Effects of Antenatal Maternal Depressive Symptoms and Socio-Economic Status on Neonatal Brain Development are Modulated by Genetic Risk

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Abstract

This study included 168 and 85 mother–infant dyads from Asian and United States of America cohorts to examine whether a genomic profile risk score for major depressive disorder (GPRSMDD) moderates the association between antenatal maternal depressive symptoms (or socio-economic status, SES) and fetal neurodevelopment, and to identify candidate biological processes underlying such association. Both cohorts showed a significant interaction between antenatal maternal depressive symptoms and infant GPRSMDD on the right amygdala volume. The Asian cohort also showed such interaction on the right hippocampal volume and shape, thickness of the orbitofrontal and ventromedial prefrontal cortex. Likewise, a significant interaction between SES and infant GPRSMDD was on the right amygdala and hippocampal volumes and shapes. After controlling for each other, the interaction effect of antenatal maternal depressive symptoms and GPRSMDD was mainly
shown on the right amygdala, while the interaction effect of SES and GPrSMDD was mainly shown on the right hippocampus. Bioinformatic analyses suggested neurotransmitter/neurotrophic signaling, SNAP RReceptor complex, and glutamate receptor activity as common biological processes underlying the influence of antenatal maternal depressive symptoms on fetal cortico-limbic development. These findings suggest gene–environment interdependence in the fetal development of brain regions implicated in cognitive–emotional function. Candidate biological mechanisms involve a range of brain region-specific signaling pathways that converge on common processes of synaptic development.

**Key words:** amygdala, cortical thickness, depression, morphological shape, polygenic risk score

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**Introduction**

Antenatal maternal depression, independent of postnatal maternal mood, influences fetal neurodevelopment, in particular the amygdala, where alterations associate with affective disorders (Rifkin-Graboi et al. 2013; Qiu et al. 2015a; Wen et al. 2017). Since depression is a familial disorder with a significant estimate of heritability (Sullivan et al. 2000), biological pathways linking maternal mood during pregnancy to offspring neurodevelopment are likely to vary as a function of genetic vulnerability for depression. Studies of candidate genes or of family history suggest the influence of “depressogenic” environmental conditions, such as childhood maltreatment and socio-economic status (SES), is moderated by genotype (Moffitt and Caspi 2006), reflecting gene–environment interdependence. However, candidate gene approaches bear weaknesses (Duncan and Keller 2011) and gene products operate in networks, such that alterations in neural systems that enhance vulnerability for psychopathology derive from genomic variants at multiple sites and may converge to influence common biological systems. Indeed, the genetic contribution to susceptibility for depression appears to reflect the cumulative influence of multiple genetic variants (Lubke et al. 2012). This idea led to the use of methods of genomic risk profiling to examine the influence of genetic burden as reflected by a set of “risk” alleles for specific psychiatric disorders (Ripke et al. 2013). The risk alleles and effect sizes of single-nucleotide polymorphisms (SNPs) are established from existing genome-wide association studies (GWAS) for depression using relevant “discovery” samples based on their P-values below a defined threshold. A genomic profile risk score (GPRS) is calculated for each individual in the target sample as the sum of the count of risk alleles weighted by the effect size in the discovery sample. The GPRS for major depressive disorder (GPrSMDD) predicts the risk for depression (P < 10^-6) (Ripke et al. 2013) and moderates the influence of childhood maltreatment on depressive symptoms (Peyrot et al. 2014). The target phenotype can also differ from that examined in the discovery sample allowing cross-phenotype analyses and studies of individual differences in endophenotypes. For example, a higher GPrSMDD associates with cortical thinning in the medial prefrontal cortex (Holmes et al. 2012), a region strongly implicated in the pathophysiology of depression (Kupfer et al. 2012).

Given the evidence on the relationship between antenatal maternal depression and the amygdala development during the fetal period (Rifkin-Graboi et al. 2013; Qiu et al. 2015a), we examined whether such relationship was moderated by infant GPrSMDD, controlling for maternal GPrSMDD, using a longitudinal Asian birth cohort sample. We then identified candidate genetic pathways and biological processes that might mediate the influence of environmental conditions associated with the genetic risk for MDD on the amygdala. We repeated the analysis on the amygdala using a US cohort of mother–infant dyads with measures of antenatal maternal depressive symptoms, neonatal neuroimaging data, and measures of GPrSMDD (see the description of the US cohort in the Supplementary Material). In this study, we computed GPrSMDD based on the SNPs selected from a Caucasian GWAS (Ripke et al. 2013). Given the fact on population differences in genotype frequency and risk genetic variants for MDD (Woo et al. 2004; Serretti et al. 2007; Porcelli et al. 2012; Sheikh et al. 2013; Ming et al. 2013, 2015), we noted that there may be a potential limitation of using a Caucasian sample as a discovery sample for studying the GPrSMDD in the Asian sample. Nevertheless, to the best of our knowledge, the discovery sample used in this study is from one of the most robust GWAS for MDD with the largest sample size (Ripke et al. 2013) at the current stage. Despite its limitation, it is informative in the sense that individual SNPs have opposing effects between Asian and Caucasian populations but they are still risk SNPs for MDD. For instance, Caucasians with 5 allele of the serotonin transporter polymorphism (5-HTTTLPR) show higher amygdala activation in response to emotional stimuli (e.g., Hariri et al. 2002, 2005; Murphy et al. 2013), whereas in Asians, 5 allele is associated with amygdala hyperactivation (Li et al. 2012). Hence, we expected that if Asian neonates with high GPrSMDD show the positive association between antenatal maternal depressive symptoms and the amygdala volume, whereas Caucasian neonates with high GPrSMDD show the opposite association or vice versa.

