Gene-Environment Interaction and Covariation in Schizophrenia: The Role of Obstetric Complications

Vijay A. Mittal²,4, Lauren M. Ellman³, and Tyrone D. Cannon¹,2,4
²Department of Psychiatry and Biobehavioral Sciences, University of California Los Angeles, Los Angeles, CA; ³Department of Psychiatry, Columbia University, New York, NY; ⁴Department of Psychology, University of California Los Angeles, Los Angeles, CA

While genetic factors account for a significant proportion of liability to schizophrenia, a body of evidence attests to a significant environmental contribution. Understanding the mechanisms through which genetic and environmental factors coalesce in influencing schizophrenia is critical for elucidating the pathways underlying psychotic illness and for developing primary prevention strategies. Although obstetric complications (OCs) remain among the most well-documented environmental indicators of risk for schizophrenia, the pathogenic role they play in the etiology of schizophrenia continues to remain poorly understood. A question of major importance is do these factors result from a genetic diathesis to schizophrenia (as in gene-environment covariation), act additively or interactively with predisposing genes for the disorder in influencing disease risk, or independently cause disease onset? In this review, we evaluate 3 classes of OCs commonly related to schizophrenia including hypoxia-associated OCs, maternal infection during pregnancy, and maternal stress during pregnancy. In addition, we discuss several mechanisms by which OCs impact on genetically susceptible brain regions, increasing constitutional vulnerability to neuromaturational events and stressors later in life (ie, adolescence), which may in turn contribute to triggering psychosis.

Key words: schizophrenia/obstetric complications/gene-environment interaction/covariation/hypoxia/infection/stress/rGE/G x E

Introduction

Although genetic factors account for 80%-85% of liability to schizophrenia,¹,² findings that the disorder occurs in only 50% of monozygotic co-twins of an affected proband indicate that the environment plays a nontrivial role. Risk for schizophrenia is thus assumed to represent an integrated function of multiple predisposing genes as well as environmental risk factors, as instantiated in the multifactorial polygenic threshold model (MPTM) (see figure 1). A question of major importance, and the focus of this review, is how do these 2 classes of risk factors—genetic and nongenetic—contribute to susceptibility for disease onset?

Given the relative magnitude of the genetic contribution to schizophrenia, it is tempting to speculate that most or all environmental contributors have a gene-dependent relevance, as in genotype \( G \times E \). However, the \( G \times E \) model is only one of several possible mechanisms by which such factors could aggregate; other models including direct environmental influences⁴ and genotype-environment covariation (rGE) are also plausible.⁵ In fact, the answer to the question of how genetic and nongenetic factors aggregate in influencing risk for schizophrenia could vary according to the particular nongenetic and genetic factors in question. Highlighting the correct model for each contributing factor is critical given that each model makes different predictions concerning the etiological relevance of the nongenetic effects, therefore affecting ascertainment of populations at risk and prevention/intervention strategies.

Across environmental variables, obstetric complications (OCs) are among the most well-documented predictors of risk for schizophrenia.⁶⁻⁸ In this article, we evaluate 3 classes of OCs—hypoxia-associated obstetric events, maternal infection during pregnancy, and maternal stress during pregnancy—in relation to risk for schizophrenia, addressing the question how each OC coalesces with genetic influences.

Mechanisms of Genetic and Environmental Influence

Phenocopy Model

Some theorists have proposed that there may be a subgroup of schizophrenia patients who acquired the disorder...
from entirely nongenetic origins; ie, for some patients, an OC could cause schizophrenia without the participation of genetic influences. In the framework of the MPTM described in figure 1, the phenocopy model would imply that the weight of a particular environmental risk factor on disease risk is so substantial as to cause an individual exposed to it to cross the threshold on the liability continuum for overt expression of schizophrenia, even though the individual lacks genetic risk factors that account for a lion’s share of liability to the disorder in the population.

This model provides the clearest rationale for primary prevention because reducing the rate of exposure to the environmental influence in question would therefore result in a reduction in the incidence of schizophrenia in the population. The phenocopy model does not predict that all schizophrenic patients should have the exposure, just that all individuals with a history of the exposure would develop schizophrenia. One test of this model is to compare the rate of the exposure in the population with the rate of schizophrenia, such that increases in rates of the exposure should lead to measurable increases in disease risk. Using this strategy, we evaluate the phenocopy model within the framework of 3 prominent classes of OC.

