Environmental Epigenetics and Asthma
Current Concepts and Call for Studies

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Recent studies suggest that epigenetic regulation (heritable changes in gene expression that occur in the absence of alterations in DNA sequences) may in part mediate the complex gene-by-environment interactions that can lead to asthma. The variable natural history of asthma (i.e., incidence and remission of symptoms) may be a result of epigenetic changes, such as DNA methylation, covalent histone modifications, microRNA changes, and chromatin alterations, after early or later environmental exposures. Findings from multiple epidemiologic and experimental studies indicate that asthma risk may be modified by epigenetic regulation. One study suggested that the transmission of asthma risk may occur across multiple generations. Experimental studies provide substantial in vitro data indicating that DNA methylation of genes critical to T-helper cell differentiation may induce polarization toward or away from an allergic phenotype. Despite this initial progress, fundamental questions remain that need to be addressed by well-designed research studies. Data generated from controlled experiments using in vivo models and/or clinical specimens collected after environmental exposure monitoring are limited. Importantly, cohort-driven epigenetic research has the potential to address key questions, such as those concerning the influence of timing of exposure, dose of exposure, diet, and ethnicity on susceptibility to asthma development. There is immense promise that the study of environmental epigenetics will help us understand a theoretically preventable environmental disease.

Keywords: epigenetics; asthma; DNA methylation; gene–environment interactions

Asthma has long been recognized as a complex genetic disease mediated by exposures to a variety of environmental triggers. The greater concordance of asthma among monozygotic twins when compared with dizygotic twins demonstrates that both genetic and environmental influences are important to asthma pathogenesis (reviewed in Reference 1). By 2007, several dozen publications provided evidence of specific gene-by-environment interactions for this disease (Table 1), and many more can be anticipated. Yet, the clinical presentation of asthma is heterogeneous. Some individuals are affected as young children, others as adults. Furthermore, some develop asthma or wheezing illness as a result of occupational exposure, whereas others may become affected after exposure to urban air pollution (2, 3). Traditional models for gene-by-environment interactions proposed in research studies have not always matched the observations that physicians make in the clinic.

In 1992, the Barker hypothesis postulated that organs undergo developmental programming in utero that predetermines subsequent physiologic and metabolic adaptations during adult life. Early (including fetal) insults after nutritional and/or environmental exposures lead to a greater propensity to later disease (4–6). Early support for this hypothesis included evidence that severe fetal malnutrition was associated with an increased risk for multiple health problems throughout adulthood (5–7). Subsequent epidemiologic and experimental studies have found associations between the effects of a variety of prenatal environmental exposures, including allergens, antibiotics, and tobacco smoke, on disorders such as allergies, diabetes, neurodegenerative diseases, and cardiovascular diseases (5, 7, 8). This research inspired the fetal and early origins of adult disease hypothesis that proposes that prenatal or early postnatal environmental exposures influence developmental plasticity and result in altered programming. This programming is responsible for lasting functional changes of organs that lead to the development of a variety of complex diseases (reviewed in Reference 9).