We further performed an exploratory analysis on whole brain morphology, including cortical thickness and hippocampal morphology, and examined whether GPrSMDD also modulates the association of antenatal maternal depression and other brain regions, particularly in the cortico-limbic system. Due to the compatibility in the magnetic resonance imaging (MRI) acquisition and the anatomical definitions of the brain structures, this study only performed the exploratory analysis using the Asian cohort. In particular, we employed large deformation diffeomorphic metric mapping (LDDMM) (Du et al. 2011) to perform detailed analysis of the morphological shape of cortical and subcortical brain regions. This approach allows for precise anatomical examination of these structures, well beyond that examined in brain volumetric studies, and reveals specific-regional shape changes associated with disease status (Qiu et al. 2009, 2010). Based on genetic contribution to MDD (Lubke et al. 2012) and the evidence that antenatal maternal depression influences fetal cortico-limbic development (Qiu et al. 2015a), we hypothesized that neonates with high GPrSMDD may show the negative relationship of antenatal maternal depression with fetal development of the cortico-limbic structures, including the hippocampus and prefrontal cortex. To further explore the moderating influence of the GPrSMDD, we examined the effects of SES, which influences the risk for a broad range of mental disorders, including MDD (e.g., Gilman et al. 2003), and cortico-limbic development (Farah et al. 2010; Noble et al. 2015).

This study, to the best of our knowledge, is the first kind to provide a strategy for the analysis of gene–environment inter-dependence and identify candidate biological processes that might mediate the influence of antenatal maternal depressive
symptoms on fetal brain development. Because of challenges in imaging neonates shortly after birth, the sample size of this study was modest for the study of gene–environment interaction and hence our findings are preliminary but may suggest biologically plausible candidate processes for transgenerational transmission of depression from mother to child.

Materials and Methods

Subjects

One-hundred and eighty-nine infants of mothers who participated in the prospective Growing Up in Singapore Towards healthy Outcomes (GUSTO) birth cohort study (Soh et al. 2012) were recruited for neuroimaging (Asian cohort). The Asian cohort recruited pregnant women attending an antenatal ultrasound scan clinic who were Singapore citizens or Permanent Residents of Chinese, Malay, or Indian ethnic background, and had no history of psychotropic medication.

We included the 168 neonates with genetic and neuroimaging data, gestational age at birth ≥35 weeks, birth weight >2500 g, and a 5-min Appearance, Pulse, Grime, Activity, and Respiration score ≥9 and whose mothers had genetic data and returned the questionnaires stated below. The Asian study was approved by the National Healthcare Group Domain Specific Review Board (NHG DSRB) and the Sing Health Centralized Institutional Review Board (CIRB). Written consent was obtained from mothers.

Scales of Maternal Depressive Symptoms and SES

In GUSTO, the Edinburgh Postnatal Depression Scale (EPDS) questionnaire was administered to mothers at 26 weeks of pregnancy to quantify depressive symptomatology. The EPDS (Cox et al. 1987) is a widely used 10-item self-report scale designed as a screening instrument for postnatal depression and valid for use in gestation. Each item of the EPDS is scored on a 4-point scale (0–3), and items 3 and 5–10 are reverse-scored. The reliability of the EPDS was 0.82 assessed using Cronbach’s analysis for our cohort. SES was measured using monthly household income (Table 1) and was collected at 26 weeks gestation via questionnaire.

MRI Acquisition and Analysis

In the Asian cohort, axial fast spin-echo T2-weighted MRI (repetition time = 3500 ms; echo time = 110 ms; field of view = 256 mm × 256 mm; matrix size = 256 × 256; 50 axial slices with 2.0 mm thickness) were acquired in infants 5–14 days of age using a 1.5-Tesla general electric scanner at the Department of Diagnostic and Interventional Imaging of the KK Women’s and Children’s Hospital. The detailed acquisition and image quality check procedure were previously reported (Qiu et al. 2013, 2015b). No sedation was used and precautions taken to reduce exposure to the MRI scanner noise. A neonatologist was present during scanning. A pulse oximeter was used to monitor heart rate and oxygen saturation throughout the entire scans. All brain scans were reviewed by a neuroradiologist (M.V.F). To determine image motion, all axial slices of the T2-weighted MRI data were visually inspected to ensure no cross-slice motion and checkerboard patterns.

Within individual subjects, when possible, 2 T2-weighted MRI acquisitions were first rigidly aligned and averaged to increase signal-to-noise ratio. In cases where only one scan was acquired, data from one scan were used in lieu of the average axial image. The skull of the averaged axial image was removed using Brain Extraction Tool (Smith 2002). A Markov random field model was used to automatically delineate amygdala, hippocampus, cortical gray matter, white matter, and cerebrospinal fluid (CSF) from the neonatal T2-weighted MRI data. The validation of this method in our sample was reported elsewhere (Qiu et al. 2013, 2015b; Rifkin-Graboi et al. 2013). Against the manual segmentation of the 20 brain images randomly selected in our data set, the segmentation accuracy rates of the amygdala, hippocampus, cortical gray matter, and white matter were respectively 0.75, 0.71, 0.79, and 0.86.

Based on the above tissue segmentation, a cortical surface was constructed at the boundary between gray and white matter using a graph-based topology correction algorithm. Cortical thickness was measured as the distance between cortical surfaces at the boundary between gray matter and CSF and at the boundary of gray matter and white matter. We employed a LDDMM algorithm to align individual cortical surfaces to the Asian atlas that was generated based on the cortical anatomy of the 20 subjects of this study and transferred the thickness of each individual subject to the atlas (Bai et al. 2012). The cortical thickness was smoothed using the Laplace–Beltrami basis functions on the cortical surface (Qiu et al. 2006). Amygdala and hippocampal morphological shapes were represented using surfaces contoured at the structure boundary. The LDDMM-surface mapping (Qiu and Miller 2008; Zhong and Qiu 2010; Tan and Qiu 2016) was then used to take surfaces as objects and provided transformation that deformed the Asian atlas to be similar to subjects (Bai et al. 2012). Shape variations of individual subjects relative to the Asian atlas were characterized by the Jacobian determinant of the transformation in the logarithmic scale, which represents the ratio of structural volume to the atlas volume. Positive values correspond to expansion, while negative values correspond to compression of the structure relative to the atlas at each anatomical location.

