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Accessibility
Stress and Childhood Asthma Risk: Overlapping Evidence from Animal Studies and Epidemiologic Research

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Rapidly expanding evidence increasingly strengthens the evidence linking psychological factors to asthma and allergy expression. Parallel studies in animals and humans demonstrating the influence of prenatal maternal stress and early caregiving experiences on the disrupted regulation of defensive biological systems [eg, sympathetic and adrenomedullary (SAM) system and the hypothalamic-pituitary-adrenocortical (HPA) axis] provide strong proof of concept for this line of research. The consequent altered neuroimmune responses may influence the expression of immune-mediated disorders such as asthma as well as enhance an individual’s susceptibility to other environmental factors that may also contribute to asthma risk.

Key words: asthma, childhood, interactions, prenatal, stress

Introduction

Efforts to understand the role of psychological stress in asthma expression and atopy are currently undergoing rapid expansion in the context of our increased understanding of both the neurobiology of stress and asthma pathophysiology, as well as trying to determine why asthma remains a leading cause of health disparities largely unexplained by known physical environmental factors. Notably, consensus statements by both the National Academy of Science and the National Institute of Environmental Health Sciences support the position that examining disparities in environmental health requires attention to both environmental hazards and social conditions. Although a number of theoretical models explaining health disparities have been proposed, a psychosocial stress model may offer the greatest promise.

With an estimated half of all cases diagnosed by age 3 years and two-thirds diagnosed by age 5 years, asthma is a developmental disease. This developmental framework presupposes that adverse early-life experiences, including prenatal exposures, may negatively influence neuroendocrine and immune developmental processes relevant to asthma risk. Although studies of mechanisms by which perinatal stress may increase the risk of childhood asthma are only beginning to emerge, proof of concept is provided by drawing from animal studies on the effects of early-life adversity on stress neurobiology and development and more recent human data that parallel the animal research. This overview provides a framework grounded in this theoretical rationale and may guide future studies that examine the mechanisms underlying the role of stress in asthma development in epidemiologic research.

Neurobiology of Stress

Psychological stressors have been associated with the activation of the sympathetic and adrenomedullary (SAM) system and the hypothalamic-pituitary-adrenocortical (HPA) axis (see Wright for an extensive review). Negative emotional responses disturb the regulation of the HPA axis and the SAM systems; that is, in the face of stress, physiologic systems may operate at higher or lower levels than during normal homeostasis. The disturbed balance of these systems is relevant to disease. Immune, metabolic, and neural defensive biologic responses important for the short-term response to stress may produce long-term damage if not checked and eventually terminated. The potential detrimental cost of such accommodation to stress has been conceptualized as allostatic load (ie, wear and tear from chronic under- or overactivity of the allostatic system). Hormones and neuropeptides
released into the circulation when individuals experience stress are thought to be involved in regulating both immunomediared and neurogenic inflammatory processes. Studies in animals and humans have shown that prenatal maternal stress and disturbances in early caregiving may have lasting effects on the stress pathways discussed above.12–14

Critical Developmental Periods

Prenatal Maternal Stress and Perinatal Physiologic Programming

Studies suggest that characteristics of the in utero environment, independent of genetic susceptibility, influence fetal development, including immune development. The concept that non genetic factors act early in life to permanently organize or imprint physiologic systems is known as perinatal programming.15 The HPA axis seems particularly susceptible to early-life programming.13,14,16 Both nonhuman primate and rodent models of prenatal stress and early adverse caregiving13,14,16 have helped us understand the consequences of similar experiences in humans.17,18 Maternal and fetal stress stimulates placental secretion of corticotropin-releasing hormone, which, in turn, is elevated in the neonatal circulation.19–22 This may stimulate the fetal HPA axis to secrete glucocorticoids, amplifying fetal glucocorticoid excess. Although these in utero responses may be adaptive in the short term, being geared toward coping with anticipated environmental challenges, ultimately they may exact a toll in contributing to increased risk of disease in later life.15

A well-known key characteristic of HPA axis functioning is the marked inter individual variability of responses to challenge,23 and the understanding of the relevance of this neuroendocrine system in human pathophysiology (including that relevant to asthma) requires the identification of the determinants of this variability. Genetic factors, other in utero and postnatal environmental factors, and the timing of exposures likely impact the differentiation of this response. The notion that the influence of maternal stress on such physiologic programming may vary based on an individual’s genetic background is supported by noted strain differences in hormonal and behavioural responses to stress in rats and mice.15 Data suggest that fetoplacental 11β-hydroxysteroid dehydrogenase, type 2 (11β-HSD2), may play an important role in modulating the programming effects of prenatal endogenous glucocorticoid exposure.19,24 The type 2 isof orm inactivates cortisol to cortisone, plays a role in the ontogeny of the fetal pituitary-adrenal axis, and protects the developing fetus from the adverse effects of circulating maternal glucocorticoids.25 Interestingly, 11β-HSD2 has been colocalized in human lung tissue and is expressed in human placenta.26,27 Interperson and interstrain (mouse) variability in the expression and efficiency of 11β-HSD2 has been demonstrated, suggesting genetic variability.28 Glucocorticoid receptors are also highly expressed in virtually all fetal tissues from midgestation or earlier.29 Variants of the glucocorticoid receptor gene may contribute to interindividual variability in HPA axis activity and glucocorticoid sensitivity in response to stress as well.30

