Nutrigenomics and Public Health

Nutrigenomic research may uncover the keys to utilizing genetic information to create food products that improve public health through mitigation of chronic diseases.

Nutrigenomics is a relatively new and very fast-moving field of research which combines molecular biology, genetics, and nutrition. It focuses on the role of nutritional status or specific nutrients in the regulation of gene expression (Afman and Muller, 2006; Komdru et al., 2009; Mead, 2007). While one of its effects may be the development of personal diets based on an individual’s genetic code and focused on optimizing the expression of certain genes therein, other health-related outcomes show more promise at this juncture and have greater implications for public health promotion (Ordovas and Corella, 2004; Muller and Kersten, 2003). Of significant interest is the potential for greater knowledge of how diet may influence disease state at the gene or molecular levels. Many threats to public health, including cancer, diabetes, obesity and other chronic diseases, are influenced to a certain extent by genetic factors. The prevalence of cardiovascular disease, diabetes, and other related conditions in the United States is shown in Figure 1.

Traditionally, many recommendations for diet intake have been population based due to findings from epidemiology studies. For instance, recommendations to lower fat intake for a population are due to a link between fat intake and heart disease. However, not all individuals who consume a high-fat diet will develop heart disease. If we know who has a genetic susceptibility to a type of heart disease associated with elevated blood cholesterol, then perhaps diets can be tailored to those individuals. With the completion of the human genome project and continuous identification of the role of human genes, more information on who may be more likely to develop certain diseases and whether diet can prevent the expression of these diseases is an area of fundamental research.

The impact of nutrigenomics is not limited to public health. The food science and culinary industries may benefit from the information revealed by nutrigenomic research, and may also take advantage of opportunities to promote public health in a more active way. Understanding the impact of processing on food composition, added to new information regarding nutrients’ effects on genetics and disease processes, will afford new chances to improve public health through everyday manufacturing and food-processing practices. One such opportunity is the further development of functional foods with higher nutritional and sensory quality (Ferguson, 2009).

This article will focus on the potential public health impact of information gleaned from nutrigenomic research. We will explore how genes affect disease, and the ways in which nutrients affect genes. We will then discuss the potential impact of nutrient intake on disease state in light of current food consumption trends. Finally, we will outline how the food and culinary industries may take advantage of this information, and opportunities for exerting a positive influence on public health.

Genetic Regulation of Disease State

It has long been recognized that a family history of various chronic diseases increases an individual’s risk for developing such diseases. In general, the closer the genetic relationship between a patient and an affected family member, the greater the risk that the patient might develop the disease in question. The best explanation for this phenomenon is that many chronic diseases are regulated not only by lifestyle factors and environmental influences, but also by some genetic component.

Since the advent of sensitive molecular biology procedures,
Nutrigenomics and Public Health

many genes partially responsible for the development of chronic disease have been suggested, and some have been confirmed. Among these, perhaps the most well known are the oncogenes. Oncogenes are genes which code for proteins that control cell death, or apoptosis; or which control cell division and proliferation (Croce, 2008). Rapid cell proliferation or inhibition of apoptosis may cause a cell to become cancerous, and begin the growth of lesions which may, in turn, become tumors. Which mechanism causes the onset of cancer is dependent on the specific oncogene and tissue type involved (Land et al., 1983).

Another chronic disease with vast implications for public health is cardiovascular disease. Common biochemical markers for elevated cardiovascular risk include nonspecific indices of systemic inflammation, and their elevation is associated with cardiovascular risk, although they may be elevated due to more acute conditions such as infection. It has been demonstrated that the circulating levels of these markers may be influenced, at least in part, by various genetic factors (Pankow et al., 2001). Other genetic factors have also shown an effect in regulating the overall energy balance of an individual, contributing to the degree of one’s obesity or leanness (Barsh et al., 2000). These findings may influence how cardiovascular disease risk is diagnosed.

Hyperinsulinemia and diabetes are examples of chronic conditions plaguing the general public which have their origins, or partial origins, in genetics. Defective genetic regulation of several proteins which respond to the presence of insulin may be implicated in the etiology of type 2 diabetes. Ducuzseau et al. (2001) found decreased expression of key proteins and their mRNA in type 2 diabetic patients, and observed a lack of response to the presence of insulin by up-regulation of key response proteins. This lack of response is linked to defective expression of the genes for these proteins in response to insulin. Type 1 diabetes is reliant on genetic components to a greater degree than type 2 diabetes because the genetic regulation of type 1 diabetes is polygenic, or accomplished by several genes. Nutrient Regulation of Gene Expression

In some cases, intake or levels of specific nutrients have been shown to directly impact gene expression. Often, this occurs by means of transcription factors. Transcription factors are biochemical entities which bind to the DNA and either promote or inhibit transcription of genes. Several nutrients are known to bind to transcription factors and regulate gene expression in this manner (Muller and Kersten, 2003).

The regulation of gene transcription by fatty acids is perhaps the best understood of the nutrient gene-regulation pathways. A major pathway by which fatty acids modulate gene expression is the peroxisome proliferator-activated receptors (PPARs), which activate gene transcription when fatty acids bind to them (Bouwens et al., 2007). One such receptor has been shown to activate genes responsible for utilization of fatty acids for energy (Kresten et al., 1999). Circulating fatty acid concentrations are increased during fasting. These fasting-induced amplified fatty acid concentrations increase the expression of several genes involved in β-oxidation of fatty acids (Bouwens et al., 2007). One of the isoforms of PPAR is PPARα, which is a nuclear receptor that binds to the promoters of genes whose products favor fatty acid breakdown (Madrazo and Kelly, 2008). However, these receptors need activators to realize their full potential. Fatty acids can act as ligands to PPARα, such as oleic acid, to enhance activity (Michaul and Renier, 2009). Furthermore, the PPARs do not act by themselves but require another partner, the retinoic acid receptors with cis-retinoic acid as a ligand. Thus, fatty acid catabolism is somewhat dependent upon cis-retinoic acid or any other nutrient or drug that enhances the activity of the nuclear receptors (Figure 2).

