposes the offspring to metabolic disease. Exposure to environmental factors in early life can influence the developmental process as well as long-term health in humans. The famine affected fertility, weight gain during pregnancy, maternal blood pressure, infant size at birth and development of the central nervous system, are associated with an increased risk of adult-onset metabolic syndrome. The point to ponder over here is how these risk factors interact at the cellular level so as to cause disease? Here, epigenetic epidemiology enables researchers to explore critical links between genomic coding, modifiable exposures and the manifestation of the disease phenotype. Extensive epidemiologic studies have suggested that adult disease risk is associated with adverse environmental conditions (famines) to which the mother is exposed to early in development.

Keywords: Upshot; Epigenetic; Famine; Malnutrition; Metabolic Syndrome

Introduction

It is now widely acknowledged that the jeopardy of chronic diseases in adulthood may have their origins in the womb through adaptations made by the fetus in response to under-nutrition [1]. The effects of under-nutrition depend upon the time of its occurrence during gestation and the organs as well as the systems that develop during that critical time window. The fetus being vulnerable to various factors like the physical and mental stress of the mother, environmental exposure, physical

activity and nutrition becomes susceptible to long term health risks which develop after birth [2]. The impact of maternal nutrition on the etiology of chronic diseases in offspring in their adult life has been well established by various epidemiological studies [3]. Nutritional imbalances, such as under and over-nutrition during critical periods of gestation, induce persistent physiological alterations.

Metabolic disorders such as obesity and type 2 diabetes have reached epidemic rates in most developed

and developing countries, but little is known about the role of DNA methylation in metabolic disease pathogenesis [4,5]. The Syndrome X or Metabolic Syndrome (MetS), is a cluster of phenotypes that include increased abdominal fat mass, impaired insulin responsiveness, dyslipidemia with increased plasma triglycerides and decreased HDL-cholesterol increased blood pressure and elevated circulating cytokines and adipokines [6]. The mechanisms by which Syndrome X leads to the development of insulin resistance, dyslipidemia and associated phenotypes are poorly understood in molecular level, but necessarily involve long-term changes in gene regulation and gene expression. Earlier studies reported that the prevalence of MetS or Syndrome X associated factors are relatively low during early childhood but increases during adolescence and thereafter tends to persist into adulthood [7-11]. Several lines of evidence support a role for non-genetic factors in the development of insulin-resistance and indicate that epigenetic factors, possibly through DNA methylation, may play a role in metabolic diseases. Epigenetics literally means "above" or "on top of" genetics. These modifications do not change the DNA sequence, but they affect how cells "read" genes. Epigenetics is the study of mitotically heritable alterations in gene expression potential that are not mediated by changes in DNA sequence and it is in the field of genetics of cellular and physiological phenotypic trait variations that are caused by external or environmental factors that switch genes on and off [12,13].

Epigenetic and Metabolic Syndrome

In the past decade, there has been a significant increase in the number of epidemiological studies investigating metabolic risk factors and outcomes in relation to DNA methylation, but many gaps stay behind in our understanding of the underlying cause. Previous studies reported that environmental factors can alter epigenetic features and change the future behavior of target cells and may play a role in susceptibility to MetS as well as other chronic diseases [14-16]. In humans this non genetic influences to change the pattern of gene expression are heritable across more than one generation and such transgenerational tradition of epigenetic states may contribute to the hereditary risk of various metabolic disorders [17-21]. DNA methylation is one of the most extensively studied epigenetic mechanisms and plays an important role in the process of development and differentiation [22]. It is also known that DNA methylation patterns continue to change after birth, at

least partly in response to environmental influences [23-25]. There is evidence from both animal and human studies that prenatal alimentary impairment can perpetually modify DNA methylation at several loci and these modifications have a pivotal role in the observed alteration of imminent risk of chronic diseases like obesity, insulin resistance and diabetes [26-32].