The complexity of the measurement issues related to prenatal physiologic stress responses is beyond the scope of this overview. A number of recent reviews provide more detail.31,32 Others highlight particular methodologic challenges and detail strategies for studying these processes prenatally.33,34

Also central to consideration of the influence of antenatal maternal stress on the postnatal development of children are the putative effects of episodes of such stress on the fetus. These are beginning to be explored and documented.35 Prenatal cortisol dysregulation in depressed pregnant women has been linked to prematurity and low birth weight36 and postpartum depression.37 Gestational exposure to maternal stress has been shown to alter the development of humoral immunocompetence in offspring, as well as their hormonal and immunologic responses to postnatal stress.38–41 Evidence in rhesus monkeys suggests that stress experienced during pregnancy impacts the infant monkeys’ response to antigens at birth.42

In this light, there is also evidence that the asthma phenotype could be programmed before birth. Both genetic and environmental factors affecting maturation of the immune system during pregnancy and early childhood set the stage for the inflammatory processes and altered reactivity to stimuli that are characteristic of chronic asthma. Studies have shown a positive association between maternal use of antibiotics during pregnancy and the development of childhood asthma and a negative association between maternal use of probiotics during pregnancy and the development of childhood asthma.43 Others have considered the influence of maternal infections during gestation on asthma risk.44–48 Some speculate that stress triggers hormones in early life, which influence T helper (Th)2 cell predominance, perhaps through a direct influence of stress hormones on cytokine production,1,49 although this has not been studied directly to date.
In particular, alterations in stress-induced maternal cortisol levels may influence the fetal immune system development and lead to an increased risk of atopic disorders, as conceptualized in Figure 1.

In preliminary analyses in our laboratory, we examined the relationship between diurnal salivary cortisol expression and total immunoglobulin E (IgE) among 89 pregnant mothers enrolled in the Asthma Coalition on Community, Environment, and Social Stress (ACCESS) project, a prospective cohort designed to study the effects of early-life stress on urban childhood asthma risk. Salivary cortisol was collected five times per day over 3 consecutive days to assess basal awakening response, morning rise, diurnal slope, and area under the curve. Total IgE was dichotomized above or below the population mean (48.95 IU/mL). Repeated measures mixed models were run controlling for race, income, and weeks pregnant at the time of cortisol sampling. Higher levels of maternal total IgE were significantly associated with a flatter diurnal cortisol slope ($p = .05$). Examination of the cortisol curves showed that those with higher IgE demonstrated less of a decline during the evening. Thus, blunted HPA functioning in pregnant women was related to higher maternal total IgE expression. Other evidence suggests that elevated maternal IgE in utero may potentiate fetal sensitization to allergens and enhance atopic risk in infancy. Stress-induced altered activity of the maternal HPA axis may have immunomodulatory effects that influence expression of IgE during pregnancy, which, in turn, may have implications for fetal sensitization and childhood allergy and asthma risk. These findings warrant further study.

![Conceptual model linking prenatal maternal stress with childhood immunity. CNS = central nervous system; CRH = corticotropin-releasing hormone; NS = nervous system.](image-url)
Early Childhood Caregiving Experience

The early childhood environment and caregiving experience can also impact these processes. Studies in both rodents and primates have shown that environmental manipulations that increase maternal stress result in elevated cortisol levels and dysfunctional behaviors in offspring that are evident later in life. In parallel to the animal studies, evidence linking the social environment and stress to regulation of the HPA axis during early development in humans is also growing. Numerous retrospective studies in human HPA functioning suggest that increased reactivity of the HPA system is associated with early-life trauma and severe deprivation. Studies of infants and toddlers have linked maternal depression to dysregulation of the child’s HPA axis in both cross-sectional and longitudinal studies. Other studies of preschoolers and older children suggest that children’s cortisol levels are positively correlated with numerous social stresses and to broader family characteristics known to be associated with higher stress levels (e.g., low socio-economic status [SES]). Essex and colleagues examined the relationships of maternal stress beginning in infancy and concurrent stress on preschoolers’ (aged 4.5 years) HPA activity and later mental health outcomes. A cross-sectional analysis revealed that preschoolers exposed to high levels of concurrent maternal stress had elevated cortisol levels. Longitudinal analyses showed that concurrently stressed children with elevated cortisol also had a history of high maternal stress exposure in infancy. Importantly, children exposed only to high levels of concurrent or early stress had cortisol levels that did not significantly differ from those never exposed to stress. Also of note, further analysis of the specific components of stress indicated that maternal depression beginning in infancy was the most potent predictor of children’s cortisol.