Gene-Environment Covariation Model

The gene-environment covariation model posits that OCs are associated with the genes for the disorder. In this model, the roles of genes and OCs are confounded such that OCs may not actually play a role in the etiology of the disorder. In the framework of the MPTM, each covarying term (rGE) influences liability to the disorder, but because the G and E components are covariant with each other, it is unclear whether both causally influence liability or whether only one does, with the other representing an epiphenomenon. This model provides the least clear rationale for prevention because it is ambiguous whether reducing exposure to the environmental factor in question will result in reductions in liability to the disorder. The chief question when evaluating the covariation model is “does the exposure occur more frequently in groups at elevated genetic risk for schizophrenia than in groups without a known genetic risk?” A null result would weaken the gene-environment covariation model because gene carriers, such as patients with schizophrenia and unaffected relatives, would be no more likely to have the exposure than nongene carriers. For example, the gene-environment covariation model is weakened by the finding that mothers with a first-degree schizophrenia relative showed no significant increase in rates of OCs, suggesting that the genetic liability for schizophrenia, on its own, does not confer increased risk for OCs. Although mothers with a schizophrenia diagnosis show increased rates of OCs, these findings are difficult to interpret given that smoking during pregnancy was found to mediate the relationship between maternal schizophrenic status and decreased birth weight, suggesting that an observed increase in OCs among these women may be more related to health risk behaviors occurring during pregnancy than to genetic liability to schizophrenia. Given the aforementioned findings, examination of OCs among schizophrenic mothers is confounded by multiple intervening variables. Most studies of rGE use unaffected first-degree relatives of probands to index genetic liability, which is reasonable given that this group likely has some of the disease-producing genes. However, there is a cost in statistical power in detecting rGE to the extent that such individuals are not expected to have all the relevant predisposing genes.

Gene-Environment Interaction and Additive Influences Models

The gene-environment interaction model asserts that the environmental influence on risk for disorder depends on the presence of disease-promoting genes. According to this model, the occurrence of an OC in a genetically vulnerable individual would increase the likelihood of that individual developing the disorder. In the framework of the MPTM (figure 1), the effects of a particular G term and a particular E term are multiplicative in influencing liability to the disorder. It also is possible that genetic and obstetric risk factors for schizophrenia occur independently of each other but additively influence risk for disease expression. In both models, reducing the rate of exposure to the environmental factor in question will...
result in a reduction in liability to the disorder in the population, with the magnitude of this reduction depending on the weights of the direct effects of the particular E and G terms or the multiplicative effects of the particular G and E terms. Both the gene-environment interaction model and the additive influences model predict a relative increase in the rate of OCs among individuals who develop schizophrenia; therefore, the 2 models can be difficult to parse apart. Specifically, it is very difficult to segregate multiplicative vs additive gene-environment influences for a low-frequency disorder such as schizophrenia unless information on both family history and OC exposures are available on a very large population (tens of thousands) or unless one has directly measured the environmental and genetic influences in question. The latter approach has become more feasible with the recent availability of serological data from pregnancy and the identification of candidate genes involved in schizophrenia. Nevertheless, while the list is growing, at the present time only a handful of candidate genes have been identified, and there is clearly a plurality of susceptibility genes for schizophrenia, which further complicates detecting gene-environment interactions.

In the absence of direct measures of the genetic effects, it is still possible to test a gene-environment interaction vs additive influences model if a continuous measure of liability to the disorder is available. For example, certain structural brain abnormalities have been shown to covary with degree of genetic liability to schizophrenia. In this situation, one can perform a factorial analysis of the main effects of genetic and environmental risk and then test whether the multiplicative (interactive) term between the 2 classes of risk factors accounts for a significant proportion of the variation in the liability indicator (in this example, in the severity of structural brain abnormalities).