Epigenetics is the study of heritable changes in gene expression that occur without directly altering the DNA sequence. One mechanism is DNA methylation, the covalent addition of a methyl group to a cytosine residue in a CpG site (i.e., where a cytosine lies next to guanine in the DNA sequence). CpG sites generally are clustered in high frequency near gene promoters and these regions are referred to as CpG islands. The methylation states of CpG islands in turn may affect gene activity and expression. Another epigenetic mechanism responsible for modulation of gene expression is post-translational modification of histones, including, but not limited to, acetylation, methylation, phosphorylation, and ubiquitylation (10, 11). Concurrent, multiple modifications of various histones create a complex pattern, often referred to as the histone code. These modifications permit transitions between chromatin states and alterations in transcriptional activity. DNA methylation usually works hand in hand with histone modifications to activate or silence genes by influencing chromatin structure and stability and therefore its accessibility by transcriptional factors (12). Both DNA methylation and histone modification are heritable from one cell generation to the next (13). As described by Callinan and Feinberg (12), potentially hundreds of methylated cytosines in multiple genes and dozens of post-translational chromatin modifications can arise. Epigenetic alterations are believed to occur predominantly prenatally and shortly after birth. However, recent evidence suggests they can occur during later periods (see below), influencing gene expression differentially throughout the lifespan. As such, epigenetic regulation provides an attractive mechanistic explanation, in addition to gene-by-environment interactions, for some of the molecular events linking early exposures with later disease.
Early seminal work performed by Cooney and colleagues, and by Waterland and Jirtle, demonstrated that dietary methyl supplementation during pregnancy with folic acid, vitamin B12, and other agents influenced the heritable phenotype of the agouti mice offspring (i.e., coat color distribution) via increased CpG methylation of the region upstream of the agouti gene (14, 15). Subsequent research demonstrated that the early postnatal diet induced epigenetic regulation of the imprinted gene Igf2, associated with several human cancers, in murine models (16, 17). Additional environmental exposures implicated in epigenetic regulation include xenobiotic chemicals, endocrine disruptors, maternal stress and nurturing, and low-dose radiation (reviewed in References 9 and 18). Importantly, epigenetic regulation, particularly DNA methylation, also appears to be inherited transgenerationally such that parental (or grandparental) exposures or phenotypes impact gene expression in the offspring (or offspring’s offspring) without changing the DNA sequence. This phenomenon has been demonstrated for the kinked tail (Axin(+/–) allele) and agouti-viable yellow (A(+) allele, and for associations between endocrine disrupters and male infertility, hormone-dependent cancer risk, and obesity (18).

Arguably, the variable natural history of asthma (i.e., incidence and remission of symptoms) may be a result of epigenetic regulation after early or later environmental exposures (Figures 1 and 2). A large body of research has implicated specific time periods when individuals seem to be more susceptible to the effects of environmental exposures and other asthma triggers. These include prenatal development, early childhood, and adolescence (19–21). During these time periods, epigenetic modifications may be more likely to develop. In other work, adult rats underwent reversal of stable epigenetic programming associated with glucocorticoid receptor expression and hypothalamic-pituitary–adrenal and behavioral responses to stress after exposure to histone deacetylase (HDAC) inhibitors or methionine (i.e., donor of methyl groups for DNA methylation) (22). This study raises the question of whether adulthood, a period when asthma often remits (23), may also be a period when critical epigenetic changes can be deprogrammed.

This perspective will describe evidence addressing the hypothesis that asthma risk may be influenced by environmentally induced epigenetic changes. Several epidemiologic studies, including one that suggests transmission of asthma risk after maternal exposure to environmental tobacco smoke (ETS) may continue across multiple generations, will be described. Experimental studies also will be reviewed that provide substantial in vitro data demonstrating that DNA methylation of genes critical to Th-helper (Th) differentiation may induce polarization toward or away from an allergic phenotype. Despite this early progress, fundamental questions remain that need to be addressed by well-designed research studies. Currently, data generated from controlled experiments using in vivo models and/or clinical specimens after environmental exposure monitoring are limited. Importantly, cohort-driven epigenetic research has the potential to address key questions, such as those concerning the influence of timing of exposure, dose of exposure, diet, and ethnicity, on susceptibility to asthma development.

