Emerging Role of Nutri-Epigenetics in Inflammation and Cancer

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Disease risk and development are influenced by many factors (e.g., lifestyle, environment, nutrition) and genetics. Evidence shows that these factors interact with one another in ways that are unique to each person, making quantification of the risks of developing diseases such as cancer challenging. Cancer is a metabolically driven process with dynamic nutrient-responsive alterations within the human genome (Vander Heiden, Cantley, & Thompson, 2009). The epigenetic machinery acting on the human genome is heavily susceptible to alterations in metabolism and nutrition, particularly during periods of inflammation (Keating & El-Osta, 2015). This emerging knowledge leads to current interest in nutri-epigenetics or nutri-epigenomics. Epigenetics focuses on processes that regulate how and when certain genes are turned on and off, whereas epigenomics refers to analysis of global epigenetic changes across many genes, made possible by high-throughput methods. Therefore, nutri-epigenetics focuses on the process by which nutrition regulates how one specific gene is turned on or off, whereas nutri-epigenomics refers to the analysis of the interaction among multitudes of genes and nutrition, as well as the effects on global gene expression, which may vary among different tissues. This article aims to provide a brief overview of epigenetics and how it can be affected by metabolism and nutrition, discuss nutri-epigenetics and cancer, and consider the implications of nutri-epigenetics knowledge and evidence for nursing practice.

Background

The fields of both epigenetics and epigenomics have been empowered by publicly available databases (e.g., Human Epigenome Browser from Washington University School of Medicine in St. Louis; epigenome gateway.wustl.edu/info) that have focused on gathering epigenetics sequences for epigenome-wide association studies. These studies have demonstrated that epigenetic alterations related to food components and environmental (nutrition) factors, along with other genetic mutations, play a role in the development of inflammatory diseases, such as cancer (van Velden et al., 2015). The field of nutri-epigenomics provides new insights into diet–genome interactions, and these insights have permitted explorations of the use of metabolically based drugs, such as metformin (Glucomphage©), as part of cancer therapy (Kasznicki, Sliwinska, & Drzewoski, 2014) and the design of nutrition- and lifestyle-based cancer management strategies (Richman et al., 2013).

The term epigenetics was introduced to describe the interactions between genes and the environment that gave rise to phenotypes during development; it has been expanded to include environmentally responsive cellular processes that can be heritable and have long-term...
effects on gene expression without alterations in the DNA sequence (change in phenotype without a change in genotype) (Dupont, Armand, & Brenner, 2009). Through epigenetics, sets of genes may be expressed or silent, and this determines which proteins are transcribed in response to an environmental cue.

Two major epigenetic mechanisms that have been found to influence cellular metabolism and nutrient components are DNA methylation and histone modification. Each leads to either silencing or activation of gene expression (see Table 1). DNA methylation is the incorporation of a methyl group on the 5-carbon position of the cytosine ring of DNA by DNA methyltransferases (DNMTs), resulting in 5-methylcytosine. Methylation of cytosine often occurs adjacent to a guanine nucleotide within a gene; consequently, the site is called a CpG site (cytosine [C] lies next to guanine [G] in the DNA sequence, and a phosphodiester [p] bond joins the two nucleotides) (Kulis & Esteller, 2010). The insertion of methyl groups changes the structure of the DNA strands, modifying the condition in which a gene may be accessible to transcription machinery. DNA methylation functions to regulate gene expression, and this regulation has been found to be responsive to environmental cues, such as the presence of interleukins or other cytokines during inflammation (Suzuki, Toyota, Kondo, & Shinomura, 2009).

Dietary methyl group consumption, primarily involving methionine-containing protein products, provides an important source of methylation components for the enzyme DNMT through cellular conversion of methionine to S-adenosylmethionine (SAM) (Li & Tollefsbol, 2010). To balance the activity of DNMT, SAM can be converted in the cells to S-adenosylhomocysteine, which is a potent inhibitor of DNMT (Stempak, Sohn, Chiang, Shane, & Kim, 2005). In normal dietary metabolism, when choline and folate (vitamin B₁₂) are present, S-adenosylhomocysteine is recycled back to methionine, which can then be converted back to SAM. The DNA methylation process is sensitive to cellular metabolic reactions, called the methionine cycle, which ultimately relies on dietary methionine, folate, and choline. In contrast to DNA methylation, which modifies the bases within the DNA sequence, histone modifications are changes on the DNA-associated histone proteins. Histones are proteins that chemically bond with DNA molecules to form the chromatin structures that are condensed into chromosomes within the nucleus of the cell (Marín-Ramírez, Kann, Shoemaker, & Landsman, 2005). Modification of the histone proteins occurs primarily through methylation or acetylation of the amino acids arginine or lysine within the histone proteins. The acetylation of histone occurs by the enzymatic addition of an acetyl group from acetyl coenzyme A, a molecule that is generated by the metabolism of glucose and lipid-derived fatty acids (Verdin & Ott, 2015). The activity of histone acetylation enzymes (histone acetyltransferases) is balanced by histone deacetylases, which determine the transcriptional competency of a gene (Talbert & Henikoff, 2010). An imbalance in the equilibrium of histone acetylation/deacetylation is associated with tumorigenesis and cancer progression (Sobolewski, Sanduja, Blanco, Hu, & Dixon, 2015).