Multiple Models

In a recent review, Ellman and Cannon proposed that multiple models might be operating simultaneously in relation to a given genetic and/or environmental risk factor. Specifically, the authors point out that a genetic polymorphism associated with a magnified inflammatory response to infection has been linked to schizophrenia, such that in the presence of infection there is an exaggerated inflammatory response (gene-environment interaction), possibly leading to fetal neuronal injury, as discussed below. The authors also note that this polymorphism is associated with increased incidence of certain OCs, such as preterm delivery (gene-environment covariation); this relationship raises the possibility that a gene associated with schizophrenia could have multiple functions, some of which may contribute to the etiology of the disorder and some of which may be unrelated to etiology.

Obstetric Complications

Hypoxia-Associated OCs

A number of studies and reviews have concluded that fetal hypoxia likely is involved in a variety of OCs associated with risk for schizophrenia. Specifically, OCs associated with schizophrenia, such as emergency cesarean section, bleeding during pregnancy, and preeclampsia, have all been associated with fetal hypoxia. In order to test the strength of the relationship between hypoxia-associated OCs and schizophrenic onset in offspring, one series of investigations used a hypoxia-associated OCs scale including both indirect and direct indicators of hypoxia.

Indirect complications were selected based on validation with direct measures of hypoxia and from previous studies. OCs associated with hypoxia were related to schizophrenia, onset, especially among the cases who had an earlier age of onset.

Phenocopy Model of Hypoxia-Associated OCs. Are hypoxia-associated OCs capable of causing schizophrenia on their own, without the participation of genetic factors? Animal models (sheep and rats) indicate that prenatal exposure to hypoxia is associated with multiple brain abnormalities that have been found in schizophrenia patients, including decreased N-methyl-D-aspartate (NMDA) receptor binding; increased transcript expression of NR1 subunit in frontal and temporal regions, nucleus accumbens, and hippocampus (while NR2A subunit expression was found to be downregulated in hippocampal subregions); increased dopamine (DA) release including persistent alterations in function of mesolimbic and mesostriatal pathways as well as the way dopaminergic function is regulated by stress later in adulthood; neuronal death (cerebellum, hippocampus, and cerebral cortex); white matter damage; reduced growth of neural processes; disrupted development of mesotelencephalic pathways modulating the expression of sleep/wakefulness; and cognitive deficits in areas such as working memory and locomotion. Therefore, even in the absence of a genetic liability for schizophrenia, there are independent contributions of fetal hypoxia on neuronal and behavioral alterations similar to those found among schizophrenic patients.

Nevertheless, the prevailing evidence suggests that a phenocopy model is unlikely because the rate of hypoxia-associated OCs in the general population (10%-15%) is far higher than the rate of schizophrenia (approximately 1% of the population). That is, the vast majority of individuals who suffer hypoxia-associated OCs (97%) do not develop psychosis. It could be argued that a more limited phenocopy model might apply to the most severe cases of perinatal hypoxia. However, even among the most severely cyanotic infants who require
prolonged efforts to invoke breathing, the lifetime prevalence of schizophrenia does not exceed 6%. Although the phenocopy model is unlikely, we cannot fully reject the model because the currently available data are only relevant to OCs that are presumed to be related to hypoxia; to our knowledge there is yet to be a published study that directly measures fetal hypoxia using molecular indicators.

**Gene-Environment Covariation Model of Hypoxia-Associated OCs.** The preponderance of studies has found no increase in the frequency of hypoxia-associated OCs in groups at high vs low genetic risk for schizophrenia.8 One such study examined the incidence of hypoxia-associated OCs among individuals with schizophrenia, their nonschizophrenic siblings, and demographically matched nonpsychiatric controls drawn from the National Collaborative Perinatal Project cohort in the United States.25 The authors found a higher incidence of hypoxia-associated OCs in schizophrenia patients than in their nonsympothetic siblings and in nonpsychiatric controls. In contrast to what would be expected under a gene-environment covariation model, there was no difference in the incidence of such OCs between the nonpsychotic siblings and controls, suggesting that the incidence of OCs does not increase with genetic loading.25 If the covariation model were correct, siblings of schizophrenic patients, who share some of the disease-promoting genes, would have an increased risk of hypoxia-associated OCs. It is also possible that siblings have less of the disease-promoting genes, and therefore, the covariation is obscured due to decreased power to detect effects. Nevertheless, the prevailing evidence does not support the covariation model.

