Common and Dissociable Regional Cerebral Blood Flow Differences Associate with Dimensions of Psychopathology Across Categorical Diagnoses

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Abstract

The high comorbidity among neuropsychiatric disorders suggests a possible common neurobiological phenotype. Resting-state regional cerebral blood flow (CBF) can be measured noninvasively with MRI and abnormalities in regional CBF are present in many neuropsychiatric disorders. Regional CBF may also provide a useful biological marker across different types of psychopathology. To investigate CBF changes common across psychiatric disorders, we capitalized upon a sample of 1,042 youths (ages 11 to 23 years) who completed cross-sectional imaging as part of the Philadelphia Neurodevelopmental Cohort. CBF during a resting state was quantified on a voxelwise basis using arterial spin labeled perfusion MRI at 3T. A dimensional measure of psychopathology was constructed using a bifactor model of item-level data from a psychiatric screening interview, which delineated four factors (fear, anxious-misery, psychosis,
and behavioral symptoms) plus a general factor: overall psychopathology. Overall psychopathology was associated with elevated perfusion in several regions including the right dorsal anterior cingulate cortex (ACC) and left rostral ACC. Furthermore, several clusters were associated with specific dimensions of psychopathology. Psychosis symptoms were related to reduced perfusion in the left frontal operculum and insula, whereas fear symptoms were associated with less perfusion in the right occipital/fusiform gyrus and left subgenual ACC. Follow-up functional connectivity analyses using resting-state fMRI collected in the same participants revealed that overall psychopathology was associated with decreased connectivity between the dorsal ACC and bilateral caudate. Together, the results of this study demonstrate common and dissociable CBF abnormalities across neuropsychiatric disorders in youth.

**Keywords**
cerebral blood flow; perfusion; psychopathology; development; anterior cingulate

**INTRODUCTION**

Co-morbidity among psychiatric disorders is highly common. Epidemiological research confirms that of the 26.2% of adults with a mental health diagnosis in the U.S., 55% have one diagnosis, while 22% carry two mental health diagnoses, and 23% carry three or more.\(^1\) Comorbidity and a lack of clear diagnostic boundaries may be particularly challenging in children, where clinical phenotypes are often less distinct.\(^2\) For example, a meta-analysis of community sample based studies on comorbidity between childhood psychiatric disorders showed that diagnoses such as attention-deficit/hyperactivity disorder (ADHD), conduct disorder, depression, and anxiety were all highly comorbid.\(^3\)

The comorbidity among neuropsychiatric disorders suggests common neurobiological origins, which may confer vulnerability to a range of symptoms. Prior neuroimaging studies have typically focused on adults in a case control design that compares a specific disorder to a control group. Given the high comorbidity among disorders, studies limited to those with “pure” singular diagnoses may not be representative of the actual presentation of psychiatric symptoms. Therefore, the search for common neural abnormalities that are present across disorders has recently accelerated.\(^4\)–\(^13\) One possible common neuroanatomical substrate was reported by Goodkind and colleagues,\(^10\) who showed that gray matter loss in the anterior insula and dorsal anterior cingulate cortex (ACC) were common across psychotic and nonpsychotic disorders in adults. Thus, abnormalities in the ACC may represent a risk factor shared across diagnoses.

While most prior across-disorder studies have focused on brain structure or signals from fMRI, regional cerebral blood flow (CBF) has been much less studied. Regional CBF, which can be quantified noninvasively using arterial spin labeled (ASL) perfusion MRI, is tightly coupled to regional brain metabolism.\(^14\) Alterations in regional CBF have been demonstrated in neuropsychiatric disorders, such as generalized anxiety disorder and posttraumatic stress disorder,\(^15,16\) and during stress provocation paradigms.\(^17\) However,
these case-control studies did not measure psychopathology symptoms across multiple disorders.