A family cohort study of Northern European descent, observed a significant portion of the epigenome is heritable, including genes known to play roles in obesity and Met S [33]. Increasing evidence shows that environment-induced genetic effects can pass transgenerationally without changes occurring in the primary DNA sequence and this epigenetic trait can be transmitted up to the fifth generation or more [34-36]. Some of the familial risks of MetS may be epigenetic in origin. One of the best-characterized epigenetically regulated loci is insulin-like growth factor II (*IGF2*) gene; is characterized by allele methylation pattern dependent on the nutritional stimuli received by the growing organism during early life development [37,38]. *IGF2* is a key factor in human growth and development and is maternally imprinted [39]. Further, childhood diet could contribute to *IGF2* loss of imprinting in individuals [38]. Imprinting is preserved by the hypomethylation of differentially methylated region (DMR) of *IGF2*, which ultimately progress for bi-allelic expression of that gene [26,40]. A recent investigation disclosed that paternal pre-conceptual obesity was associated with hypomethylation of *IGF2* in newborns [19].

There are several studies on humans and animals which suggest that the early nutrition and poor growth in-utero is associated with an increased risk of coronary disease, hypertension, type 2 diabetes and obesity in adulthood [41,42]. Maternal nutritional imbalance was shown to exhibit transgenerational effects through epigenetic and metabolic changes [43,44] and the most consistently-observed epigenetic association with adiposity has been with that of methylation at the *IGF2/H19* imprinting region [45,46]. Higher methylation levels at specific genes, including *IL10*, *LEP*, *ABCA1*, *GNASAS* and *MEG3*, were closely linked with nutrition metabolism, cardiovascular function and inflammation [47].

Famine and Metabolic Syndrome

The hardships of war (World War II) invoked thought that the body's reaction to starvation could be genetic.

However, at that time, it was not known that starvation of pregnant women affected their children. At present, with advanced technological aids as well as extensive knowledge, researchers have delved into the secrecy of inheriting life experiences. A panel of investigators discovered that the mothers who have experienced famine conditions undergo a change in their DNA, which passes on for up to three generations. It has been further speculated that the mechanism functions as a way for parents to "prepare their progeny for hardships similar to the ones that they have experienced", which would give them a better chance of survival. The first study about the possible association between prenatal exposure to famine and health reported during 1972-1975 used the "Dutch Famine 1943" to analyze adult health outcomes in relation to specific periods of gestation [48,49]. However, another study reported that exposure to famine in early and mid-gestation was associated with the prevalence of obesity [50]. Findings of The Dutch Hunger Winter Family Study suggest that the effects of famine can be observed some sixty years later [51].

The famine affected fertility, weight gain during pregnancy, maternal blood pressure, infant size at birth and central nervous system development are associated with an increased risk of adult-onset metabolic syndrome post birth [49,52-56]. It can be taken into consideration that the famine condition composes a 'natural experiment' in which outpouring to famine was attributed based upon an individual's birth place and time.⁵¹ This design was used to investigate how maternal undernutrition during specific gestational time windows may affect the subsequent life course of the offspring who has experienced the famine in-utero and their next generation. One of the most significant studies was conducted on children who were born to women exposed to severe undernutrition during pregnancy as a result of the Dutch Hunger Winter during World War II. The study reported reduced methylation of the imprinted gene IGF2 in these individuals during adulthood [26]. Two well-cited human population studies are those of the Dutch Famine cohort [57-59] and Overkalix cohorts [60], where both cohorts have been linked to transmission of ill-health into the F2 generation and the fetal origins hypothesis is broadly supported by the findings of the Dutch Famine birth cohort study. Another study from Northern Europe proposed that poor social conditions during childhood may become potent risk factors for obesity, diabetes and

other cardiovascular diseases (CVDs) in later life [61].

Upshot

Maternal under-nutrition during early pregnancy, in relation to low birth weight and uterine growth restriction, may adversely influence offspring metabolic health and are associated with an increased risk of adult-onset metabolic syndrome. On the other hand poor nutrition during pregnancy or post natal development may be a risk factor for irreversible health issues including obesity, type 2 diabetes, hypertension hypercholesterolemia and other metabolic diseases in adult life (Figure 1). To evaluate the function of maternal health and nourishment in the instigation and advancement of ailments in childhood as well as adulthood, it is necessary to identify the physiological and pathological roles of specific nutrients on the epigenome. An increased insight into dietary interventions in-utero as well as early life could modulate risk of diseases through epigenomic alteration. Early embryonic development is of special interest in this respect because it is a crucial period for establishing and maintaining epigenetic marks [62]. Gene-gene and gene-environment interaction are important processes for initiation of particular symptoms associated with metabolic syndrome and its progression. Mainly the explanation for the adaptation to the environment could be used as an example for epigenetics, especially, if epigenetic trans-generational inheritance exists [63,64]. However, a study systematically explained with the schematic diagram relating the genome, epigenome and environment with respect to trans-generational phenotypic characters [36].