**Gene-Environment Interaction/Additive Influences Models of Hypoxia-Associated OCs.** In a prior review of the hypoxia-associated OC literature, Cannon8 found that regardless of whether risk is based on having an affected parent,5,31 sibling,32–35 or co-twin,36 studies using objective birth records have found that those high-risk subjects with a history of hypoxia-associated OCs are significantly more likely to become schizophrenic than those without such a history.

In magnetic resonance imaging studies of siblings from a Finnish birth cohort, hypoxia-associated OCs were significantly related to more severe brain abnormalities among patients with schizophrenia.37,38 Specifically, hypoxia-associated OCs were related to reduced gray matter and increased cerebrospinal fluid throughout the cortex in cases and their nonaffected siblings, as well as ventricular enlargement among the cases.37 Similarly, a stepwise decrease in hippocampal volumes according to genetic liability was found among cases, nonaffected siblings, and controls, with the most pronounced decreases among the cases exposed to hypoxia-associated OCs38 (see figure 2).
It is important that both studies are limited in that they utilized maternal and patient recall of OCs, and although both studies included hypoxia-associated OCs, it is unclear whether hypoxia is the primary mechanism of action underlying these epidemiological associations.

Prenatal Maternal Infection

In a landmark study, Mednick et al\(^43\) found that individuals whose mothers were exposed to an influenza epidemic in Finland during the second trimester of pregnancy exhibited substantially higher rates of schizophrenia. A multitude of studies have replicated this finding, and subsequent epidemiological studies have also linked fetal exposure to upper respiratory infections, genital tract infections, and rubella to risk for schizophrenia.\(^44\)–\(^50\) In addition, a new line of research utilizing serological confirmation of infection from archived maternal sera (not susceptible to retrospective biases) has indicated that in addition to influenza infection, \textit{Toxoplasma gondii} (\textit{T. gondii}), toxoplasmosis, and herpes simplex virus type 2 are linked to psychosis.\(^51\)–\(^53\)

With the exception of the parasitic infection by \textit{T. gondii}, most viral infections rarely cross the placenta, and evidence suggests that the teratogenic effects of fetal exposure to infection involves other mechanisms, such as the mother’s immune response to infection.\(^54\) This idea is supported by 2 studies that have found that fetal exposure to increases in proinflammatory cytokines during pregnancy resulted in increased incidence of psychosis in offspring.\(^51\)–\(^53\)

Phenocopy Model of Prenatal Infection. A phenocopy model predicts that prenatal exposure to a viral teratogen would result in a schizophrenia outcome independently of genetic contributions. Consequently, prenatal virus exposure would reliably lead to expression of schizophrenia. This model is supported by animal studies that have found neuronal and behavioral abnormalities similar to those found among schizophrenic patients after exposure to infection and/or proinflammatory cytokines.\(^55\)–\(^57\) For example, pregnant mice exposed to the human influenza virus yielded offspring displaying deficits in prepulse inhibition, as well as responses to acute administration of antipsychotic and psychomimetic drugs similar to adults with schizophrenia (ameliorations and exacerbations, respectively); further, at least some of these behavioral changes were attributable to the maternal immune response, given that a group of mice injected with sham viruses yielded offspring with acoustic startle response deficits.\(^56\) In a related rodent study, investigators examined the role of maternal exposure to human influenza virus on day 9 of pregnancy and found altered pyramidal and nonpyramidal cell density values as well as atrophy of pyramidal cells (despite normal cell proliferation) indicative of both acute and long-lasting effects characteristic to what would be observed in individuals with schizophrenia.\(^55\) Further, rat models of perinatal viral infection have revealed a pattern suggestive of disruption in developing neurocircuitry, yielding eventual death of dentate granule cells secondary to excitotoxicity.\(^58\)

In humans, researchers have also linked proinflammatory cytokines to neonatal white matter damage during prenatal maternal infection.\(^59\)

Despite these findings, only a small minority of mothers who are exposed to infection give birth to schizophrenic offspring.\(^43\),\(^46\),\(^60\) Even studies that have found increased rates of schizophrenia following flu epidemics, \textit{T. gondii}, toxoplasmosis, and herpes simplex virus type 2 note only an incremental increase.\(^60\)–\(^62\) The consistent findings showing that a large majority of individuals prenatally exposed to maternal infection do not develop schizophrenia weaken the phenocopy model.