There is a growing recognition that psychiatric symptoms exist on a continuum in contrast to a categorical approach to diagnosis. Thus, dimensional measures of psychopathology that cut across categorical clinical diagnoses are needed.\textsuperscript{18,19} Such measures may enhance the power to detect differences compared to healthy controls and may improve interpretation of these differences.\textsuperscript{19} A number of models have been proposed to identify broad traits shared across psychiatric disorders, such as measures of neuroticism and negative emotionality.\textsuperscript{20,21} In particular, Lahey and colleagues used a bifactor model to quantify psychopathology dimensionally across psychiatric disorders.\textsuperscript{22} They identified a general factor associated with psychopathology as well as dimensions such as externalizing, distress, and fear. Other studies have shown a similar bifactor structure, with factors for anxious-misery, psychosis, behavioral (externalizing), and fear as well as a general factor representing overall psychopathology.\textsuperscript{12} Bifactor analysis has the advantage of identifying a general factor and orthogonal factors among highly correlated symptoms. Previously, we have used this approach to document that overall psychopathology was associated with reduced activation within executive regions including the ACC on a dimensional basis across psychiatric symptoms in youth.\textsuperscript{12}

In contrast to existing studies of brain structure\textsuperscript{10} and executive function,\textsuperscript{12} few studies have examined associations between CBF and dimensional measures of psychopathology across clinical diagnoses, and fewer still have done so in youth. In one previous study,\textsuperscript{23} we examined the association between CBF and a dimensional measure of mood and anxiety using the State Trait Anxiety Inventory. We found that greater mood/anxiety traits were associated with elevated CBF in the left amygdala, bilateral insula, and left fusiform gyrus in adolescence.\textsuperscript{23} This suggests that CBF may be a useful biological marker of dimensions of emotion. However, we only examined anxiety and mood traits in that study and did not directly examine overall symptoms of psychopathology.\textsuperscript{23}

In this study, we investigated a potential common neurobiological phenotype related to regional resting-state CBF using a dimensional measure of psychopathology symptoms across a broad range of neuropsychiatric disorders in a large sample of children, adolescents, and young adults imaged as part of the Philadelphia Neurodevelopmental Cohort (PNC). We applied a bifactor analysis to dimensionally quantify psychopathology.\textsuperscript{12} As suggested by both our prior work on executive function\textsuperscript{12} and the results of Goodkind et al.,\textsuperscript{10} we hypothesized that greater overall psychopathology would be associated with CBF changes within previously-implicated regions such as the ACC. In order to further understand the functional implications of such differences, we conducted follow-up analyses examining changes in functional connectivity in cingulate regions where alterations in CBF were found. Finally, while the focus of this study was on regional CBF changes associated with overall psychopathology, we also explored dissociable relationships between regional CBF and specific dimensions of psychopathology.
METHODS AND MATERIALS

Participants

Of the total sample of 1,601 youths imaged in the Philadelphia Neurodevelopmental Cohort (PNC), we included participants age 11 years and older who completed a psychiatric assessment interview (n=1,288); participants under 11 who only received a collateral assessment were not included. An additional 127 participants were excluded for: medical disorders that could affect brain functioning (n=68), non-psychiatric medication use that could impact CNS functioning (n=53), or substantial structural brain abnormalities (n=16); several subjects were excluded for multiple criteria. Of the remaining participants, 119 participants were excluded for missing clinical data (n=2), failing to complete perfusion imaging (n=5), or failing to meet ASL and structural image quality assurance protocols (n=117); 5 subjects were excluded for multiple reasons. This yielded a final sample of 1,042 participants (mean age = 16.12 years; range = 11–23 years; SD = 2.82 years; 468 males). Among this final sample, 120 participants (12%) were taking psychiatric medications at the time of imaging and were evaluated in sensitivity analyses, as described below.

Clinical assessment

The institutional review boards of both the University of Pennsylvania and the Children’s Hospital of Philadelphia approved all study procedures. Informed consent was obtained from all participants. As described previously, psychiatric symptoms were assessed using a structured screening interview (GOASSESS) based on a modified version of the Kiddie-Schedule for Affective Disorders and Schizophrenia (see Supplementary Methods). Demographic data and lifetime prevalence are summarized in Table 1.

Factor analysis to create a dimensional measure of psychopathology

As described in detail elsewhere, (see Supplementary Methods) we used a confirmatory bifactor analysis implemented in Mplus to quantify a dimensional measure of psychopathology. This analysis yielded four orthogonal dimensions of psychopathology (anxious-misery, psychosis, behavioral (externalizing), and fear), plus a fifth factor that was common across all psychiatric disorders, which we termed overall psychopathology (Figure 1).

