Associations Between Brain Structure and Connectivity in Infants and Exposure to Selective Serotonin Reuptake Inhibitors During Pregnancy

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IMPORTANCE Selective serotonin reuptake inhibitor (SSRI) use among pregnant women is increasing, yet the association between prenatal SSRI exposure and fetal neurodevelopment is poorly understood. Animal studies show that perinatal SSRI exposure alters limbic circuitry and produces anxiety and depressive-like behaviors after adolescence, but literature on prenatal SSRI exposure in humans is limited and mixed.

OBJECTIVE To examine associations between prenatal SSRI exposure and brain development using structural and diffusion magnetic resonance imaging (MRI).

DESIGN, SETTING, AND PARTICIPANTS A cohort study conducted at Columbia University Medical Center and New York State Psychiatric Institute included 98 infants: 16 with in utero SSRI exposure, 21 with in utero untreated maternal depression exposure, and 61 healthy controls. Data were collected between January 6, 2011, and October 25, 2016.

EXPOSURES Selective serotonin reuptake inhibitors and untreated maternal depression.

MAIN OUTCOMES AND MEASURES Gray matter volume estimates using structural MRI with voxel-based morphometry and white matter structural connectivity (connectome) using diffusion MRI with probabilistic tractography.

RESULTS The sample included 98 mother (31 [32%] white, 26 [27%] Hispanic/Latina, 26 [27%] black/African American, 15 [15%] other) and infant (46 [47%] boys, 52 [53%] girls) dyads. Mean (SD) age of the infants at the time of the scan was 3.43 (1.50) weeks. Voxel-based morphometry showed significant gray matter volume expansion in the right amygdala (Cohen $d = 0.65$; 95% CI, 0.06-1.23) and right insula (Cohen $d = 0.86$; 95% CI, 0.26-1.14) in SSRI-exposed infants compared with both healthy controls and infants exposed to untreated maternal depression ($P < .05$; whole-brain correction). In connectome-level analysis of white matter structural connectivity, the SSRI group showed a significant increase in connectivity between the right amygdala and the right insula with a large effect size (Cohen $d = 0.99$; 95% CI, 0.40-1.57) compared with healthy controls and untreated depression ($P < .05$; whole connectome correction).

CONCLUSIONS AND RELEVANCE Our findings suggest that prenatal SSRI exposure has an association with fetal brain development, particularly in brain regions critical to emotional processing. The study highlights the need for further research on the potential long-term behavioral and psychological outcomes of these neurodevelopmental changes.

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The prescription of selective serotonin reuptake inhibitor (SSRI) medications for pregnant women has accelerated over the past 30 years. To some extent, this rise may be attributable to increased awareness of the detrimental effects of untreated prenatal maternal depression (PMD) on women and children, along with early studies failing to document immediate effects of SSRI exposure in offspring (although later rodent studies document postpubertal alterations). However, little is known about the association between prenatal SSRI exposure and human fetal neurodevelopment.

Serotonin (5-hydroxytryptamine [5-HT]) plays a vital role in neurodevelopment. In the fetal brain, 5-HT signaling affects cell proliferation, differentiation, neuronal migration, network formation, and synaptogenesis. The 5-HT transporter is widely expressed in the fetal brain in both serotonergic and nonserotonergic neurons, thus providing a developmentally transient target for SSRIs. Atypical serotonergic signaling resulting from prenatal SSRI exposure may alter fetal brain development and subsequent functioning.

Animal studies support this idea. Perinatal SSRI exposure in rodent studies is associated with delayed motor development, reduced pain sensitivity, disrupted thalamocortical organization, reduced dorsal raphe neuronal firing, reduced arborization of 5-HT neurons, and altered limbic and cortical circuit functioning. Rodent studies also suggest behavioral consequences of early-life SSRI exposure, including increases in anxiety and depression-like behaviors in adulthood (eg, impaired stress response and grooming, decreased play), and suggest that early perturbations in 5-HT signaling may be associated with neurodevelopment, giving rise to atypical emotion-related behaviors later in life.