Gene-Environment Covariation Model of Prenatal Infection. A gene-environment covariation model posits that prenatal infection is associated with the genes for schizophrenia but that the exposure itself may not exert any etiological role. If this model were correct, there would be an increased incidence of prenatal virus exposure in individuals carrying a genetic loading for schizophrenia, regardless of whether or not the individuals developed the disorder.

One strong area of support for gene-environment covariation comes from recent studies examining the incidence of genetic variation in genes associated with immune responses. Common proinflammatory cytokine polymorphisms (eg, interleukin IL-1 complex [IL-1\(\alpha\) (-889) allele 2, IL-1\(\beta\) (-511) allele 1, and IL-1RA allele 1] and tumor necrosis factor alpha [TNF-\(\alpha\)] have been found to be associated with schizophrenia outcome\(^63\)–\(^64\) and associated with particular OCs.\(^65\) This is a particularly powerful methodology; where we were once only able to index genetic risk using high-risk designs, the recent ability to examine specific genetic variations holds promise for elucidating mechanisms behind this complex phenomenon. In addition, these cytokines are associated with premature delivery,\(^66\) an OC that has been consistently reported to be elevated in the birth histories of schizophrenia patients.

As noted, preterm delivery has been found to be associated with aberrant developmental processes also implicated in the etiology of schizophrenia;\(^12\) these findings provide support for covariation but do not rule out the possibility of another mechanism of gene-environment influence. For example, a particular gene or gene cluster may contribute to both OCs and schizophrenia separately (as in pleitropy), but OCs may not have a causal effect on schizophrenia. For example, if a hypothetical gene is associated with low birthweight, and this same gene, independently of its effects on low birthweight, also results in an irregular DA metabolism that contributes significantly to psychotic symptoms, then it may appear.
as if low birthweight is associated with psychotic symptoms when in fact it is not. In this hypothetical example, it appears as though schizophrenia and low birthweight are related, whereas, they in fact represent different expressions of the same gene.

Some genes may contribute to schizophrenia not directly, but indirectly, by causing certain OCs that in turn affect brain structure and function in a way that increases risk for schizophrenia. For example, a hypothetical gene might increase rates of hypoxia, subsequently damaging the hippocampus, and this insult may participate with other risk factors resulting in the onset of psychotic symptoms. Such a gene would not be causally related to schizophrenia independently of its effects on OCs.

Each of these 2 models of rGE may apply in relation to different sets of OCs and schizophrenia susceptibility genes. OCs are heterogeneous in origin and mechanisms of effect and can therefore have a number of different causal relationships with schizophrenia, each through a different gene-environment mechanism of origin. Using the 2 examples noted above, the same person could carry a hypothetical gene resulting in low birthweight and independently of this effect, dysregulation of the dopamine system (which in turn increases risk for psychosis), and a second hypothetical gene that results in hypoxia-related hippocampal damage, increasing vulnerability for schizophrenia. Because schizophrenia is likely to be the result of numerous interacting genetic and environmental factors, this hypothetical person may in fact represent the rule rather than the exception.

Another likely possibility also serves as an illustration of the complexities faced in OC and biobehavioral genetics research. More specifically, OCs that directly effect etiological processes (like hypoxia in the second example above) might cause secondary OCs that are not directly related to etiological processes. For example, a gene related to hypoxia (which can indeed affect hippocampal development and subsequently play a significant vulnerability factor) may also indirectly lead to low birthweight, which may or may not have an independent etiological role in schizophrenia. Such interactions between OCs have been rarely studied to date. Consistent with the notion of multiple cooccurring mechanisms, in the following section we also discuss how these polymorphisms also support an interactive model.