SNP Genotyping and GPRS Computation for MDD

In the Asian cohort, genomic DNA was extracted from frozen umbilical cord specimens for infants and from blood for mothers according to standard procedures. The samples were genotyped

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### Table 1 Demographics of the Asian cohort

|                          | Mean   | SD    |
|--------------------------|--------|-------|
| Gestational age (week)   | 38.69  | 1.13  |
| Birth weight (gram)      | 3117.85| 391.76|
| Birth weight adjusted by gestational age (gram) | 3118.14 | 379.01 |
| Postconceptional age at the MRI visit (week) | 40.08 | 1.21 |
| Antenatal maternal depression | 8.59 | 4.46 |

| Sex, %                      | N      | %     |
|----------------------------|--------|-------|
| Male                       | 168    | 53.6  |
| Ethnicity, %               |        |       |
| Chinese                    | 168    | 44.0  |
| Malay                      | 39.3   |       |
| Indian                     | 16.7   |       |
| Monthly household income (US$), % |  |       |
| ≤768                       | 168    | 3.0   |
| 769–1358                   | 13.7   |       |
| 1538–3076                  | 39.3   |       |
| 3077–4615                  | 23.2   |       |
| ≥4616                      | 13.7   |       |
| Unreported                 | 7.1    |       |

Note: SD, standard deviation.
on Illumina Omni express arrays, shown to perform well and with better coverage than alternatives in Asian populations (Jiang et al. 2013), and on Illumina Exome arrays, following the manufacturer’s instructions (Expression Analysis Inc). Further quality control on the genotyping calls were previously described (Qiu et al. 2015b). SNPs were verified for a genotyping rate ≥95% and no deviation from Hardy-Weinberg equilibrium (P < 0.001), and minor allele frequency ≥0.05, in PLINK (Purcell et al. 2007).

The GPRS<sub>MDD</sub> of infants and mothers was created using PLINK based on their SNPs and meta-analysis results from the Psychiatric Genomics Consortium (PGC; European ancestry, 9240 MDD cases, and 9519 controls) (Ripke et al. 2013). The GPRS<sub>MDD</sub> was a cumulative summary score computed as the sum across SNPs of the number of risk alleles (0, 1, and 2) weighted by odd ratio. Odd ratios were obtained from the meta-analysis on MDD from the Psychiatric Genomics Consortium website (http://www.med.unc.edu/pgc) (Ripke et al. 2013) and represented the allele-specific risk for MDD. We selected the SNPs with low linkage disequilibrium to each other (r<sup>2</sup> < 0.25 within 500 kb window, filtering for significance; PLINK-command -clump-p 1 -clump-p2 1 -clump-r2 0.25 -clump-kb 500) (Peyrot et al. 2014) and that survived at P-value thresholds (0.05, 0.1, and 0.2) to incorporate the proportion of variation in disease risk explained through their additive effects (Purcell et al. 2009; Holmes et al. 2012; Peyrot et al. 2014). The resulting numbers of selected SNPs for infant GPRS<sub>MDD</sub> were 2766, 5081, and 8935. The numbers of SNPs for maternal GPRS<sub>MDD</sub> were 2769, 5078, and 8902. This GPRS<sub>MDD</sub> was then standardized to a mean of zero and standard deviation of one to aid interpretation of results.

Assume that genetic risks for rheumatoid arthritis (RA) should not contribute to the risk for MDD and hence should not modulate the relationship between maternal depressive symptoms and neonatal morphology. In this study, we computed GPRS scores for RA (GPRS<sub>RA</sub>) as a negative control. We anticipated that GPRS<sub>RA</sub> would not modulate the relationship between GPRS<sub>MDD</sub> and neonatal brain morphology.

**Statistical Analysis**

Linear regressions were used to examine interaction effects of infant GPRS<sub>MDD</sub> with antenatal maternal depressive symptoms (or SES) on amygdala and hippocampus volumes and shapes, as well as cortical thickness of the whole brain in the Asian sample. The EPDS scores (or SES) and the GPRS<sub>MDD</sub> of the infant were included as main factors, which were entered into the first block of linear regressions along with covariates of gender, maternal ethnicity, birth weight adjusted by gestational age, postconceptual age on the MRI day. The interaction of the 2 main factors (EPDS score/SES and GPRS<sub>MDD</sub>) was then entered into the second block of the regression. For statistical analysis on shapes and thickness, the data were smoothed on the atlas surfaces using a heat kernel with 30-mm full-width-at-half-maximum (Chung et al. 2005). Statistical results were illustrated with P-value < 0.001 at the vertex level and with the cluster level of significance (P < 0.05) after correction for multiple comparisons based on random field theory (Chung et al. 2005, 2010). The correction of multiple comparisons was applied to one interaction model at a time, but not across genomic polygenic risk scores (GPRS<sub>MDD</sub> and GPRS<sub>RA</sub>) and different brain measures (volume, shape, and thickness).

If the interaction of the EPDS score (or SES) and GPRS<sub>MDD</sub> was significant, we performed post hoc analyses where the sample was then divided into 2 groups, one with a GPRS<sub>MDD</sub> greater than its mean value, 0 (high GPRS<sub>MDD</sub> group) and the other ≥0 (low GPRS<sub>MDD</sub> group). We examined associations of antenatal maternal depressive symptoms (or SES) and brain measures in each GPRS<sub>MDD</sub> group using regression with EPDS (or SES) as a main factor and covariates of gender, maternal ethnicity, adjusted birth weight, and postconceptual age on the MRI day. The brain measures used in this post hoc analysis were computed as average values of thickness, or shape deformation of the brain regions (colored regions shown in Fig. 2A and Fig. 3A) with the significant interaction of GPRS<sub>MDD</sub> with antenatal maternal depressive symptoms (or SES). The same analysis was repeated while additionally controlling for maternal GPRS<sub>MDD</sub>.

For the interaction of GPRS<sub>MDD</sub> and antenatal maternal depressive symptoms, the same analysis was also repeated while additionally controlling for SES and vice versa.

**Genetic Pathway Annotation Analysis**

Genetic pathway enrichment was used to explore the underlying biological processes associated with genes involved in the GPRS<sub>MDD</sub> computation. We first selected the top SNPs that moderated the relationship of antenatal maternal depressive symptoms with individual brain regions. These SNPs were mapped to genes using the batch query function in UCSC Genome Bioinformatics (http://genome.ucsc.edu). PANTHER (http://pantherdb.org) was then used to identify pathways that were statistically overrepresented against the reference genes involved in the same genetic pathways based on fold enrichment analysis.