Although different epigenetic mechanisms follow distinctive pathways to regulate gene expression, they tend to be interrelated as part of a harmonized regulation of chromatin/chromosome structural dynamics. Altering structures that are known to affect epigenetic alterations may cause unusual gene activation or silencing of genes. Epigenetic changes have been related to cancer (breast, colon, lung, and leukemia) and to other diseases (asthma, autism, Fragile X syndrome, multiple sclerosis, obesity, rheumatoid arthritis, type 2 diabetes mellitus). Cellular processes, such as aging, host defense system, and imprinting, also play a role. These epigenetic processes are key determinants of gene expression, providing new dimensions of interest in epigenetic mechanisms in health outcomes.

### Nutri-Epigenetics, Inflammation, and Cancer

Nutrients, such as glucose, lipid-derived fatty acids, and methionine or other amino acids, can affect epigenetics by promoting or inhibiting enzymes involved in DNA methylation and histone modifications (see Figure 1). In normal cells, energy in the form of adenosine triphosphate is produced from glucose and fatty acids through a mitochondrial respiratory chain reaction that requires

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**TABLE 1. Epigenetic Mechanisms and Their Effects in Gene Expression**

<table>
<thead>
<tr>
<th>Epigenetic Mechanism</th>
<th>Effect</th>
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<tbody>
<tr>
<td>DNA methylation</td>
<td>Silencing</td>
</tr>
<tr>
<td>Histone acetylation or histone</td>
<td>Transcriptional activation</td>
</tr>
<tr>
<td>phosphorylation</td>
<td></td>
</tr>
<tr>
<td>Lysine methylation in histones</td>
<td>Silencing or transcriptional</td>
</tr>
<tr>
<td>RNA interference: siRNA, microRNA</td>
<td>activation</td>
</tr>
</tbody>
</table>

*Note. Based on information from Callinan & Feinberg, 2006; Keating & El-Osta, 2015.*
When oxygen is absent, lactate is produced from anaerobic metabolism of glucose. In cancer cells, whether oxygen is absent or present, lactate is continually produced from glucose metabolism, and this overrides the mitochondrial respiratory chain reaction of energy production. This unique preference of cancer cells to metabolize glucose independently of mitochondria, termed the Warburg effect of aerobic glycolysis, has a profound impact on epigenetics because it produces a shift in the availability of acetyl coenzyme A and other molecules needed for epigenetic enzymes (Gogvadze, Zhivotovsky, & Orrenius, 2010). The continual cellular uptake of glucose by cancer cells is indirectly associated with the expression of proinflammatory genes, such as interleukin-8 (Rofstad, Mathiesen, Kindem, & Galappathi, 2006). Evidence of epigenetic dysregulation in cancer is accumulating. The decrease in methylated cytosine results in global hypomethylation. Reduced cytosine methylation characteristically affects DNA sequences and CpG sites. Some cancers also show focal hypermethylation in CpG islands. Genes that are affected become permanently silenced. Methylation affects each patient with cancer differently. Although some patients have negligible variations, others display concordant hypermethylation of several genes, which is seen in colorectal cancer and other cancers (Stirzaker, Zotenko, & Clark, 2016). A buildup of genetic and epigenetic inaccuracies alters normal cells into aberrant tumor cells. In addition, DNA methylation patterns may produce atypical expression of cancer-related genes. Remembering that these DNA methylation arrangements are tissue- and cell-specific is important.