Gene-Environment Interaction/Additive Influences Models of Prenatal Exposure to Virus. This model would predict that prenatal virus exposure in genetically high-risk individuals would increase the odds of a schizophrenia outcome. In addition to genetic polymorphisms leading to OCs, like preterm delivery, polymorphisms in both interleukin-1 complex [IL-1β (-889) allele 2, IL-1β (-511) allele 1, and IL-1RA allele 1] and TNF-α have been associated with increased inflammation in response to infection, as well as increased basal levels of proinflammatory cytokines. Thus, carriers of these polymorphisms are likely to be more vulnerable to the deleterious effects of prenatal infection, which could increase the likelihood of psychotic onset in adulthood, although there have been no published studies directly testing this potential gene-environment interaction.

One strong area of support comes from Brown et al., who found that the rubella-exposed individuals who eventually developed schizophrenia demonstrated a decline in IQ from childhood to adolescence as well as increased neuromotor and behavioral abnormalities when compared with rubella-exposed controls. Because rubella exposure uniquely affected the premorbid functioning of the schizophrenia group, it is likely that the genetic or environmental vulnerability associated with schizophrenia made the fetal brain more susceptible to the effects of rubella with observable cognitive effects appearing prior to the onset of the disorder.

Similarly, a recent study found that among ultrahigh-risk individuals, maternal recall of prenatal viral exposure was reported in the histories of 50% of those patients who converted to psychosis compared with only 20% of the high-risk individuals who did not convert to psychosis. Even though this study is limited by retrospective reports of prenatal virus exposure and the use of clinical high-risk status exclusively (without consideration of family history of psychosis in modeling genetic risk), it is suggestive of a link between prenatal viral exposure and increased risk of conversion among individuals at high risk for psychosis.

Our understanding of gene-environment interaction influence and prenatal exposure to infection remains limited. Studies looking at specific genetic polymorphisms that play a key role in the immune response combined with serological exposure data are needed to directly test gene-environment interactions.

Maternal Stress Exposure
Exposure to war, unwanted pregnancy, and death of a loved one or spouse are among the list of maternal stressors during pregnancy that have been associated with increased risk of schizophrenia among offspring. It is likely that prenatal stress triggers the release of maternal stress hormones, which have been found to disturb fetal neurodevelopment and subsequent functioning of the hypothalamic-pituitary-adrenal (HPA) axis, which in turn influences behavior and cognition. This mechanism is complex however, as effects appear to vary upon timing and duration of the stressor. Nevertheless, this effect has been demonstrated in numerous animal models, such that prenatal exposure to maternal stress and the administration of glucocorticoids (GCs) and other substances leads to changes in the structure/function of the HPA system in offspring that often persist through adulthood.
Findings from several lines of investigation (animal and human) indicate that persistently elevated GC levels can affect multiple neuronal regions that have been implicated in schizophrenia research. Animal studies have shown that, at heightened levels, GCs can inhibit neurogenesis in the dentate gyrus and contribute to neuronal death. In rodents, prenatal exposure to stress hormones has been associated with abnormalities in the adult brain including increases in dopamine D2-like receptors in the dorsal frontal cortex, medial prefrontal cortex, hippocampus, and nucleus accumbens. In addition, animal models have shown prenatal stress to be associated with decreased dopamine turnover and alteration of genes responsible for dopaminergic function in the midbrain.

In humans, research suggestive of possible alterations in learning beginning in utero point to the vulnerability and susceptibility of this delicate period. For example, salivary cortisol in late pregnancy has been associated with lower mental scores at 3 months. In addition, persistently heightened GC secretion also has implications for cognitive functions, in that the hippocampus plays a critical role in memory and research has shown that volumetric reductions of the hippocampus are associated with memory deficits. Prenatal inhibition of hippocampal neurogenesis has also been associated with persistent learning deficits. Further, fetal exposure to increases in stress hormones have been associated with OCs found in the history of schizophrenic patients, including shortened gestational length, fetal growth restriction, and decreased infant physical and neuromuscular maturation. These effects appear to depend on the gestational timing of the exposure, and emerging evidence suggests that the deleterious effects of fetal exposure to stress hormones may be moderated by gender. Taken together, the evidence suggest that prenatal stress exposure can result in lasting synaptic and neurotransmitter alterations, potentially altering personality and predisposing individuals to attention deficits as well as mental illness.