In this context, it is notable that our laboratory has also linked early-life caregiver stress to repeated wheeze and dysregulation of immune function in a birth cohort predisposed to atopy. Specifically, caregiver stress assessed during infancy predicted antigen-specific T-cell proliferative responses in children at approximately 2 years of age. There is growing evidence that these cytokine patterns are already present in the first year of life and may have their roots in utero.

Need to Consider Environmental Interactions with Stress

Another notion that we have explored in an ongoing Boston cohort is whether there is increased asthma risk in lower-income urban environments that can be explained, in part, by a combination of increased contaminant exposures and greater susceptibility to their effects. It has long been noted that air pollution, for instance, may be higher near major roadways, power plants, and industrial sites, where property values are lower and lower-income populations reside. Increased life stress among subgroups living in lower-income neighborhoods has also been proposed as a primary pathway through which socioeconomic position (SEP) impacts health. These observations suggest that chronic social stressors, such as violence, may be more prevalent or severe in the same communities where pollution is elevated, resulting in both greater exposures and greater susceptibility.

Traffic-related air pollution is linked to asthma exacerbation and respiratory outcomes. Traffic-related air pollution is linked to asthma symptoms in cross-sectional population studies and in prospective studies linking caretaker stress to infant wheeze and IgE and immune mediator production. Some evidence indicates that violent events may trigger asthma episodes. Chronic stress may induce HPA axis and cortisol dysregulation, glucocorticoid resistance, SAM activation, catecholamine production, immune mediator function, inflammation, and cytokine production. Finally, stress and pollution impact common physiologic systems, facilitating synergistic effects; early-childhood environmental exposures and catecholamines affect Th1-Th2 balance. For example, studies show that psychological stress, diesel exhaust, cigarette smoke, and ozone affect oxidative stress, asthma, and chronic obstructive pulmonary disease.

Traffic-health relationships have been examined using many traffic indicators, with no consensus on which best capture variability in traffic-related pollution or health outcomes in different settings. Previous studies have successfully extrapolated traffic-related exposures from sampling homes to larger cohorts using predictive land-use regression (LUR) models. LUR shows strong predictive power for intraurban nitric oxide (NOx) variability using traffic and land use characteristics (i.e., population density, major sources).

My research group investigated the potential for exposure to violence (ETV), as a chronic stressor, to increase pollution susceptibility. We developed GIS-
based models to retrospectively estimate residential exposures to traffic-related air pollution for 413 children in a community-based pregnancy cohort recruited in East Boston, Massachusetts, between 1987 and 1993, using monthly NO₂ measurements for 13 sites collected over 18 years. Merging pollution estimates with questionnaire data on lifetime ETV (considered here as a chronic stressor) and prospectively collected repeated measures data on asthma onset in these urban children, we explored the hypothesis that stress may enhance the susceptibility to air pollution in childhood asthma etiology.

After correcting for potential confounders, including gender, SES, race/ethnicity, tobacco smoke exposure, and lower respiratory tract illnesses, we found an elevated risk of asthma with a 1 SD (4.3 ppb) increase in NO₂ exposure solely among children with above-median violence exposures (odds ratio [OR] = 1.63; 95% confidence interval [CI] = 1.14–2.33; p = .03). Among children always living in the same community, with lesser measurement error, this association was magnified (OR = 2.40; 95% CI = 1.48–3.88; p = .0009).

Summary

Evidence from both animal and human studies supports the notion that HPA functioning and other stress pathways may be altered by in utero stress and early caregiving experiences. Both nonhuman primate and rodent models of early adverse caregiving have helped us understand the consequences of similar experiences in humans. Disturbed regulation of stress systems (eg, HPA axis and the SAM system) related to chronic stress suggests that immune function, which is modulated by these systems, may also be disrupted in these individuals. This, in turn, may have implications for asthma development. Future studies that incorporate these strands of overlapping scholarship and strategies for studying stress reactivity during pregnancy, infancy, and early childhood are needed to continue to elucidate the mechanisms underlying the links between stress and asthma development. Specifically, these studies need to address how fetal exposure to stress may influence human immune and neuroendocrine development, whether such effects are independent of postnatal exposures, and how these pathways may, in turn, influence asthma development. The role of exposure timing and critical windows of development will need to be considered in these study designs. By so doing, we will be better able to translate this research into more effective intervention strategies and treatments.

Moreover, given the potential spatial covariance across exposures, and because stress and physical environmental factors (eg, pollution) may influence common physiologic pathways (ie, oxidative stress) and health outcomes (ie, respiratory disease), stronger methods are needed to disentangle their effects and investigate synergies. Similar hypotheses could be developed in relation to stress and tobacco exposure.

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