The "*Barker hypothesis*" [65] or "*developmental programming hypothesis*" has opened up a new research paradigm for understanding chronic disease risk that has moved beyond the simplistic explanations based on genetic and lifestyle influences. A more integrated approach has developed which examines the relations between genetic inheritance and lifestyle factors, thereby incorporating the role of developmental plasticity i.e., the ability of changes in gene function to generate a range of phenotypic outcomes based on environmental exposures [66]. The hypothesis suggests that the influences of environment during early life of the offspring may induce susceptibility to the onset of obesity and related metabolic disorders during subsequent life post birth.

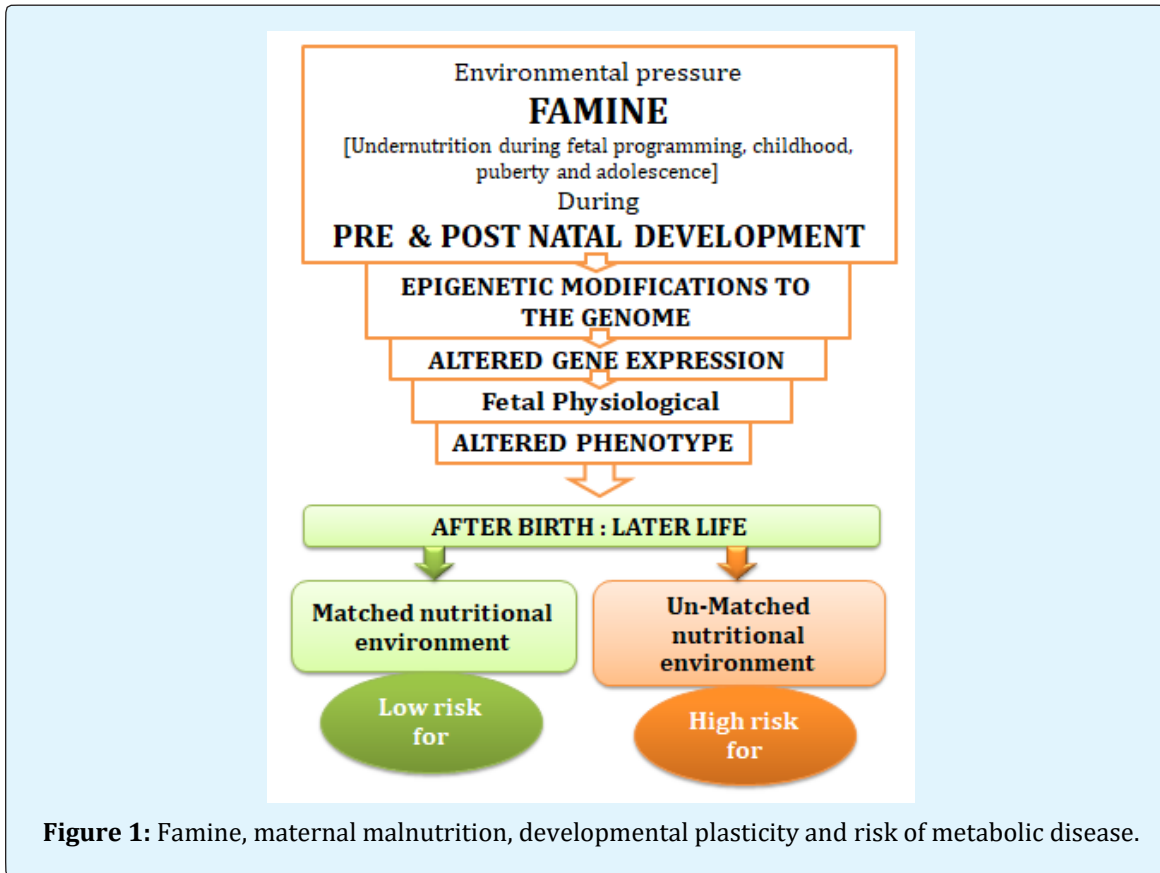


Figure 1: Famine, maternal malnutrition, developmental plasticity and risk of metabolic disease.

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Conflicts of Interest

The authors declare that they have no competing interests.

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