**Phenocopy Model of Maternal Stress.** Studies supporting a phenocopy model would yield results suggesting that exposure to prenatal stress in itself, without any genetic contribution, would reliably result in offspring with schizophrenia. As stated above, fetal exposure to stress hormones affects multiple areas of the brain implicated in schizophrenia. Nevertheless, while studies examining rates of schizophrenia in cohorts of individuals who experience a stressful event such as a war or a death of a spouse or close relative show an incremental increase in schizophrenia rates following stressful events, the findings are presented within the framework of small percentages. For example, van Os and Selten found a significant increase in the incidence of schizophrenia among individuals prenatally exposed to stress during a Nazi invasion, but the relative risk (1.15, 95% CI 1.03-1.28) indicates only an incremental increase in rates of schizophrenia. These results indicate that stress exposure is likely to play an etiological role, but other etiological events are likely to be necessary for a schizophrenia outcome.

**Gene-Environment Covariation Model of Maternal Stress.** Research supporting a gene-environment covariation model would yield results suggesting that maternal susceptibility to stress is associated with a genetic loading for schizophrenia but that the stress itself may not be a contributing etiological factor. The most appropriate basis for examining this question would be to select unaffected siblings or offspring of schizophrenics and then to compare severity of maternal stress during pregnancy in this group with that in a sample of unaffected individuals without affected relatives. However, the idiopathic nature of stress adds a level of complexity that renders such a design difficult to implement and interpret. Given the methodological limitations, testing a gene-environment covariation model in humans is difficult. Prospective studies, examining cortisol levels of schizophrenic and nonschizophrenic individuals, who have experienced a stressful event (naturalistically), and following their offspring stand to help elucidate our understanding of this complicated issue.

**Gene-Environment Interaction/Additive Influences Model of Maternal Stress.** A gene-environment interaction model predicts that individuals at genetic high risk for developing schizophrenia who also were exposed to acute or chronic prenatal stress would be at an increased risk of developing the disorder, when compared with genetically high-risk individuals not exposed to maternal stress or with healthy controls regardless of their exposure status. The best design study for investigating this model would consist of a prospective longitudinal study, assessing maternal cortisol levels throughout the pregnancy, following high-risk offspring (including multiple siblings) from the prenatal period to adulthood. If the high-risk children who developed schizophrenia were exposed to a greater degree of prenatal stress than their unaffected siblings, this would provide strong direct evidence for an interaction. However, this design is prohibitive in terms of cost and time and involves many uncontrollable naturalistic factors, and there have been no such studies to date.

**How Do OCs Influence the Pathophysiology of an Adult-Onset Disorder?**

Given the foregoing, it is imperative to identify the mechanisms through which OCs interact with the developing brain to increase the likelihood of schizophrenia in late adolescence and early adulthood. One theory is that OCs confer a latent constitutional vulnerability that become evident when interacting with later neuromaturational
processes. Within this framework, we review evidence suggesting that early hippocampal lesions (associated with OCs) can interact with normative adolescent increases in (1) synaptic pruning and (2) stress hormone levels and thereby contribute to the onset of psychosis.

In healthy development, the peak of synaptic density occurs at 2 years of age and is followed by a slow drop during childhood followed by a steep decline in adolescence. Early in development many synaptic connections are superfluous, and the increase in synaptic pruning during adolescence, particularly in the prefrontal brain regions, is thought to be related to efficiency and adult reasoning capabilities. Indeed, abilities to solve abstract and complex problems coincide with the pruning during this period. McGlashan and Hoffman proposed that an abnormally aggressive synaptic pruning process, which would lead to a reduction in synaptic connectivity, may be a strong contributing factor to psychosis. This idea is supported by postmortem studies, which have shown reduced neuropil, representing a loss of connections between neurons, without neural loss.

In continuing within the framework of this model, OCs disrupting a particular neural structure(s) (eg, hypoxia affecting the hippocampus) would, in effect, reduce the amount of synaptic pruning necessary to pass a threshold of synaptic loss, resulting in psychosis. This prediction is congruent with research of individuals with a history of OCs, who typically have an earlier age of onset and more pronounced neuroanatomical abnormalities.