Literature on prenatal SSRI exposure in humans is limited and mixed. Studies have most consistently reported that prenatal SSRI exposure is associated with a shorter gestational period, lower birth weight, lower Apgar scores, and neonatal abstinence syndrome. Initial studies on longer-term neurodevelopmental consequences have yielded mixed findings; some studies suggest increased internalizing and externalizing behaviors during early childhood, whereas others fail to find such associations. However, consistent with animal studies, a recent national registry study (including >15,000 perinatally SSRI-exposed offspring) found increased rates of depression in early adolescence in youth with perinatal SSRI exposure.

Brain imaging provides a window into neurodevelopment, yet human infant and fetal imaging studies of prenatal SSRI exposure are scarce. A recent electroencephalography study found reduced interhemispheric connectivity and lower cross-frequency integration in SSRI-exposed infants, suggesting uncoupling of frontal circuitry. Two infant magnetic resonance imaging (MRI) studies documented changes in gray matter (GM) and white matter (WM) tissue properties in perinatally SSRI-exposed infants (eg, altered fractional anisotropy of the thalamostriatal GM and superior WM fascicule and increased mean diffusivity in several major fasciculi). Although these studies suggest an association between prenatal SSRI exposure and variation in fetal brain development, they are confounded by sample characteristics (eg, inclusion of very preterm-born infants) or the lack of an untreated PMD comparison group.

Based on prior animal studies, we hypothesized that perinatally SSRI-exposed infants would demonstrate altered GM morphology and WM connectivity within the corticolumbic circuit. To test this, we used deformation-based GM morphometry and diffusion probabilistic WM tractography. To more accurately assess the association between prenatal SSRI exposure and the infant brain, we considered the following methodological advances: 2 comparison groups (healthy controls [HCS] and infants exposed to untreated PMD), optimization of image analytics for the infant brain, and enhanced connectivity measures.

Methods

Participants
Data were collected between January 6, 2011, and October 26, 2016. Participants (pregnant women, aged 18-45 years) were recruited through obstetricians, midwives, and psychiatrists. A total of 204 mothers were recruited; 103 infants underwent an MRI scan (eMethods and eTable 1 in the Supplement). Group membership was determined after mothers completed a prenatal mood and medication assessment (between 19 and 39 weeks’ gestation). Mothers were assigned to the SSRI group if they self-reported receiving an SSRI at some point in their pregnancy. Sleeping, nonsedated infants underwent an MRI session when they were approximately 3.43 (SD 1.50) weeks of age. The New York State Psychiatric Institute Institutional Review Board approved all procedures and participants provided written informed consent. Participants received financial compensation for their participation.

PMD and Psychiatric Symptoms
Group membership (PMD vs HC) was determined during the prenatal assessment based on the mother’s depression scores, assessed via the Center for Epidemiological Studies Depression scale (scores ≥16 were considered indicative of clinically significant depression). Women completed the Schedule for Affective...
Disorders and Schizophrenia (SADS), a semistructured diagnostic interview.6 Owing to time limitations, 20 of the 98 mothers with usable infant MRI data did not complete the SADS (SSRI, 2; PMD, 5; and HC, 13). Postnatal depression was assessed via the CES-D, which was completed by mothers again at the time of their infant’s MRI session.

Infant 5-HT Transporter–Linked Polymorphic Region Genotype

To determine infant 5-HT transporter-linked polymorphic region (5-HTTLPR) genotype, saliva samples were collected and genotyped (eMethods in the Supplement). Genotype data were missing for 9 infants, who were excluded from 5-HTTLPR analyses.

MRI Acquisition and Analysis

Structural and diffusion MRI was acquired on a whole-body scanner (MR 750 3T; GE Healthcare) with an 8-channel head coil; eMethods in the Supplement provides details.