Image acquisition, preprocessing, and perfusion quantification

Image acquisition and processing are reported in detail elsewhere; see Supplementary Methods for details. Perfusion was measured using a pseudo-continuous arterial spin labeling (pCASL) sequence sampled with spin-echo echoplanar imaging. These data were then pre-processed using standard tools included with FSL. Following distortion correction using the B0 map with FUGUE, the first four image pairs were removed, the timeseries were realigned in MCFLIRT, the skull was removed with BET, and the images were smoothed at 6mm FWHM using SUSAN. Resting-state CBF was quantified from control-label pairs using ASL Toolbox. As prior, the T1 relaxation parameter was modeled on an age- and sex-specific basis. This model accounts for the fact that T1 relaxation time differs according to age and sex, and has been shown to enhance the
accuracy and reliability of results in developmental samples.\textsuperscript{39} Participant-level CBF images from each subject were co-registered to the T1 structural image using boundary-based registration,\textsuperscript{40} and normalized to the PNC adolescent template using the topperforming deformable registration included in Advanced Normalization Tools.\textsuperscript{41–43} Images were downsampled to 2-mm resolution prior to group-level analysis. All transformations were concatenated so that only one interpolation was performed. After statistical testing, images were registered to the Montreal Neurologic Institute (MNI) 152 1-mm template space for reporting of standard coordinates and display.

**Group-level analyses**

We conducted voxelwise analyses that modeled overall psychopathology as well as each of the four psychopathology sub-factors. We included model covariates to control for subject demographics as well as data quality. Specifically, due to known nonlinear age-by-sex interactions in the development of CBF during adolescence,\textsuperscript{37} the model included age, age squared, sex, an interaction between sex and age, and an interaction between age squared and sex. Furthermore, to control for the known confounding effects of motion in developmental populations,\textsuperscript{31,44} motion during the ASL acquisition (mean frame displacement) was also included as a covariate. Thus, our group level model was as follows:

\[
\text{CBF}_{\text{voxelwise}} = \text{age} + \text{sex} + \text{age}^2 + \text{age} \times \text{sex} + \text{age}^2 \times \text{sex} + \text{in-scanner motion} + \text{anxious-misery} + \text{psychosis} + \text{behavioral} + \text{fear} + \text{overall psychopathology}
\]

We also examined age by sex effects in regions found to be significantly associated with overall psychopathology, as well as interactions between overall psychopathology and both age and sex. Based on recent evidence that lower thresholds provide inadequate Type I error control,\textsuperscript{45} we used AFNI 3dClustSim (version 17.0.13) with a cluster defining threshold of \( p = 0.001 \ (z = 3.09) \) and a corrected cluster significance of \( p < 0.01 \). Data were normally distributed. For additional details, see the Supplementary Methods section.

**Sensitivity analyses**

Sensitivity analyses were conducted after excluding the minority (12%) of participants who were treated with psychiatric medication at the time of imaging and adding both race and maternal level of education as additional covariates. Lastly, due to the known relationship between brain structure and CBF, we re-evaluated associations between psychopathology and CBF while including gray matter density within each cluster as an additional model covariate (see Supplementary Methods).

**Resting state functional connectivity**

As a final step, in order to further understand the functional implications of our results, we conducted seed-based connectivity analyses that evaluated the functional connectivity of the dorsal ACC seed, which showed a significant association between CBF and overall psychopathology and has been implicated in prior studies.\textsuperscript{10} Functional connectivity was available for 833 participants who received both ASL and resting-state imaging as part of the PNC; these data were processed using a previously-described pipeline that minimizes the
Results

Elevated CBF in the ACC is associated with overall psychopathology

We began by testing the hypothesis that greater overall psychopathology would be associated with regional resting-state CBF differences. Voxelwise analyses revealed that greater overall psychopathology was associated with elevated perfusion in several regions including the dorsal anterior cingulate cortex (ACC) and rostral ACC (Figure 2). Other regions that also showed higher CBF included the right postcentral gyrus, parahippocampal cortex, and midbrain (Table S1). We found significant associations between age and perfusion in many of these regions, with CBF declining with age in the right postcentral gyrus, dorsal ACC, and rostral ACC. Furthermore, consistent with our prior work, linear age by sex interactions were found in all clusters: perfusion declined in males, while perfusion either declined more slowly or increased in females over the age range sampled (Figure S1). However, there were no significant two-way or three-way interactions between overall psychopathology, age, and sex in regions where a main effect of overall psychopathology was present. Sensitivity analyses showed that these results were similar when participants taking psychiatric medications were excluded from the analysis, and when participant race and maternal education were included in the model (Table S2). Notably, results were unchanged when cluster gray matter density was added as a model covariate (Table S3).