We further explored biological functions of each genetic pathway using GeneMania (Mostafavi et al. 2008: http://Genemania.org). GeneMania was developed as a heuristic algorithm, derived from ridge regression, to integrate multiple functional association networks and predict biological functions from a single process-specific pathway. GeneMania was used to assess candidate biological functions associated with the genes in the pathways identified by PANTHER using multiple functional association data sets, including co-expression analysis, protein–protein interactions, co-localization, gene–gene interactions and protein domain similarity.

**Results**

**Demographics**

Table 1 and Supplementary Table S1 list the demographic information of the Asian and US cohorts. In the Asian cohort, antenatal maternal depressive symptoms did not vary as a function of infant gender (t<sub>166</sub> = 0.11, P = 0.92) and ethnicity (F<sub>2,165</sub> = 2.26, P = 0.11; Table 1). Antenatal maternal depressive symptoms did not vary as a function of infant or maternal GPRS<sub>MDD</sub> based on the 3 thresholds (all P > 0.2). But, SES was significantly correlated with antenatal maternal depressive symptoms (r = −0.273; P = 0.001) and significantly differed among the 3 maternal ethnicity (F<sub>2,155</sub> = 4.726, P = 0.01).

Among 168 neonates, 85 had 1 good T2-weighted MRI scan and 83 had 2 T2-weighted MRI scans. Between these 2 groups, we did not observe differences in birth weight (t<sub>166</sub> = −0.833, P = 0.406), postconceptual age on the MRI day (t<sub>166</sub> = 0.180, P = 0.857), gender (X<sup>2</sup> = 1.908, P = 0.216), maternal ethnicity (X<sup>2</sup> = 0.825, P = 0.662), and antenatal maternal depressive symptoms (t<sub>166</sub> = 1.140, P = 0.256).

**Interaction of GPRS<sub>MDD</sub> with Antenatal Maternal Depressive Symptoms on Neonatal Brain Morphology in the Asian cohort**

Regression analysis revealed a significant interaction effect between antenatal maternal depressive symptoms and infant GPRS<sub>MDD</sub> on the right, but not left amygdala and hippocampal...
volumes (Table 2) after adjusting for gender, maternal ethnicity, adjusted birth weight, and postconceptional age on the MRI day. The interaction effect was apparent at all 3 threshold levels of the GPRS_{MED}, with the largest interaction effect found using the infant GPRS_{MED} at \( P = 0.2 \). These findings remained significant when additionally controlling for the mother GPRS_{MED} (the Supplementary Material; Table 2) and also the finding for the right amygdala remained significant when additionally controlling for SES (the Supplementary Material). Subsequent post hoc analysis revealed a significant positive association between antenatal maternal depressive symptoms and right hippocampal volume \( (t_{8} = 2.04, P = 0.045; \text{Fig. 1B}) \) and the marginal, positive association between antenatal maternal depressive symptoms and right amygdala volume \( (t_{8} = 1.83, P = 0.076; \text{Fig. 1A}) \) in neonates with high GPRS_{MED} (\( > 0 \)). No associations were found between antenatal maternal depressive symptoms and right amygdala volume \( (t_{9} = -1.62, P = 0.109) \) and between antenatal maternal depressive symptoms and right hippocampal volume \( (t_{9} = -1.62, P = 0.109) \) in neonates with low GPRS_{MED} (\( < 0 \)).

Similarly, we found significant interaction effects between infant GPRS_{MED} and antenatal maternal depressive symptoms on the right amygdala shape (corrected cluster \( P = 7.86 \times 10^{-5}; \text{Fig. 2A} \)) and anterior right hippocampal shape (corrected cluster \( P = 0.016; \text{Fig. 2A} \)), with the largest interaction effect at GPRS_{MED} of \( P = 0.2 \) among the 3 thresholds (0.05, 0.1, and 0.2). This interaction effect was apparent for GPRS_{MED} of all 3 threshold levels. These findings remained significant when controlling for maternal GPRS_{MED} and the same finding on the right amygdala shape remained significant when controlling for SES (the Supplementary Material). The post hoc analysis revealed a significant positive association between antenatal maternal depressive symptoms and right anterior hippocampal shape \( (t_{9} = 1.84, P = 0.070; \text{Fig. 2B}) \) in neonates with high infant GPRS_{MED} (\( > 0 \)). No associations were found between antenatal maternal depressive symptoms and right amygdala shape \( (t_{9} = -1.77, P = 0.08) \) and between antenatal maternal depressive symptoms and right anterior hippocampal shape \( (t_{9} = -1.63, P = 0.108) \) in neonates with low infant GPRS_{MED} (\( < 0 \)).

We found a significant interaction effect between antenatal maternal depressive symptoms and infant GPRS_{MED} on thickness of left orbitofrontal cortex (OFC) and ventromedial prefrontal cortex (vmPFC) (corrected cluster \( P = 0.031; \text{Fig. 3A} \)), with the largest interaction effect using the GPRS_{MED} at \( P = 0.1 \) among the 3 thresholds (0.05, 0.1, and 0.2); the interaction was also apparent using the GPRS_{MED} at \( P = 0.2 \). These findings remained significant when additionally controlling for the maternal GPRS_{MED} (the Supplementary Material). These findings became a trend of significance after additionally controlling for SES (the Supplementary Material). The post hoc analysis further revealed the significant positive association between antenatal maternal depressive symptoms and cortical thickness of left OFC and vmPFC \( (t_{9} = 2.63, P = 0.010) \) in neonates with high infant GPRS_{MED} (\( > 0 \)), suggesting that a greater maternal depressive score was associated with thicker cortex in left OFC and vmPFC (see Fig. 3B). No association was found between antenatal maternal depressive symptoms and cortical thickness of left OFC and vmPFC (see Table 3). Moreover, the GPRS_{RA} did not affect the association between cortical thickness in the vmPFC and OFC and antenatal maternal depressive symptoms (cluster \( P \)-value > 0.05).