A T2-weighted structural MRI (single run) and diffusion-weighted images (2 runs) were obtained from 98 infants; 80 infants had usable T2-weighted scans and 91 had at least 1 usable diffusion-weighted imaging run (eTable 2 in the Supplement). The T2-weighted images underwent voxel-based morphometry. After preprocessing, diffusion-weighted imaging underwent probabilistic tractography and a recently developed filtering algorithm to curtail false-positive streamline estimates and improve the quantitative interpretability of streamline-based connectivity measures (eMethods in the Supplement provide details).

Statistical Analysis

For both GM morphometry and WM connectivity, linear regression with permutation testing (nonparametric) was used. To examine the effects of SSRI exposure beyond the effects of PMD exposure, primary contrast maps compared the SSRI group vs both PMD and HC infants. Three follow-up contrasts were then conducted, comparing (1) SSRI-exposed vs HC infants, (2) SSRI- vs PMD-exposed infants, and (3) PMD-exposed vs HC infants. The following covariates were included in the initial regression model: infant sex, age at scan, birth weight, and mother’s postnatal depression score, indexed via the CES-D. Significance of effects was determined using nonparametric permutation tests, which do not assume Gaussian distributions. To control for type I error in voxel-based morphometry, we used conditional Monte Carlo permutation testing (randomize program in Functional MRI of Brain Software Library [FSL] v5.0; 10,000 permutations) with the cluster-extent threshold option (a cluster-forming threshold of \( z = 3.1 \); whole-brain correction).

For WM connectivity data, we used both connection-level and whole-brain connectome-level analysis. Connection-level analysis used the same linear regression model as described above with exhaustive permutation testing in the ImPerm r package (https://cran.r-project.org/web/packages/ImPerm/index.html). Whole-brain connectome-level analysis was done with the Network-Based Statistics Toolbox (NBS, version 1.2).20 We used 2 methods to control for type I error: false discovery rate and network-based statistics. These methods are complementary because false discovery rate tests the null hypothesis at the individual connection level, whereas network-based statistics tests at the network level using familywise error. False discovery rate is more sensitive to focal effects, while network-based statistics is more sensitive to distributed network effects; 10,000 permutations were used to determine significance.

Results

Demographics

Magnetic resonance imaging scans were collected from 103 term infants, 98 of whom had usable MRI data. Five infants were excluded owing to apparent imaging artifacts resulting from excessive head motion. Specific subsamples for each imaging modality were structural MRI in 80 infants (SSRI, 14; PMD, 19; and HC, 47) and diffusion MRI in 92 infants (SSRI, 14; PMD, 20; and HC, 58). Groups did not differ significantly on infant gestational age at birth, sex, and birth weight (all \( P > .05 \); analysis of variance) (Table 1). No significant group differences were detected in the number of nondepressive psychiatric disorders documented with the SADS measure. Group differences were detected for infant age at MRI scan, maternal age, maternal race/ethnicity, and total family income (Table 1).

SSRI Exposure and GM Volume

Compared with infants not exposed to SSRIs (ie, PMD and HC), SSRI-exposed infants showed a significant GM volume expansion in the right amygdala and insula with medium to large effect sizes (SSRI vs PMD and HC; right amygdala, Cohen \( d = 0.65 \); 95% CI, 0.06–1.23; right insula, Cohen \( d = 0.86 \); 95% CI, 0.26–1.14) as well as in the right superior frontal gyrus and the left occipital gyrus at a whole-brain corrected \( P \) value < .05 (randomization permutation; adjusted for standard covariates) (Figure 1 and Table 2). An unadjusted model also showed a significant increase in volume in the right amygdala and right insula at whole-brain corrected \( P < .05 \).

Compared with PMD infants alone, SSRI-exposed infants showed a significant increase in volume in the right amygdala, right insula, right superior frontal gyrus, and right precentral gyrus at whole-brain corrected \( P < .05 \) (Figure 1 and Table 2). Furthermore, compared with HC infants alone, SSRI-exposed infants showed a significant increase in volume in the right amygdala, right insula, and right caudate. No regions showed a decrease in GM intensity in the SSRI-exposed group at whole-brain corrected \( P < .05 \). There were no significant differences in GM volumes between the PMD and HC groups.