Regional cerebral blood flow differences associated with the symptom-specific sub-factors

Next, we examined each of the orthogonal sub-factors representing distinct dimensions of psychopathology (anxious-misery, psychosis, behavioral, and fear) to investigate dissociable associations with resting-state CBF. We found that psychosis-spectrum symptoms were related to reduced perfusion in the left frontal operculum/left insula, whereas fear symptoms were associated with reduced perfusion in the right occipital/fusiform gyrus and left subgenual ACC (Figure 3 and Table S1). No significant results were found for the anxious-misery or behavioral factors. As expected, age by sex interactions were also found in these regions, with perfusion declining more steeply in males than in females (Figure S2). No significant interactions between sub-factors and age or sex were found in these regions. Results remained similar after a sensitivity analysis that excluded participants taking psychiatric medications and after including participant race and maternal education as covariates (Table S2), or when cluster gray matter density was added as a covariate (Table S3).

Dimensional psychopathology is associated with reduced connectivity between dorsal ACC and bilateral caudate

Finally, based on prior work implicating the dorsal ACC in psychopathology, we conducted seed-based connectivity analyses to further understand the functional implications...
of CBF abnormalities in the dorsal ACC. These seed-based analyses revealed that higher levels of overall psychopathology were associated with diminished connectivity between dorsal ACC and bilateral caudate (Figure 4). Overall psychopathology was also associated with diminished connectivity between the dorsal ACC and other regions including the right thalamus, supramarginal gyrus, and right putamen, as well as increased connectivity with the dorso-medial frontal cortex (Table S4).

**DISCUSSION**

In this study, we provide novel evidence that higher resting-state CBF in multiple regions including the ACC is associated with overall psychopathology. This dimensional measure of overall psychopathology was also associated with reduced dorsal ACC functional connectivity with regions such as the caudate. Finally, specific dimensions of psychopathology such as fear and psychosis were associated with diminished CBF in several regions. Taken together, the results suggest the existence of both common and dissociable perfusion abnormalities across traditional categorical psychiatric diagnoses.

**Advantages of a dimensional approach to psychopathology**

To dimensionally quantify overall psychopathology, we used a bifactor analysis\(^{28,29}\) of the item-level data from a clinician-administered screening interview. As we have shown previously,\(^{12}\) this identifies orthogonal latent dimensions of psychopathology (anxious-misery, psychosis, behavioral, and fear) as well as a dimensional measure of general psychopathology. This approach is advantageous for several reasons. First, the bifactor model allows us to estimate variance that is shared across highly comorbid disorders.\(^ {29} \) Studies comparing psychopathology across more than one disorder typically measure comorbidity as shared variance, which is controlled for but not directly evaluated. Second, this approach yields continuous dimensions of psychopathology, rather than “all or none” diagnostic labels. These dimensions allow evaluation of the degree to which individual differences in perfusion are associated with both sub- and supra-threshold symptoms. Third, traditional designs that compare multiple categorical diagnoses are often hindered by varying frequency of, and co-morbidity among, disorders. A bifactor model accounts for such structured covariance and yields orthogonal dimensions that can be included as predictors in a single model.\(^ {29} \)