**Interaction of GPRS_{MED} with SES on Neonatal Brain Morphology in the Asian Cohort**

Likewise, a significant interaction effect between SES and infant GPRS_{MED} on the right, but not left amygdala and hippocampal

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**Table 2 Statistical results for amygdala and hippocampal volumes in the Asian sample**

| Structure                  | Mean (SD) (mm³) | Interaction effects |
|----------------------------|-----------------|--------------------|
| Left amygdala              | 211.41 (34.48)  | \( F_{50} = 2.96, P = 0.087 \) |
| Right amygdala             | 187.09 (33.77)  | \( F_{50} = 6.86^{*}, P = 0.009 \) |
| Left hippocampus           | 779.40 (108.18) | \( F_{50} = 0.002, P = 0.964 \) |
| Right hippocampus          | 781.76 (109.11) | \( F_{50} = 4.15^{*}, P = 0.043 \) |

Note: \( F \)-values indicate the interaction effects of infant GPRS_{MED} with antenatal maternal depressive symptoms on the structural volumes.

\(^{*}\)\( P \)-value < 0.05.

\(*\)The result remains significant when maternal GPRS_{MED} was entered in regression analysis as an additional covariate.

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**Figure 1. Interaction of the infant’s GPRS_{MED} with antenatal maternal depressive symptoms on the amygdala and hippocampal volumes.** Scatter plots in panels (A) and (B) illustrate the relationship of antenatal maternal depressive symptoms with right amygdala volume and right hippocampal volume in neonates with high GPRS for major depressive disorder (GPRS_{MED}, blue asterisks) and those with low GPRS_{MED} (magenta circles).
volumes and shapes (Supplementary Table S2) after adjusting for gender, maternal ethnicity, adjusted birth weight, and postconceptional age on the MRI day. Such interaction effects remained significant for the right hippocampus but not the right amygdala when additionally controlling for antenatal maternal depressive symptoms. No interaction effect between SES and infant GPRS was observed on cortical thickness.

Brain Region-Specific Gene Networks and Biological Processes

Among the SNPs involved in the GPRS, the top 1551 showed the most significant interaction with antenatal maternal depressive symptoms ($P < 0.2$) on the right amygdala, 1170 SNPs on the right hippocampus, and 1236 SNPs on the left OFC and vmPFC thickness, which were mapped to 629, 508, and 503 genes, respectively. PANTHER analysis revealed biological pathways for the 3 structures. These pathways were statistically over-representative in relation to those in the PANTHER pathway database (Table 4). For the right amygdala, 78 genes could not be found in the PANTHER database. The remaining 551 genes formed one statistically over-represented pathway ($P = 0.003$) labeled by PANTHER as “metabotropic glutamate receptor group III”. For the left OFC and vmPFC, 61 genes could not be found in the PANTHER database. The remaining 442 genes formed 5 statistically over-represented pathways, including “heterotrimeric G-protein signaling pathway-Gq alpha and Go alpha” ($P = 0.004$), “endothelin signaling” ($P = 0.007$), “histamine H1 receptor” ($P = 0.005$), 5-HT2 type receptor ($P = 0.013$), and “oxtocin receptor” ($P = 0.044$). For the right hippocampus, 69 genes could not be found in the PANTHER database. The remaining

| Threshold for the GPRS computation | Right amygdala | Right hippocampus |
|-----------------------------------|----------------|-------------------|
| $P < 0.05$                        | $F_{1159} = 0.867$, $P = 0.353$ | $F_{1159} = 1.457$, $P = 0.229$ |
| $P < 0.1$                         | $F_{1159} = 1.662$, $P = 0.199$ | $F_{1159} = 1.267$, $P = 0.262$ |
| $P < 0.2$                         | $F_{1159} = 1.662$, $P = 0.199$ | $F_{1159} = 1.267$, $P = 0.262$ |

Note: $F$-values indicate the interaction effects of the infant’s GPRS with antenatal maternal depression on the structural volumes.

Figure 2. Statistical maps of the interaction effects of the infant GPRS with antenatal maternal depressive symptoms (A) on the amygdala and hippocampal shapes. Panels (B, C) illustrate the relationship of antenatal maternal depressive symptoms with right amygdala shape (left panel) and right hippocampal shape (right panel) in neonates with high GPRS for major depressive disorder (GPRS, blue asterisks) and those with low GPRS (magenta circles). Antenatal maternal depressive symptoms are represented by the score of EPDS, with a higher score indicating more severe of depressive symptoms.

Figure 3. Statistical maps for interaction of the infant GPRS with antenatal maternal depressive symptoms (A) on cortical thickness. Panel (B) illustrates the relationship of antenatal maternal depressive symptoms with thickness of left OFC and vmPFC in neonates with high GPRS (blue asterisks) and those with low GPRS (magenta circles). Antenatal maternal depressive symptoms are represented by the score of EPDS, with a higher score indicating more severe of depressive symptoms.
439 genes formed 2 statistically over-represented genetic pathways, including “axon guidance mediated by Slit/Robo” (P = 0.027) and “Alzheimer’s disease-amyloid secretase” (P = 0.047). Supplementary Table S3 lists the genes involved in each pathway.

The same analysis for SNPs in GRDMS that nominally moderate the association between SES and fetal neurodevelopment did not reveal any significantly enriched pathways for the right amygdala. For the right hippocampus, 40 genes could not be found in the PANTHER database and the remaining 311 genes formed 4 statistically over-represented pathways (Table 4), including “CCK receptor signaling” (P = 0.009), “Beta1” (P = 0.021) and “Beta2” (P = 0.021) adrenergic receptor signaling”, and “5-HT2 type receptor” mediated signaling pathways (P = 0.021). Supplementary Table S4 lists the genes involved in each pathway.

We then entered the genes from the individual pathways identified by PANTHER into GeneMania and identified candidate biological processes (Table 5). GeneMania identified “Glutamate receptor activity” as the primary function of the metabotropic glutamate receptor group III pathway, which included genes that encode for subunits of the metabotropic-D-aspartate (GRIN1, GRIN2A, GRIN2B, GRIN2C, and GRIN2D), Kainate (GRK1-5), and a-alpha-3-hydroxy-5-methyl-4-isoxazol-lepropionic acid (GRIA1-4) receptors, as well as for the metabotropic receptors (GMR1-8). This same pathway was also significantly associated with the SNAP receptor (SNARE) complex function. Likewise, the SNARE complex emerged as the primary biological function associated with both the 5-HT2-type receptor- and the oxytocin receptor-mediated signaling pathway identified as candidate moderators of the effect of antenatal maternal depressive symptoms on cortical thickness (Table 5).