Review of the Literature

Chronic inflammation is a major risk factor for tumor formation, and studies have noted inflammatory signals as a new epigenetic mechanism that silences specific genes that cause inflammation-induced cellular changes (Abu-Remaileh et al., 2015). Nutrition is thought to be a factor involved in inflammation (Galland, 2010) and a modulator of risk toward some cancers (Day et al., 2013). The complexity of linkages between dietary components and epigenetics mechanisms,
such as DNA methylation, histone modification, and chromatin remodeling, including how these may affect the inflammation phenotype and the development of cancer, has been revealed (Keating & El-Osta, 2015). Diets rich in polyunsaturated fatty acids, plant-derived dietary phytochemicals, and macro- and micronutrients may produce mutagenic free radicals, oxidative stress, and inflammatory signaling, which have been implicated in epigenetic alterations (Alegria-Torres, Baccarelli, & Bollati, 2011). DNA modulation of endothelial cells with arachidonic acid promotes upregulation of tumorigenesis (Rodriguez-Blanco et al., 2014). However, diet high in polyunsaturated fatty acids may play a role through reduction of inflammation to reduce tumor formation (Liang et al., 2016).

Fruits and vegetables, which include polyphenolic compounds, such as resveratrol, tea catechins, and flavonoids, have shown chemoprotective properties (He et al., 2011); however, the nutritional consequences may be organ-specific. Studies of tea catechins demonstrate inhibitory activity against cancer development and cancer cell growth (Rahmani, Alshabrami, Allemailem, Aly, & Khan, 2015; Xiang et al., 2016). In addition, sulforaphane, mostly found in cruciferous vegetables (broccoli and broccoli sprouts), inhibits enzymes involved in cancer initiation (Gills et al., 2006), is chemoprotective, and, in animal studies, has been shown to reduce tumor growth (Myzak, Tong, Dashwood, Dashwood, & Ho, 2007). In the same study, healthy volunteers who consumed a single portion of broccoli were found to suppress histone deacetylase a few hours following consumption, with simultaneous activation of histone acetylation (Myzak et al., 2007). Likewise, another study in human tumor colon cell lines found that elevated doses of diallyl disulfide (found in garlic) increased histone acetylation (Robert, Mouillé, Mayeur, Michaud, & Blachier, 2001). Yi and Su (2013) discussed the molecular mechanisms by which diallyl disulfide aids in reduction of cancer proliferation. Studies focused on folate and vitamin B<sub>12</sub> intake, which is important for DNA methylation, have shown that diets low in folate are associated with risk for colon cancer (Kim, 2003), whereas diets high in folate are associated with a decrease in breast cancer (Chen et al., 2014). The effects of folate on epigenetic mechanisms and potential relationship to cancer remain to be seen.

Selenium is another dietary compound that epigenetically modulates DNA and histones by activating methylation-silenced genes (Speckmann & Grune, 2015). Biobehavioral interventions, including nutritional bioactives, are under investigation. Although findings in the area of nutri-epigenetics are exciting and compelling, additional clinical trials and epidemiologic research are needed.

Implications for Nursing

The opportunity exists to optimize patient outcomes through an interplay of host and genomic interactions. Nurses are at the junction to transform health care using genetics and genomics (Tully & Grady, 2015). Knowledge of epigenetic changes in disease processes sheds light into the complex mechanisms underlying tumorigenesis and has unlocked novel therapeutic options and targets for treating malignancies and their associated symptoms. Nurses possess the skills and are strategically positioned to lead research designed to better understand how the environment (i.e., nutrition) and epigenomics may interact mechanistically to add or reduce risks of certain cancers. The knowledge and awareness of epigenetics’ influence on inflammation and cancer and the effects of nutrients and metabolism on this process is critical for nurses who are at the forefront of patient care and education, particularly in the era of personalized medicine. That nutrition influences the epigenome is indisputable. This dynamic link between nutrition and genes provides great insight into how to target health promotion and disease prevention from a nutritional perspective. Nutrient and metabolic needs among individuals may vary greatly, depending on their specific disease process; therefore, the potential role of precision nutrition is key for modulating cancer risk and development.

Conclusion

Epigenetic modifications are key regulators of developmental processes involving differentiation and growth. Differences in epigenetic modifications may contribute to genetic diversity; however, aberrant epigenetic changes often lead to developmental abnormalities and diseases, such as cancer. Knowledge of epigenetic modifications in the development and progression of cancers will not merely allow nurses to recognize different high-risk profiles and to monitor response to drug therapy; it may also provide a stage from which they can design individual lifestyle interventions. Epigenetic changes are easier to modify than modifications that affect the genome. Therefore, nutri-epigenetics is an attractive biological target for modifiable behavioral nutritional interventions. Nurse-led clinical trials with diet modifications will enrich scientific understanding of how nutritional modulation concurrent with pharmacotherapies may enhance health outcomes in cancer.

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