SSRI Exposure and WM Connectivity

Linear regression of the structural connectome (connectivity was defined as streamline counts) revealed a significant increase in connectivity in the SSRI group relative to all infants not exposed to SSRIs (SSRI vs PMD and HC) in the following 4 connections: right amygdala-right insula, left anterior cingulate cortex-left thalamus, right precentral gyrus-right cuneus, and left insula-right precuneus at \( P < .05 \) (permutation testing).
testing; null hypothesis testing at the individual connection level using false discovery rate). An unadjusted model similarly showed a significant increase in connectivity between the right amygdala and right insula and between the left insula and right prefrontal cortex at the network level (net-1-ed education, maternal comorbid psychiatric disorders, and infant 5-HTTLPR genotype (eTable 3 and eResults in Supplement) in the first (n = 1), second (n = 2), and third (n = 13) trimesters. 

Table 1. Demographic Data

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>SSRI* (n = 16)</th>
<th>PMD (n = 21)</th>
<th>HC (n = 61)</th>
<th>Test Statistic (df)</th>
<th>P Value</th>
</tr>
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<tbody>
<tr>
<td>Age at scan, mean (SD), wk</td>
<td>4.29 (1.81)</td>
<td>3.03 (1.65)</td>
<td>3.30 (1.27)</td>
<td>F2,61 = 3.82</td>
<td>.02</td>
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<tr>
<td>Gestational age at birth, mean (SD), wk</td>
<td>38.71 (1.00)</td>
<td>39.32 (1.04)</td>
<td>39.43 (1.06)</td>
<td>F2,61 = 3.08</td>
<td>.05</td>
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<tr>
<td>Sex</td>
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<td></td>
<td></td>
<td>X2 = 0.84</td>
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<td>Male</td>
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<td>8</td>
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</tr>
<tr>
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<td>8</td>
<td>13</td>
<td>31</td>
<td></td>
<td>.65</td>
</tr>
<tr>
<td>Infant birth weight, mean (SD), g</td>
<td>3754.19 (1320.23)</td>
<td>4000.57 (755.81)</td>
<td>3888.78 (746.28)</td>
<td>F2,61 = 0.37</td>
<td>.69</td>
</tr>
<tr>
<td>Maternal age, mean (SD), y</td>
<td>33.12 (4.20)</td>
<td>27.55 (6.57)</td>
<td>31.04 (5.75)</td>
<td>F2,61 = 4.72</td>
<td>.01</td>
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<td>Maternal race/ethnicity</td>
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<td>X2 = 20.91</td>
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<td>Hispanic/Latino</td>
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<td>White</td>
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<td>8</td>
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<td>Other</td>
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<td>9</td>
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<td>Total family income, $</td>
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<td>0-25 000</td>
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<td>26 000-50 000</td>
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<tr>
<td>51 000-100 000</td>
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<td>2</td>
<td>20</td>
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<td>&gt;100 001</td>
<td>12</td>
<td>2</td>
<td>11</td>
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<td>Infant SERT genotype</td>
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<td>X2 = 2.66</td>
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<td>Long/long</td>
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<td>8</td>
<td>21</td>
<td></td>
<td></td>
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<tr>
<td>Short/long</td>
<td>4</td>
<td>7</td>
<td>28</td>
<td>.61</td>
<td></td>
</tr>
<tr>
<td>Short/short</td>
<td>4</td>
<td>2</td>
<td>9</td>
<td></td>
<td></td>
</tr>
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<td>Depressive disorder</td>
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<tr>
<td>Women with nongadular disorder</td>
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<td>3</td>
<td>5</td>
<td>X2 = 2.91</td>
<td>.23</td>
</tr>
<tr>
<td>Maternal CES-D, prenatal, mean (SD)</td>
<td>12.63 (12.12)</td>
<td>24.33 (7.22)</td>
<td>7.22 (3.99)</td>
<td>F2,61 = 51.80</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Maternal CES-D, postnatal, mean (SD)</td>
<td>10.84 (11.44)</td>
<td>15.8 (9.39)</td>
<td>6.98 (4.86)</td>
<td>F2,61 = 11.54</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

Abbreviations: CES-D, Center For Epidemiologic Studies Depression scale; HC, healthy control; MDD, major depressive disorder; PMD, prenatal maternal depression; SADS, Schedule for Affective Disorders and Schizophrenia; SERT, serotonin transporter; SSRI, selective serotonin reuptake inhibitor.