### PRENATAL ENVIRONMENTAL EXPOSURES, GENE INTERACTIONS, AND ASTHMA RISK

A preponderance of evidence from multiple large prospective studies indicates that prenatal exposure to ETS is associated with impaired respiratory function, transient wheeze, asthma, and/or respiratory infections in infants, young children, and adolescents (24, 25). It is evident that prenatal exposure to ETS alters airway structure, because greater distances between alveolar attachments in intraparenchymal airways have been measured among exposed infants who died of sudden infant death compared with unexposed infants (26). This body of work provides the most convincing evidence that prenatal environmental exposures can influence the risk for subsequent asthma. In addition to ETS, low maternal intake of foods containing vitamin E and zinc (27), or use of antibiotics during pregnancy (28), may increase the risk for childhood asthma. In contrast, maternal intake of probiotics, and higher levels of fruits, vegetables, and oily fish during pregnancy, may decrease the risk (29, 30). A prospective study from the Columbia Center for Children’s Environmental Health showed that, at age 2 years, more difficulty breathing and probable asthma was reported among children jointly exposed to prenatal polycyclic aromatic hydrocarbons (PAHs) and postnatal ETS (31). Experimental animal models also have shown that intrauterine exposure to airborne pollutants may increase the risk for respiratory disease in offspring. Hamada and colleagues developed a mouse model that demonstrated that exposure during preg-
nancy to residual oil fly ash (extracted from a precipitator unit that removes particulate contaminates of an oil-fired power plant) led to a greater susceptibility to an asthmalike phenotype in the offspring mice, including airway hyperresponsiveness and inflammation, such as eosinophil infiltration and goblet cell hyperplasia (32). In other experimental studies, prenatal exposure to endotoxin and probiotic bacteria protected mice against an asthmalike phenotype (33, 34). Most recently, prenatal exposure of mice to either diesel exhaust particles or control “inert” titanium dioxide led to greater airway inflammation and hyperresponsiveness in the offspring (35).

PARENT-OF-ORIGIN EFFECT

Another set of studies suggested that inheritance of particular genetic polymorphisms from the mother is more likely to be associated with asthma in the child than inheritance from the father. This parent-of-origin effect points to a significant role of the maternal prenatal environment and gene interactions on later asthma risk in the offspring. As examples, polymorphisms in the β-chain of the high-affinity receptor for IgE (FcεRI-β) demonstrated stronger associations with positive allergy skin prick tests and greater allergen-specific IgE levels when they were inherited from the mother (36). In two separate cohorts, the HLA-G –964G allele was overexpressed in children with bronchial hyperresponsiveness if the mother was affected by bronchial hyperresponsiveness (37). The North Staffordshire cohort of subjects with mild asthma found that maternal, but not paternal, transmission of the glutathione-S-transferase P1 (GSTP1) Val105/Val105 and Ala114/Val114 genotypes was associated with greater lung function (i.e., FEV1/FVC) in children. The phenotype was not more frequent after maternal inheritance of GSTP1 Ile105/Ile105, Ile105/Val105, or Ala114/Ala114 (38). The North Staffordshire cohort of subjects with mild asthma found that maternal, but not paternal, IgE was positively associated with elevated IgE levels in cord blood and at age 6 months (40). The preferential association of children’s outcomes with maternal history of asthma or elevated IgE implies that the disease phenotype during pregnancy also may mediate the fetal response to environmental triggers and presumably the offspring’s subsequent asthma risk.

MULTIGENERATIONAL TRANSMISSION OF ASTHMA

Recent data also suggest that the risk of ETS exposure on asthma may be transmitted across two generations. The Children’s Health Study conducted a nested case-control study of 338 children with asthma who had been diagnosed by age 5 years versus 570 control subjects who were matched on in utero exposure to maternal smoking within grade, sex, and community of residence. Maternal and household smoking histories were obtained by telephone interviews. Grandmaternal smoking during the mother’s fetal period may have been associated with a greater asthma risk in the grandchildren, independent of maternal smoking exposure. The risk for asthma in grandchildren was heightened if both the grandmother and mother smoked during pregnancy (OR, 2.6; 95% CI, 1.6–4.5).
Although much of the data were collected retrospectively, the findings lend support to the possibility that environmental exposures may alter asthma risk across generations.

**EARLY POSTNATAL ENVIRONMENTAL EXPOSURES AND BOTH EARLY- AND LATE-ONSET ASTHMA RISK**

In addition to prenatal exposures, multiple cohort studies suggest that early postnatal exposures modify the risk for developing later childhood or adult-onset asthma. As an example, exposure to dust mite allergen during infancy may be an important determinant for later childhood asthma (42). Exposure to dog or cat allergen is associated with protection from later childhood wheeze in some (43), but not all (44), cohort studies. Exposure during infancy to indoor combustion-related pollutants has been associated with later childhood sensitization to dust mite and a reduction in FEV1 (45). Despite these associations between early postnatal environmental exposures and later disease, direct links to epigenetic mechanisms have not been established to date.