The Findings from the US Cohort

We also examined the neonatal amygdala and hippocampus structures using the US sample of 85 dyads with available neonatal brain imaging data of the amygdala and hippocampal volumes, SNP data genotyped using the Illumina Omni express arrays, and measures of maternal depressive symptoms assessed using the Center for Epidemiologic Studies Depression scale during the third trimester (the Supplementary Material). We used the same statistical model as that employed with the Asian cohort data and found a statistically significant interaction effect between infant GRDMS and antenatal maternal depressive symptoms on the right (F1,77 = 4.09, P = 0.047), but not left amygdala volume (F1,77 = 2.11, P = 0.275), with the largest interaction effect at GRDMS of P = 0.1 among the 3 thresholds (0.05, 0.1, and 0.2). Interestingly, the US cohort showed the significant positive association between antenatal maternal depressive symptoms and right amygdala volume (t176 = 2.21, P = 0.034) in neonates with low infant GRDMS (<0) and the marginal negative association between antenatal maternal depressive symptoms and right amygdala volume (t176 = −1.66, P = 0.105) in neonates with high infant GRDMS (>0), which is opposite to that seen in the Asian sample. This could be due to differences in populations, which has been reported in other studies on Asian and Caucasian populations based on individual SNPs (Sheikh et al. 2013) (Woo et al. 2004; Serretti et al. 2007; Forcelli et al. 2012; Ming et al. 2013, 2015). Nevertheless, interaction effects between infant GRDMS and antenatal maternal depressive symptoms were not observed on bilateral hippocampal volumes (left: F1,77 = 0.064, P = 0.801; right: F1,77 = 0.903, P = 0.345).

Among the SNPs involved in the GRDMS for the US sample, the top 747 SNPs with the most significant interaction with antenatal maternal depressive symptoms (P < 0.1) on the right amygdala volume were mapped to 594 genes. For the right amygdala, 65 genes could not be found in the PANTHER database. The remaining 529 genes formed 2 statistically over-represented pathway labeled by PANTHER as “metabotropic glutamate receptor group III” (P = 0.002) and “metabotropic glutamate receptor group I” (P = 0.003). Thus, despite the differences in sample size and ethnicity, these biological pathways were strikingly similar to those identified in the Asian cohort for the interaction between antenatal maternal depressive symptoms and GRDMS on the right amygdala volume.

Discussion

This study provides evidence for the role of infant genotype as a moderator of the association of antenatal maternal mental health with fetal development of cortico-limbic structures.

Table 4 Genetic pathways associated with interaction of GRDMS and antenatal maternal depressive symptoms in the Asian sample

| Structure            | Genetic pathways                                                                 | Fold enrichment | P-value*   |
|----------------------|----------------------------------------------------------------------------------|----------------|------------|
| Interaction of GRDMS and antenatal maternal depression | Metabotropic glutamate receptor group III                                 |                | 0.003      |
| Right amygdala       | Metabotropic glutamate receptor group III                                 | >5             | 0.004      |
|                      | Heterotrimeric G-protein signaling pathway-Qg alpha                            | >5             | 0.005      |
|                      | and Go alpha mediated pathway                                                  | >5             | 0.007      |
|                      | Histamine H1 receptor-mediated signaling                                       | >5             | 0.013      |
|                      | Endothelin signaling                                                          | >5             | 0.044      |
|                      | 5-HT2 type receptor-mediated signaling                                        | >5             | 0.027      |
|                      | Oxytocin receptor-mediated signaling                                         | >5             | 0.047      |
| Right hippocampus    | Axon guidance mediated by Slit/Robo                                            | >5             | 0.009      |
|                      | Alzheimer’s disease-amyloid secretase                                         | >5             | 0.021      |

Notes: The Fold Enrichment is defined as the ratio of the proportion of our input genes involved in a particular genetic pathway to the proportion of the reference genes involved in the same genetic pathway.

*Bonferroni corrected P-value.
including structures such as the amygdala, hippocampus, OFC, and vmPFC that are implicated in affective disorders. These interaction effects were apparent using measures of depressive symptoms across a community sample and remained significant even after controlling for maternal GRPS$_{MDD}$ thus revealing selective moderation by infant genotype. This study further revealed over-representative genetic pathways and biological processes formed by the genes included in the GRPS$_{MDD}$ computation for which variations best contributed to modulating the relationship of antenatal maternal depressive symptoms and the morphology of the aforementioned brain structures.

The amygdala, hippocampus, OFC, and vmPFC are cortico-limbic structures implicated in mood disorders and central to emotion and social regulation. Abnormal morphology and activity in these regions are reported in children (Thomas et al. 2001; Pagliaccio et al. 2014) and adults with MDD (Johnstone et al. 2007). Smaller hippocampal volume and greater activity in response to sad faces predict increased depressive symptoms in children (Suzuki et al. 2013). In depressed children severity correlates with activity of the amygdala–hippocampal complex and medial PFC during sad mood elaboration (Pagliaccio et al. 2012) and amygdala reactivity during emotional face viewing (Gaffrey et al. 2011). Likewise, children with early-life stress show smaller OFC (Hanson et al. 2010). Children of parents with MDD also show altered amygdala activity in the emotional facial expression (Monk et al. 2008). Increased amygdala activation in response to novelty or threatening faces in children is associated with measures of negative emotionality (Perez-Edgar et al. 2007), which in turn predicts a greater risk for mood disorders (Bruder-Costello et al. 2007). Moreover, altered functional connectivity between the amygdala and medial PFC, including OFC and vmPFC, is observed in 6-month-old infants born to mothers with increased antenatal maternal depressive symptoms (Qiu et al. 2015a). These findings suggest that the cortico-limbic systems may be particularly vulnerable to maternal influences during fetal development and reflect, in part, the neural basis for the familial transmission of phenotypes associated with depression.

Our study revealed an interaction effect between maternal depressive symptoms and infant GRPS$_{MDD}$ that was apparent in the right amygdala and hippocampus. Individual differences in stress reactivity associate with morphology and function of the prefrontal cortex, the hippocampus and the amygdala, as well as the connections between these regions (Pruessner et al. 2008; van Marle et al. 2009) and there is compelling evidence implicating the right hemisphere, notably the right amygdala and right hippocampus in the regulation of stress responses (Abercrombie et al. 1998; Carrion et al. 2007; Hamilton and Gotlib 2008). The risk for depression is linked to heightened stress reactivity, which associates with a familial risk for depression: individuals with increased genetic vulnerability for depression show greater negative affect in response to daily life stress, which then predicts depression (Wichers et al. 2009). Our findings suggest the familial transmission of individual differences in neural systems that underlie stress reactivity may occur during fetal development.