**Overall psychopathology is associated with common perfusion abnormalities**

Using this bifactor approach, the present study demonstrated that common perfusion abnormalities are present across psychiatric diagnoses. Overall psychopathology was associated with elevated resting-state CBF in a network of regions including the anterior cingulate cortex (ACC). These ACC perfusion abnormalities are consistent with prior metanalytic investigations of case-control structural and functional imaging studies in major depressive disorder,\(^ {49,50} \) bipolar disorder,\(^ {5,51} \) anxiety disorders,\(^ {4,52} \) ADHD,\(^ {53,54} \) and psychosis.\(^ {5,55} \) This also accords with several across-disorder studies that have provided evidence for structural,\(^ {10} \) task-based fMRI,\(^ {12} \) and resting state functional connectivity\(^ {13} \) abnormalities in the ACC. Given that the ACC is central to numerous processes including cognitive control,\(^ {56} \) reward-related decision-making,\(^ {57} \) and affect regulation,\(^ {58} \) these results
suggest that dysfunction in the ACC may impact multiple cognitive processes, and confer vulnerability to a wide variety of psychiatric symptoms.

The traditional conceptualization of the role of the ACC suggests that the dorsal anterior cingulate is involved in effortful regulation of emotional responses, thereby modulating affective limbic regions. Thus, activation of the dorsal ACC might be anticipated to potentially reduce a broad range of psychiatric symptoms, which would result in a negative correlation between ACC activation and symptomatology. However, meta-analyses of case-control studies implicate increased activation in the ACC during task-based fMRI studies in anxiety and depression, symptoms that are common across many disorders. The results of these meta-analyses are consistent with the findings of the current study and suggest that higher ACC perfusion may be shared across psychopathology.

Beyond the ACC, we also found elevated perfusion in the postcentral gyrus, parahippocampal cortex, and midbrain. Prior studies document sensorimotor abnormalities in regions such as the postcentral gyrus in incarcerated juveniles with high impulsivity, attention-deficit/hyperactivity disorder, autism spectrum disorders, and schizophrenia. Furthermore, hyperactivation of the parahippocampal cortex has been hypothesized to be a signature of stress in studies of childhood maltreatment.

In addition to associations between overall psychopathology and CBF, functional connectivity analyses revealed reduced connectivity between the dorsal ACC and the caudate. Abnormalities of the dorsal ACC within a cortico-striatal circuit have been implicated in the pathogenesis of depression, anxiety, ADHD, and psychosis. No interactions between overall psychopathology and age or sex were found. This suggests that ACC abnormalities are established early in development and remain stable over time. Dysfunctional ACC perfusion and connectivity between the ACC and caudate may represent important risk factors for many types of psychopathology in youth of both sexes.

**Orthogonal dimensions of psychopathology have dissociable relations with cerebral perfusion**

The current study extends prior work on neurobiological phenotypes associated with psychopathology by showing that several clusters of reduced perfusion were associated with specific dimensions of psychopathology. We found that psychosis symptoms were related to reduced perfusion in the left frontal operculum and insula, consistent with prior findings linking schizophrenia to structural and functional abnormalities in the insular cortex. Although there is a large body of work demonstrating a connection between psychosis and abnormalities in the cingulo-opercular network, which includes the anterior insula/operculum, dorsal anterior cingulate cortex, and thalamus, we found an association between psychosis and only one region in this network.

We also found that fear symptoms were associated with less perfusion in the left subgenual ACC and right occipital/fusiform gyrus. Prior work has implicated subgenual ACC hypofunction in anxiety and fear. The subgenual ACC is thought to regulate limbic regions (such as the amygdala) that are important for generating fear responses. Therefore, we speculate that reduced activity of the subgenual ACC, as manifested by reduced subgenual
ACC perfusion, may result in decreased regulation of these affective regions, leading to greater fear symptoms. Additionally, we found perfusion abnormalities in the occipital fusiform cortex, a region that is important for processing socially-relevant visual information such as faces, and is implicated in fear and anxiety.\textsuperscript{73} Taken together, these results suggest that in addition to common perfusion abnormalities associated with overall psychopathology, there are also dissociable perfusion abnormalities specific to psychosis and fear symptoms.