The links between early environmental exposure and adult-onset asthma are even more indirect, but several studies suggest possible relationships. Very low birth weight and prematurity, two disease phenotypes that have been associated with prenatal exposure to pollutants such as ETS and PAHs (46, 47), were associated with reduced FEV1.75-% (forced expiratory flow, mid-expiratory phase) FEV1/FVC, chronic cough, wheeze, and asthma during subsequent adolescence (48). Two other studies reported that respiratory infections during infancy were associated independently with a greater incidence of chronic obstructive lung disease (49, 50). Combined, both studies point to an enduring effect of respiratory infections during infancy on the risk for developing later childhood asthma. Therefore, exposures during either of these critical periods for lung development appear to affect the risk for developing asthma at later life stages, including during childhood and later adulthood.

**PROPOSED EPIGENETIC MECHANISMS**

The traditional view that interindividual risk for asthma, like other complex diseases, is determined solely by interactions between genetic polymorphisms and environmental exposures needs to be reconciled with new findings suggesting that epigenetic mechanisms also may contribute. These mechanisms include genomic imprinting, histone modification, altered DNA methylation of regulatory sequences in Th and other genes, and regulation by microRNA (miRNA), which may change asthma risk after conception via environmentally mediated epigenetic disruption of gene expression (18). In genomic imprinting, unequal expression of the maternal and paternal alleles occurs, presumably due to reversible modification of gene activity in association with the sex of the parent. Differential DNA methylation of promoter regions of reprogrammable genes may be an important mechanism in establishing the imprint (reviewed in Reference 51). For example, infections incurred during pregnancy have been hypothesized to transmit transgenerationally the imprints of infections and inflammation, reduce the offspring’s ability to withstand environmental pathogens, and in turn impact later morbidity and mortality (52). Direct mechanistic links to asthma pathogenesis still need to be determined.

Another proposed mechanism is induction of histone modifications, a process that is reversible and may be associated with chromatin remodeling and gene transcription. Oxidant-generating systems and proinflammatory mediators, some of which are implicated in asthma, may regulate histone acetylation (53). As an example, exposure to H2O2 (a reactive oxygen species) caused an increase in histone acetyltransferase (HAT) activity that promoted acetylation and induced chromatin remodeling (53). In other work, untreated subjects with asthma possessed greater levels of HAT and reduced levels of HDACs in bronchial biopsies, with levels that reversed after treatment with inhaled steroids (54). In addition, endotoxin exposure, associated with protection from developing atopy or asthma in several studies (55, 56), binds histones. It may disrupt chromatin remodeling as well and participate in gene silencing. The evidence is the finding in a septicemia cell model in which endotoxin exposure altered nuclear factor-κB binding and chromatin remodeling of the proinflammatory gene IL-1β promoter nucleosome (57).

Methylation of DNA and resulting changes in chromatin structure have been shown to initiate the process by which the Th cells lose their plasticity and differentiate productively toward the Th1 versus the proallergic Th2 pattern of cytokine gene expression. Demethylation of sites at the proximal promoter and conserved intrinsic regulatory element (CIRE) in the first intron of the proallergic IL-4 gene, and hypermethylation of sites in the counterregulatory IFN-γ promoter, all result in greater IL-4 production and Th2 differentiation (58–61). Methylation of a highly conserved DNaseI-hypersensitive region at the 3’ end of IL-4 is associated with Th1 differentiation (58). Interestingly, many CpG sites found in Th genes are highly conserved across species, including CpG53 and CpG190 of the IFN-γ promoter (61, 62). CpG53 resides in a proximal activator protein 1 (AP1)-binding site, and when methylated, can change transcription factor binding (63). In addition, methylation of CpG53 in the IFN-γ promoter was associated with Th2 polarization after inhibition of cAMP response element binding protein (CREB) and activating transcription factor 2 (ATF2)/c-Jun binding to the CpG-containing AP1 site (61). Therefore, mounting evidence from in vitro experimental studies suggests that the development of a polarized Th2 phenotype associated with atopy may result from altered DNA methylation at genes regulating Th differentiation. Moreover, recent work by our group demonstrated that inhaled diesel exposure and intranasal Aspergillus fumigatus induced hypermethylation at the CpG45, CpG53, and CpG205 sites of the IFN-γ promoter and hypomethylation at CpG408 of the IL-4 promoter in mice. Altered methylation of both gene promoters was correlated significantly with changes in IgE levels, suggesting that inhaled environmental exposures can induce methylation of Th genes and IgE regulation in vivo (64). In addition, ETS has been shown to induce gene hypermethylation in a gene associated with lung cancer (i.e., p16INK4a) tumor suppressor gene involved in cell cycle control (65). Whether these observations occur in association with clinical asthma still needs to be elucidated.