We explored gene networks and their associated biological processes as candidate mediators of the influence of clinically relevant environmental factors on fetal neurodevelopment. The gene pathways (metabotropic glutamate receptor group pathways) derived from SNPs that best mediated the interaction of antenatal maternal depressive symptoms and infant GRPS$_{MDD}$ on the right amygdala in both the Asian and US cohorts were strongly associated with glutamate receptor activity. Multiple approaches, including genome-wide analyses, implicate glutamatergic synaptic transmission in the etiology of depression (Skolnick et al. 2009; Lee et al. 2012) and patients with MDD exhibit abnormal glutamate concentrations in the amygdala (Michael et al. 2003). Postmortem studies with MDD suggest altered glutamatergic synaptic signaling (Hashimoto et al. 2007) and ketamine, a glutamatergic intervention, is a treatment for MDD (Caddy et al. 2014). The antidepressant, tianeptine, blocks stress-induced increases in glutamate release in the amygdala (Pirolli et al. 2013) and moderates behavioral responses to stress. This glutamate receptor activity included the GRM1 and GRM8 genes, which are nominally associated with MDD (Terracciano et al. 2010). Knockout mouse models of genes coding for metabotropic glutamate receptor proteins, which modulate postsynaptic glutamate signaling, associate with increased anxiety (Bures et al. 2005). The “G-protein signaling” network also identified as a candidate mediator for the effects of maternal depression on cortical structure (Supplementary Table S3) includes the GRM1, GRM4, GRM7, and GRM8 genes.

The genetic pathway analysis in the right amygdala identified the “SNARE complex” biological process, which is a fundamental component of neural connectivity and neurotransmitter release comprised of syntaxin, SNAP-25, and VAMP. Indeed, perhaps the most compelling finding from our analysis was the broad implication of the SNARE complex as a biological process mediating the impact of antenatal maternal depressive symptoms on multiple brain structures (Table 5). Postmortem studies revealed abnormal expression of SNARE proteins in the hippocampus and frontal cortex of subjects with MDD (Scarr et al. 2006). Likewise, proteomic analyses implicate synaptic

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**Table 5 Biological functions of genetic pathway in the Asian sample**

| Genetic pathway | Biological function | FDR corrected P-value | Coverage |
|-----------------|---------------------|-----------------------|----------|
| Metabotropic glutamate receptor group III | Glutamate receptor activity | 7.3e–46 | 20/20 |
| G-protein signaling | SNARE complex | 3.2e–18 | 12/32 |
| Histamine H1 receptor-mediated signaling | G-protein coupled receptor signaling | 1.2e–32 | 29/125 |
| Endothelin signaling | Inositol phosphate metabolic process | 6.0e–19 | 13/55 |
| 5-HT2 type receptor-mediated signaling | Fibroblast growth factor signaling | 1.5e–43 | 35/170 |
| Oxytocin receptor-mediated signaling | SNARE complex | 3.1e–25 | 15/32 |
| Axon guidance mediated by Slit/Robo | SNARE complex | 5.0e–26 | 15/32 |
| Amyloid secretase | Small GTPase mediated signal transduction | 3.9e–8 | 12/291 |
| Beta1 adrenergic receptor signaling | Neurotrophic signaling | 2.4e–20 | 23/277 |
| Beta2 adrenergic receptor signaling | SNARE complex | 1.6e–27 | 15/32 |
| CCKR signaling | Intracellular ligand-gated calcium channel activity | 3.1e–14 | 7/10 |

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dysfunction in MDD and identify altered expression of SNARE complex proteins (Martins-de-Souza et al. 2012). Administration of ketamine as well as a broad range of antidepressant medications alters the accumulation of SNARE complexes in synaptic membranes (Bonanno et al. 2005), which then associates with altered capacity for stimulated glutamate release (Bonanno et al. 2005). Two SNP’s (rs3787283 and rs3746544) in the SNAP-25 gene have been associated with MDD in a Han Chinese sample (Wang et al. 2015), an ethnic group that accounted for the majority of the Asian subjects in our study.

Our findings suggest that the influence of antenatal maternal depressive symptoms on the cortico-limbic structure is mediated by brain-specific signaling pathways (oxytocin, 5-HT2, and glutamate receptor pathways). These pathways are linked to processes underlying synaptic development. In addition to association with the glutamate receptor activity, the SNARE complex was the top biological process for the oxytocin and 5-HT2 gene networks (Table 5). There is strong evidence for the importance of serotonin, including 5-HT2 receptor activation, in neonatal brain structure (see Lesch and Waider 2012 for a review). While oxytocin receptor activation is not extensively described in terms of brain development, there is a period of transient neocortical expression of oxytocin receptor binding in rodent brain that suggests a developmental influence (Hammock and Levitt 2013).

Finally, our analyses identified neurotrophic signaling systems as candidate mediators of the effects of antenatal maternal depression on hippocampus (Table 4). The Fibroblast growth factor signaling system is particularly interesting to consider in light of the findings with rodent models that early postnatal administration of FGF2 decreases fear behavior in animals genetic predisposed to increased expression of anxiety-like behaviors (Eren-Kocak et al. 2011). Sustained FGF2 effects on genes expression included targets involved in cell survival, an effect that is consistent with the role for neurotrophic signaling pathways as a mediator of the effects of antenatal maternal depression. Neurotrophic signaling pathway was the top biological process emerging from the amyloid secretase network and includes genes coding for proteins that mediate the effects of brain-derived neurotrophic factor signaling, implicated in the pathophysiology of MDD and associated with the activation of inositol phosphate metabolism (Duman and Voleti 2012), which was also a top biological process identified in our analysis. Taken together, our findings are consistent with the hypothesis that environmental conditions that increase the risk for MDD associate with alterations in mechanisms controlling synaptic plasticity in cortico-limbic brain regions (Duman and Aghajanian 2012).

