Role of diet in epigenetics: a review

Abstract

Epigenetics refers to the mechanisms that alter gene expression without altering the primary DNA sequence. The epigenetic modifications may be due to cytosine base methylation in DNA or by post-translational biochemical modifications of histones, the core proteins of nucleosomes. Epigenetic mechanisms are heritable and reversible. Epigenetic modifications can occur in response to environmental stimuli, of which the most important one is diet. Nutrients can modify physiologic processes through epigenetic mechanisms. The dietary nutrients enter metabolic pathways and get modified to simpler forms for the body to utilize. One carbon cycle is such a unique pathway which produces methyl groups to silence the genes. Nutrients like methionine, Vitamin B<sub>12</sub>, folic acid, and Vitamin B<sub>6</sub>, are the key components of this pathway. These methyl-donating nutrients in the diet can alter gene expression. The knowledge regarding nutritional epigenetics is still limited. As Epigenetics is reversible in nature, it is now considered as an attractive field of nutritional intervention. In the future, we need to investigate on more nutrients or bioactive food compounds to find the ones that can improve human health.

Keywords: epigenetics, diet, nutrients, one carbon cycle

Introduction

The word epigenetic seems to be a new word for us, but it has been studied a long while ago. The name epigenetics was coined by Conrad Waddington in 1942. “Epi”, in Greek, means “above” or “beyond”. Epigenetics was explained as a multifaceted developmental process connecting genotype and phenotype. DNA is the genetic blueprint to create a living organism, while the epigenetic information provides the message on how, where and when the genetic material should be used. Nutrients can modify physiologic processes through epigenetic mechanisms that alter the gene expression. The link between nutrition and epigenetics can be elucidated by metabolic pathways especially the one carbon metabolism. Nutrients from our food are modified in one carbon pathway that extracts methyl groups and then attaches them to our DNA or proteins. One carbon metabolism involving the folate and methionine cycle, integrates carbon units from amino acids and generates substrates for methylation of nucleic acid and protein. Nutritional deficiencies in folate, choline, methionine, vitamins B<sub>6</sub> and B<sub>12</sub>, perturb the complex regulatory network that maintains one-carbon metabolism. Polymorphisms in genes encoding the enzymes involved in metabolic pathways can also alter one carbon metabolism. Aegigenetics is reversible in nature, it is now considered as an attractive field of nutritional intervention. In this review, we are focusing mainly on the nutrients and metabolic pathways leading to epigenetic modification by DNA methylation.

Epigenetic mechanisms affecting gene expression

Nucleosomes, the fundamental repeating units of chromatin can be modified by many types of biochemical processes such as methylation, acetylation, phosphorylation, ubiquitylation, and sumoylation. The epigenetic modifications may be due to

i. cytosine base methylation in DNA

ii. post-translational modifications of histone proteins

iii. Positioning of nucleosomes along the DNA.

Epigenetic modifications can regulate the state of chromatin into a euchromatic (highly active) or heterochromatic (completely silent) state. Epigenetic modification of DNA at CpG islands modifies cytosine base and opposes transcription in most cases. Promoter regions with CpG islands become methylated during development, which results in long-term transcriptional silencing and inactivation of X-chromosome and imprinted genes which are classic examples of naturally occurring CpG island methylation during development. In the current scenario, DNA demethylation mechanism is gaining importance as it is necessary for cellular processes during development stages. The importance of the DNA methylation is that it facilitates the silencing of the imprinted genes and mainly focused on the role of early life nutrition causing epigenetic modifications. DNA methyltransferases (Dnmnts) are the enzymes those control DNA methylation and regulate gene expression patterns. Currently, five different Dnmnts are known: Dnmt1, Dnmt2, Dnmt3a, Dnmt3b and DnmtL. Dnmt1 are the maintenance DNA methyl transferase and Dnmt3a, 3b, and L are the de novo ones.

The histones are the major protein components of chromatin, and they have a significant role in gene regulation. The role of histone modifications in RNA synthesis has been very well reported. The five types of histones in chromatin are H1/H5, H2A, H2B, H3, and H4, and together with DNA, they form the nucleosome, which forms the chromatin. It is suggested that their modification can result in both repressive and stimulatory effects on genomic functions in vivo. It has been reported that histone modifications can either activate or repress the genes based on the type of residues which are modified and also the type of modifications occurred. By definition, epigenetic changes are heritable changes, which can be reversed. This heritability is established during differentiation stage and through multiple cycles of cell division it is being maintained with same genetic information, and thus the identity of the cells do not change.
Role of diet in epigenetics

Many scientists have studied the influential role of environment (mainly the diet) in changing the phenotype of an individual. The studies conducted in honey bees are one of the best examples to show the role of diet in epigenetics. Both fertile queens and sterile workers develop from the same ancestor, but they differ only because of the diet they are being fed. The genetically identical honeybee queen and workers get differentiated only because of the special diet called as ‘royal jelly’ fed to the larvae. In 1944 a famine, later named as Dutch Hunger Winter Famine, affected Netherlands. The environmental influence on the growth rate of that time (Barker hypothesis) was proposed which established that the men having lowest weights at early stages of development with poor nutrition had the highest death rates from cardiac diseases. If nutrition is not good enough during these stages, imperfections will be there in the growing person, even with the right genetic makeup, and this will make someone more prone to illness later in life. Studies were conducted in adult periconceptional individuals exposed to famine and found significant differences in IGF2 DMR methylation in adulthood. These findings demonstrate that even with the right genetic makeup, diet is one of the factors which can modify gene expressions and improper diet may be an etiology for many of adulthood diseases. The traditional concept of the role of early life nutrition in changing the epigenetic mechanisms causing disease condition has been substantiated. These data were the first to prove direct link between epigenetic modifications and diet that persist throughout life.

Future Perspectives

“You are what you eat” and also epigenetics prove that you are what your ancestors ate. Individual programmed traits which we inherit get altered by environmental factors including diet which may lead to diseases. Bacalini et al., has recently reviewed the inheritance pattern of epigenetics and the importance of environmental factors during different stages of development in utero. Recent discoveries have helped focus attention on the outstanding contributions of epigenetics in preventing diseases. As per Bacalini et al., age advances, epigenetic modifications also are positively related to diet, physical exercise, and lifestyle habits. However, we are still in the infancy of such studies, and that much has to be done in order to unravel the complex relationship between epigenetics, nutrition and ageing. Limited knowledge on the effects of nutrients or bioactive food components on histone modifications or chromatin remodeling complexes paves the way for further investigation on nutrients effect to improve human health.

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Conflict of interest

The author declares no conflict of interest for this manuscript.

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