A rapidly emerging area of epigenetics research relates to the role played by miRNA in regulating asthma-related gene expression. Since the first miRNA, Lin 4, was discovered in 1993 (66), over 300 miRNAs have been identified in humans. Each miRNA may regulate up to 200 target genes by blocking the translation of the target protein (67). These molecules are believed to regulate up to one-third of all human genes by promoting the degradation of target messenger RNA. Aberrant expression of miRNA has been shown to contribute to the pathogenesis of many human diseases and may serve as valuable diagnostic or prognostic disease markers (68). However, studies relevant to asthma or asthma risk are still lacking, except for a recent report demonstrating that a single nucleotide polymorphism at the 3’ untranslated region of HLA-G, an asthma-susceptibility gene, affects the binding of three miRNAs to this gene (69, 70). This report raises the exciting possibility of a gene-by-epigenetics interaction. Perhaps future discoveries will implicate specific environmental exposures that might disrupt miRNA expression patterns in cells and tissues relevant to asthma pathogenesis.
WHAT ARE THE KEY RESEARCH QUESTIONS?

The first important question for epigenetic studies is to determine whether any in vivo environmental exposure can induce epigenetic alterations that underlie asthma. The clinical presentation of asthma is mediated by exposure to allergens, air pollutants, endotoxin, viruses, and ETS. Which of these are important? What are the effects of exposure dose or diet (i.e., methyl donors)?

Second, when are the time periods during fetal development and afterward when individuals are more susceptible to epigenetic modifications that influence the clinical course of asthma? How do they differ by environmental exposure, and how do they explain the induction or remission of asthma that is observed at different time points and ages? Can epigenetic changes be reversed?

Third, how do genetics and epigenetics interact? For example, how do polymorphisms in asthma-related genes, many of which have been identified (Table 1), interact with epigenetic mechanisms to confer heightened susceptibility to environmental influences?

Fourth, can epigenetic changes provide measurable biomarkers that may predict later asthma and greater need for immediate intervention? Can one identify an epigenome that is susceptible to, for example, air pollution-related asthma, and enact specific interventions to reduce asthma risk?

POTENTIAL APPROACHES TO FUTURE EPGENETIC STUDIES

Both in vivo experimental animal models and cohort-driven epigenetic research have great potential to answer many of the above questions. Mouse models may provide the experimental controls needed to examine the main effects of a variety of airborne exposures, including allergens and air pollutants, on selected candidate asthma genes. The modifying effects of diet enriched with methyl donors or HDAC inhibitors also can be tested. In addition, mice of different genetic background and those with unique transgenes could illuminate further these issues.

More exciting may be the potential to apply newly developing technologies to epidemiologic studies. Longitudinal cohort studies that feature repeated measures of specific environmental exposures, collection of biological specimens over time, and comprehensive clinical outcomes assessment have enormous potential to add a wealth of information on specific environmental triggers and critical time windows of susceptibility. In partnership with global, unbiased screening methods, such as methylation-sensitive restriction fingerprinting, restriction landmark genomic scanning, methylation of CpG island amplification–representational difference analysis, and methylation target microarrays to identify candidate regions, as well as validation assays to reassess genes of interest (71), epigenomes susceptible to ETS-induced asthma, for example, may be identifiable in the near future.

It is unlikely that any individual study will be able to account for all of the potentially confounding factors, limiting one’s ability to examine epigenetic pathways in observational studies. These include environment-by-environment interactions, traditional gene-by-environment interactions, effects of birth order, and other host factors such as obesity. Despite these limitations, there is great promise that the study of environmental epigenetics will help us understand a theoretically preventable environmental disease.

Conflict of Interest Statement: Neither author has a financial relationship with a commercial entity that has an interest in the subject of this manuscript.

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