An example through which OCs could fit within this framework also relates to the effects of early hippocampal damage. More specifically, a body of animal literature suggests that particular structures such as the hippocampus, and particular neural systems such as the HPA axis, are particularly susceptible to OCs, as discussed above. This work is of interest to our understanding of the etiology of psychotic disorders because the hippocampus regulates the HPA axis that governs the release of cortisol. Several lines of research suggest that prenatal insult can produce hippocampal damage in the fetus, and this can render the organism hypersensitive to stress postnatally. During adolescence, a time period also associated with the synaptic pruning noted above, there is also a well-documented normative increase in cortisol. Although the mechanisms are not yet clearly understood, it is clear that GC secretion augments dopamine activity in certain brain regions, especially the mesolimbic system. Because cortisol is known to augment dopamine activity, it is believed that the hippocampus and HPA axis act as a nonspecific moderating system potentiating expression of psychotic disorders as well as mediating the effects of stress on symptom expression. Within this framework, OCs resulting in hippocampal and/or HPA axis abnormality could result in a constitutional vulnerability or lesion, which may later interact with normative maturational processes leading to psychosis.

By acting through these distinct pathways, OCs may result in neural changes that place individuals at higher risk for schizophrenia. The brain may be "preprogrammed," in part via the impact of OCs on genetically susceptible brain regions such as the hippocampus, to be vulnerable to later life (ie, adolescent) stress and/or neurobiological influences (ie, circulating neurohormonal surges during pubertal maturation), which may subsequently trigger psychosis.

Conclusion and Future Directions

This present review provides support for the notion that different forms of OCs can influence liability to schizophrenia through disparate but potentially overlapping mechanisms of gene-environment coalescence. The findings have several broad-ranging implications. First, it may be possible to decrease the risk of schizophrenia in some genetically at-risk individuals with careful prenatal and perinatal monitoring. Further, a new generation of high-risk studies suggests that including OCs in models combining multiple risk indicators holds promise in enhancing our ability to identify those individuals who stand to most benefit from early intervention.

Second, some component of the genetic diathesis to schizophrenia would appear to render the fetal brain particularly susceptible to the neurotoxic consequences of OCs, and thus, search for candidate genes that mediate the brain’s vulnerability to early injury continues to be highly warranted. The emergence of candidate disease genes, as well as the advances in mapping out molecular pathways, will likely elucidate our understanding and treatment of psychosis. Although the list of candidates is long, prominent suspects might be located within the glutamatergic NMDA receptor system because overstimulation of such receptors represents an early event in the sequence leading from hypoxia to neuronal death. Moreover, as mentioned previously, genes that may be involved in mechanisms linking OCs to the etiology of schizophrenia have been identified, such as IL-1B and TNF-α polymorphisms. This research can lead to a new depth of epidemiological understanding and by linking the disease, environmental insult, and specific genetic polymorphisms help to highlight new promising gene clusters.

Although we have thus far focused on G × E interactions, another possibility is that OCs could alter the function and structure of genes leading to pathogenic consequences, such as schizophrenia. Specifically, in analyzing the phenomena of increased concordance rates of schizophrenia following population-sized famines such as the Dutch Hunger Winter and the Chinese famine of 1959–1961, McClellan et al. introduce the theory that a potential molecular basis for the increased risk
may be the occurrence of new mutations in genes critical for brain development and that famine-induced folate deficiency might be a mediator by impairing DNA repair. The authors point out that identifying disease-causing de novo mutations is particularly informative to the larger population of individuals with schizophrenia because a gene that harbors one disease-associated mutation is likely to harbor other illness-associated mutations. Further, because folate is necessary for normal DNA methylation, and folate regulates the expression of many genes involved in neurodevelopment, a deficiency could also lead to epigenetic alterations (stable changes in gene expression not dependent on mutations) that could predispose an individual to the later development of schizophrenia. This area of inquiry represents a significant future direction for OC research as well as epidemiological and behavioral genetics.

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