Also, PPAR activity activation may lead to weight loss (Verreth et al., 2004). However, much has been done in developing drugs to act as ligands to both PPARα and

Figure 1. Prevalence of chronic diseases and related states in the United States (2006).

![Figure 1](image-url)
PPARγ; perhaps the same approach may be taken by the food industry (Robinson and Grieve, 2009).

Dietary carbohydrate intake has also been implicated in the regulation of gene expression. Glucose consumption is a crucial determinant of the rate of transcription in the liver of genes for several glycolytic enzymes and enzymes for fatty acid synthesis, as well as lipoproteins and insulin (Clarke and Abraham, 1992). However, this effect has also been observed in other tissues such as the pancreas (German et al., 1990).

Vitamins and minerals may also have transcriptional regulation properties. Retinoic acid, a form of vitamin A, forms part of transcription factors which may bind to the promoter region of genes involved in embryonic development and cell differentiation (DeLuca, 1991). Divalent minerals, such as iron, copper, and zinc, alter the transcription rates of genes for proteins such as metallothionein, a crucial transport and storage protein for these minerals (Bremener and Beattie, 1990). Probably the most notable nutrient gene interaction deals with vitamin D and the vitamin D nuclear receptor. Here, 1, 25 dihydroxycholecalciferol binds with the nuclear receptor, and in the nucleus this complex binds to promoters of genes that encode for many proteins. In essence, the role of vitamin D appears to favor pathways involved with cell differentiation and inhibits pathways favoring cell proliferation. Hence, this may have an impact on cancer as cell proliferation versus differentiation must be kept in balance (Samuel and Sitrin, 2008).

Folate metabolism may have variations among populations. When there is a mutation in one nucleotide base pair that alters the function of the protein that occurs in more than 1% of the population, this is termed a single nucleotide polymorphism or SNP. Folate synthesis in some populations may be limited due to SNP and thus more dietary folate may be required (Martínez de Villar-SNP and thus more dietary folate populations may be limited due to SNP. Folate synthesis in some a single nucleotide polymorphism 1% of the population, this is termed there is a mutation in one nucleotide variations among populations. When differentiation must be kept in bal- cancer as cell proliferation versus Hence, this may have an impact on differentiation and inhibits path- favor pathways involved with cell development and cell differen- (DeLuca, 1991). Divalent minerals, such as iron, copper, and zinc, alter the transcription rates of genes for proteins such as metallothionein, a crucial trans- port and storage protein for these minerals (Bremener and Beattie, 1990). Probably the most notable nutrient gene interaction deals with vitamin D and the vitamin D nuclear receptor. Here, 1, 25 dihydroxycholecalciferol binds with the nuclear receptor, and in the nucleus this complex binds to promoters of genes that encode for many proteins. In essence, the role of vitamin D appears to favor pathways involved with cell differentiation and inhibits pathways favoring cell proliferation. Hence, this may have an impact on cancer as cell proliferation versus differentiation must be kept in balance (Samuel and Sitrin, 2008).

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and consumers, and also more information should be made available to them. Although genetically sensitive foods are different from genetically modified foods, consumer skepticism cannot be ruled out if the information about nutrigenomics is not conveyed properly to consumers (Archibald, 2004). According to Archibald (2004), a genetic counselor or a physician could help an individual understand his/her genetic makeup, and then the nutritional professional could tailor diets accordingly (Figure 3).

Let us suppose for example that since oleic acid enhances PPARα activity and therefore fatty acid oxidation, should food and culinary entities choose food high in oleic acid (i.e., peanut or olive oils) in some of their meals or menus? Dysfunctional PPARα activity has been implicated in diabetes and thus it is not inconceivable that providing a diet to enhance PPARα activity may prove beneficial. Indeed, oleic acid and peanut oil high in oleic acid have been shown to reverse the inhibitory effect upon insulin production and can have a beneficial effect in type 2 diabetes (Vassiliou et al., 2009). Flavonoids have been shown to enhance PPARα expression and normalize blood triglycerides in high-fat-fed diabetic rats (Kaviarasan and Pugalendi, 2009). While trying to develop these targeted products, the developers (food and culinary industries) and the practitioners (mainly registered dietitians and nutritional professional) must consider the regulatory and legal issues associated with nutrigenomics. Since nutrigenomics is a relatively new field, work should be undertaken to develop proper ethical and legal guidelines to protect consumers (Reilly and DeBusk, 2008).

**Future Discoveries**

Current global trends in food consumption may have an impact on disease progressions observed worldwide. This impact may occur by means of nutrient regulation of genes, or by other unclear mechanisms which are yet to be discovered. There may be opportunities for the food industry to improve public health through modification of processing and production methods, as well as to capitalize on promotion of more positive trends. Nutrigenomic research may uncover the keys to utilizing genetic information to maximize public health and generally benefit the food and culinary industries.

The key to synthesizing this broad spectrum of information into a coherent picture may be the emerging field of nutrigenomics. Through nutrigenomic research, new nutritional regulation of gene expression will hopefully come to light. If specific nutrient regulation of genes closely related to disease onset or progression is identified, new arenas for disease preven-
tion and potential for treatment will come to the foreground of nutritional research and preventive medicine. Treatment and prevention plans based on nutrient consumption will necessarily rely heavily on the available food supply, which, in turn, relies upon the food and culinary industries.

Maximum utilization of the information brought forth by nutrigenomic research is crucial if these industries hope to promote public health. FT

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REFERENCES