With diagnosis based on SADS score25: major depressive disorder (n = 3) and depressive disorder not otherwise specified (n = 3).

With diagnosis based on SADS score: major depressive disorder (n = 4) and depressive disorder not otherwise specified (n = 12).

With diagnosis based on SADS score: bulimia (n = 1), panic disorder (n = 2), generalized anxiety disorder (n = 1), and obsessive-compulsive disorder (n = 1).

With diagnosis based on SADS score: posttraumatic stress disorder (n = 1), panic disorder (n = 2), agoraphobia (n = 1), obsessive-compulsive disorder (n = 1), and simple phobia (n = 1).

With diagnosis based on SADS score: posttraumatic stress disorder (n = 1) and simple phobia (n = 4).

We confirmed these results in an additional tractography analysis with a larger total streamline count (ie, 100 million for initial tractography and 5 million for the streamline filtering analysis).

Potential Confounders

Effects of SSRIs on GM volume expansion and increased WM connectivity remained significant after adjusting for maternal age, maternal race/ethnicity, total household income, maternal education, maternal comorbid psychiatric disorders, and infant 5-HTTLPR genotype (eTable 3 and eResults in Supplement).

Prediction of Brain Changes Associated With Prenatal SSRI Exposure

We assessed the capability of the selected GM and WM measures (amygdala volume, insula volume, and amygdala-insula tract estimate) to predict prenatal SSRI exposure (Figure 3). A logistic regression model with both GM and WM (GM and WM model) showed the greatest area under the curve (AUC) of 0.83 (95% CI, 0.63-0.93; leave-1-out cross-validation), significantly greater than models with either WM (P = .008, Wilcoxon rank sum test; AUC, 0.74; 95% CI,
0.51-0.87) or GM (P = .003; AUC, 0.67; 95% CI, 0.45-0.84) alone. The GM and WM model with randomly permuted group labels showed an AUC of 0.41 (1000 iterations).

**Discussion**

To our knowledge, this is the first study to report increased volumes of the amygdala and insular cortex, as well as increased WM connection strength between these 2 regions, in prenatally SSRI-exposed infants. Our findings suggest a potential association between prenatal SSRI exposure, likely via aberrant serotonin signaling, and the development of the amygdala-insula circuit in the fetal brain.

Expression patterns of the 5-HT transporter in the developing brain may underlie the association of prenatal SSRI exposure with altered brain morphology. Exclusively during the prenatal period, 5-HT transporter is expressed in...
serotonergic neurons (eg, dorsal raphe) as well as nonserotonergic regions across the brain, such as corticollimbic and sensorimotor systems, as commonly seen in rodents, nonhuman primates, and humans. Expression in nonserotonergic cells is then repressed after birth. It is thus possible that the transient prenatal expression of the 5-HT transporter in nonserotonergic systems may render the fetal vs postnatal brain differentially sensitive to SSRI exposure.

Our finding of increased GM volume in the amygdala, anterior insula, and superior frontal gyrus in prenatally SSRI-exposed infants is in line with findings from animal studies. In 5-HT transporter knockout mice, studies have shown increased dendritic spine density in the amygdala and increased dendritic branching of pyramidal neurons of the infralimbic cortex. Studies also point to effects of pharmacologic blockade of the 5-HT transporter in rodents during the
Immediate postnatal period (ie, comparable to third-trimester gestation in humans) with subsequent increased anxiety and depression-like behaviors. However, these behavioral sequelae emerged only after the rodents entered puberty, much like what has been observed in a birth cohort study of children with gestational SSRI exposure.