Interestingly, the direction of the interaction between maternal depressive symptoms and infant genotype on right amygdala volume in the US sample is in the opposite direction to that observed in the Asian sample. This is not surprising as there is now considerable evidence for opposing effects of individual SNPs between Asian and Caucasian populations. For instance, Sheikh et al. 2013 found that during childhood Caucasian children homozygous for the Catechol-O-methyltransferase (COMT) val allele show higher levels of internalizing symptoms compared to children with at least one copy of the COMT met allele. In contrast, Baumann et al. 2013 found that in Asians homozygous of the COMT met allele with adverse childhood experiences show higher anxiety sensitivity. Similar findings are apparent in studies of the serotonin transporter polymorphism (5-HTTPLR). Ming et al. 2015 report a replication of the well-established interaction between the 5-HTTPLR and stressful life events on emotional health in Han Chinese, the principal ethnic group in our Asian sample. However, in this study it was the L and not the S allele that conferred greater sensitivity to stress; studies with Caucasian samples commonly report that the S allele confers sensitivity to adversity. This study was also a replication of a previous, similar finding with the 5-HTTPLR variant (Ming et al. 2013). Meta-analyses likewise reveal differential effects of 5-HTTPLR on selective serotonin reuptake inhibitors efficacy between Caucasians and Asians (Serretti et al. 2007; Porcelli et al. 2012). Importantly, functional MRI studies show a link between S allele and higher amygdala activation in response to emotional stimuli in Caucasians (e.g., Hariri et al. 2002, 2005; Murphy et al. 2013), whereas in Asians L allele is associated with amygdala hyperactivation (Li et al. 2012). These findings imply that the same risk allele may operate in the opposite fashion as a function of ethnic background. Allele frequency may be a factor. The frequency of the L allele is much lower than that of S allele in Chinese population and Japan populations, but higher in Caucasians or African-Americans (Lesch et al. 1996; He et al. 2010).

Importantly, despite the directional difference, the pathway analyses revealed the same underlying biological processes for the amygdala effect in both samples. Indeed, PANTHER identified the same primary pathway for both samples using independent analyses. These findings are consistent with the studies noted above, that reveal effects of polymorphisms in the same genetic regions, consistent with common underlying biological processes, but with polymorphisms operating in an ethnicity-specific manner.

The relationship between antenatal maternal depression and children’s outcomes is often observed as a function of gender. Recent evidence suggests that girls born to depressed mothers may be more likely to exhibit increased cortisol reactivity in childhood and subsequently altered amygdala–prefrontal cortex activity in adolescence (Burghy et al. 2012). Nevertheless, our study did not observe the gender effect, which may partly because of the age of children in this study. It remains possible that gender differences could be observable at later stages in development.

This study is best considered as exploratory and intended to provide a strategy for the analysis of gene–environment interdependence and identify candidate biological processes that might mediate the influence of environmental conditions associated with the risk for MDD on fetal brain development. While preliminary, our findings do suggest biologically plausible candidate processes in two independent cohorts. While the neonatal neuroimaging dataset in this study is highly unique in its timing of acquisition and number of subjects, it is nevertheless a modest sample size for genomic analyses. Due to the time limit for scanning of neonates and brain immaturity, the neonatal brain MRI, in general, has relatively low contrast between gray and white matter, which could be challenging for imaging and the brain segmentation. The contrast between gray matter and white matter is relatively clear on T2-weighted MRI at the neonatal stage but is diminished around 4–9 months of age. We reported the brain segmentation accuracy in the Materials and Methods section to indicate our constraint with the neonatal brain images. An additional limitation is the use of GWAS datasets derived from Caucasians for the definition of a GPRS to study within an Asian sample. While this is an important caveat, we note that the Wang et al. (2015) study identified multiple SNPs in the SNAP-25 gene, which is a member of the SNARE complex, in the pathophysiology of MDD in Asian population. Moreover, our assessment of maternal depressive symptoms was based on a common screening tool intended to elicit a subjective report of emotional well-being, but does not constitute clinical assessment. The brain variations in the offspring are thus best considered as being associated with self-reported depressive symptoms and not to clinical depression,
per se. Furthermore, other factors, such as marital discord, paterno
tal mental illness, medication during pregnancy, maternal sub-
stance use, or breastfeeding, could also influence the fetal brain
development over and beyond antenatal maternal depression
since the brain develops rapidly in early life (Uematsu et al. 2012).
In our study, SES was highly correlated with antenatal maternal
depressive symptoms and both showed its interaction with
GPRS\textsubscript{MDD}, and influence the right amygdala and hippocampus
(Supplementary Table S2 and Figures S1 and S2). After controlling
for each other, the interaction effect of antenatal maternal depres-
sion and GPRS\textsubscript{MDD} was mainly shown on the right amygdala,
while the interaction effect of SES and GPRS\textsubscript{MDD} was mainly
shown on the right hippocampus (the Supplementary Material).
These findings suggested that environment conditions associated
with the risk for MDD interplay with genetic risks for MDD on fetal
brain development. Finally, comparing brain anatomical mea-
sures across different cohorts, particularly, on neonates, is chal-
lenging. This is partly because of different strength of the MRI
scanners and imaging acquisition protocols. More importantly,
the challenge also lies in the definition of the anatomy of individ-
ual brain structures. In this study, we obtained the comparable
volume measure for the amygdala from the two cohorts. Hence,
this study incorporated the amygdala findings from both cohorts
where we could be certain of comparable anatomical localization.
Furthermore, the US cohort was lack of maternal GPRS\textsubscript{MDD} infor-
mation and hence our reported results for the US cohort could not
count for the contribution of maternal GPRS\textsubscript{MDD}.

Conclusion
We present a hypothesis-generating approach to the study of
gene × environment interaction that uses GPRS together with
bioinformatic approaches to identify candidate biological pro-
cesses mediating the influence of environmental conditions on
brain development and function. This approach is considered as
hypothesis-generating. For example, in larger samples, with
sufficient statistical power, we predict that a multi-gene locus
score, reflecting polymorphisms in genes involved in the bio-
logical processes of glutamate receptor activity, SNARE com-
plex, 5-HT2 receptor signaling, should moderate brain region-
specific effects of “depressogenic” environmental conditions
during fetal development.

Supplementary Material
Supplementary data is available at Cerebral Cortex online.

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