Coordination of the amygdala and insula during threat awareness is essential to adaptive fear regulation. A meta-analysis of functional brain imaging studies indicates that fear conditioning is associated with task-related activations in both the amygdala and insula in healthy individuals, and analogous findings are reported in individuals with high trait anxiety. Similarly, the coordination of the amygdala and insula is essential not only for fear conditioning, but also for anticipatory anxiety, particularly under conditions of uncertainty.

Abnormalities in the amygdala-insula circuitry may be associated with anxiety and depression. Increased amygdala volumes are seen in both children and adults with anxiety disorders, heightened amygdala and insula task-related functional MRI activations are evident in adults with anxiety disorders during the presentation of fearful stimuli, and resting-state functional MRI studies show increased functional connectivity between the amygdala and insula in generalized anxiety and posttraumatic stress disorder. Similar functional connectivity findings are reported in children and adolescents across a range of anxiety disorders and symptoms. This abnormal amygdala-insula circuitry may be associated with increased vulnerability to anxiety and/or other mood disorders. Amygdala-insula hyperactivity to threats is reported in people who are at risk but not yet meeting criteria for anxiety disorders, suggesting that hyperactivity in this circuit may index increased susceptibility to anxiety disorders. Taken together, the structurally primed circuit in the infant brains could lead to maladaptive fear processing in their later life, such as generalization of conditioned fear or negative attention bias.

Prior infant neuroimaging studies report seemingly mixed findings: increased fractional anisotropy of the superior WM pathway and decreased fractional anisotropy and increased mean diffusivity of the WM pathways. However, the first study had no direct comparison of exposure to both SSRI and PMD vs exposure with PMD alone and, in the second study, the association between prematurity and neurodevelopment might confound the results.

The effects of SSRIs were present only in the right hemisphere. One study hints at asymmetric distributions of 5-HT receptors. In healthy adults, 5-HT1A receptor binding estimated by positron emission tomography is higher in the right frontal gyri relative to their left hemispheric counterparts (and higher in the left auditory cortex). It remains to be determined whether the asymmetric expression pattern of 5-HT transporter in adults is also present during the fetal period.

Limitations

Our findings should be interpreted in the context of a few limitations. First, because participants were not randomly assigned to the PMD or SSRI group, there could be unmeasured sample differences. It is possible that women who received an SSRI during pregnancy were more severely depressed than were those with PMD. Because our assessments of depression occurred when women were already receiving the SSRI, this hypothesis requires further investigation. Future human studies could include randomization and placebo control (eg, clinicaltrials.gov Identifier: NCT02185547). Second, the groups in our study differed in sociodemographics (maternal education, income, race/ethnicity, and birth weight). Although we statistically adjusted for these potential confounding variables, future research will be needed to conclusively disentangle SSRI and PMD exposure from these sociodemographic variables. Third, the behavioral and psychological correlates of our volumetric and connectivity findings need to be determined and longitudinal studies will need to examine whether developmental trajectories are affected. Last, the accuracy (eg, sensitivity, but perhaps not reliability) of the fiber orientation estimates in diffusion probabilistic tractography might be partially limited by a relatively small number of gradient directions (n = 11). However, our selection of scanning parameters was based on multiple factors, not solely on the number of gradient directions, such as spatial resolution (submillimeter in-plane resolution for the small neonatal brains), scan duration, and signal-to-noise ratio that decreases as the number of gradient directions increases.

Conclusions

Use of SSRIs during pregnancy has increased in recent decades, yet their association with fetal neurodevelopment continues to be a topic of considerable debate. Because...
untreated PMD poses risks to both the infant and mother, and the decision to initiate, continue, or suspend SSRI treatment remains a clinical dilemma. Preclinical studies of rodents indicate that dose, timing, and mechanism of action (5-HT augmenting or not) all contribute to outcomes in later life. Further study is required to better elucidate the effects of gestational SSRI exposure on fetal brain development and later life susceptibility to depressive, cognitive, and motor abnormalities. Such information may eventually allow more informed clinical decisions about how to best treat psychiatric disorders during pregnancy for the benefit of both mother and fetus.

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Study supervision: Cha, Gingrich, Posner.

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