**Limitations**

Several limitations of the present work should be noted. First, the cross-sectional nature of the current study limits the ability to study developmental changes over time. Future work would benefit from longitudinal designs that allow investigation of when abnormalities in perfusion arise during development. Second, although the use of a community sample enhances the generalizability of these results, it would also be useful to examine these perfusion abnormalities in samples with more severe clinical presentations. Third, it should be noted that this study did not find specific significant associations between CBF and anxious-misery symptoms. In our earlier work,\textsuperscript{23} we found associations between a dimensional measure of anxious-misery (the State-Trait Anxiety Inventory, STAI) and higher perfusion of regions such as the insula and amygdala. Notably, these analyses of the STAI did not adjust for overall psychopathology as in the current bifactor model, and thus constitute a mix of overall and domain-specific anxious-misery symptoms. However, at lower thresholds both overall psychopathology and anxious-misery showed non-significant relationships with CBF in similar regions. This underlines the strengths and also limitations of the bifactor approach. Of note, the effect sizes in the current study were low (partial correlations ranging between .11 and .18), which may be due to the use of a community-based sample of non-help seeking youth, rather than a clinical population with more severe symptoms. However, small sample sizes with low statistical power are prone to overestimated effect sizes with low reproducibility.\textsuperscript{74} As the effect sizes reported here were derived from a much larger sample than most prior studies, they may provide a better estimate of the underlying magnitude of each association. It should also be acknowledged that the present results were found in a single sample, and should be replicated in future independent samples. Finally, ASL MRI technology has improved since these data were collected, and future studies will benefit from sequences which provide greater sensitivity and spatial resolution.\textsuperscript{75}

**Conclusions**

The current study provides novel evidence that elevated perfusion in regions including the ACC is associated with dimensional burden of overall psychopathology across psychiatric disorders in youth. This suggests a shared mechanism for psychopathology symptoms that cuts across clinical diagnostic categories, supporting the pathophysiology-based conceptualization of neuropsychiatric disorders proposed by the National Institute of Mental Health’s Research Domain Criteria.\textsuperscript{19} Perfusion abnormalities in the ACC may also suggest possible candidate biomarkers for future work on pharmacological and clinical interventions. Importantly, interventions that target these shared circuits may be beneficial to a broad range of psychiatric disorders and not limited to a specific categorical diagnosis.
Finally, longitudinal data could identify perfusion abnormalities in youth at risk for psychopathology and allow for biomarkers to guide the development of targeted early interventions during the vulnerable period of adolescence.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Bifactor model of psychopathology reveals common and divergent dimensions of psychopathology across categorical screening diagnoses
A) Confirmatory bifactor analysis of 112 items from the GOASSESS screening interview revealed four orthogonal dimensions of psychopathology (anxious-misery, psychosis, behavioral, and fear), plus a general factor shared across disorders (overall psychopathology). B) Mean factor scores for each orthogonal dimension of psychopathology are presented by screening categories with at least 20 subjects. Overall psychopathology was common across diagnostic screening categories. ADHD = attention deficit hyperactivity
disorder; GAD = generalized anxiety disorder; MDD = major depressive disorder; OCD = obsessive-compulsive disorder; ODD = oppositional defiant disorder; PTSD = posttraumatic stress disorder; TD = typically developing.
Figure 2. Overall psychopathology is linked to elevated perfusion in the anterior cingulate circuit.
Greater overall psychopathology across categorical clinical diagnoses was associated with elevated perfusion in A) right dorsal anterior cingulate cortex (ACC) and B) left rostral ACC. Images thresholded at $z > 3.09$, cluster corrected $p < 0.01$. 
Figure 3. Dissociable dimensions of psychopathology are associated with cerebral perfusion
Greater psychosis-spectrum symptoms were related to reduced perfusion in A) left frontal operculum/left insula, whereas fear symptoms were associated with less perfusion in B) left subgenual ACC and C) right occipital/fusiform gyrus. Images thresholded at $z > 3.09$, cluster corrected $p < 0.01$. 

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Overall psychopathology is associated with diminished functional connectivity between the dorsal ACC and bilateral caudate

Resting-state functional connectivity data was collected in the same subjects (n=833), and a seed analysis from the dorsal ACC (see Figure 2A) was conducted. This data revealed diminished functional connectivity between the dorsal ACC and the head of the both A) the right caudate and B) the left caudate. Images thresholded at z > 3.09, cluster corrected p < 0.01.
Table 1

Summary of demographic data.

|                           | M   | SD  |
|---------------------------|-----|-----|
| Age (years)               | 16.12 | 2.82 |
| N                         |     |     |
| Percent                   |     |     |
| Gender                    |     |     |
| Male                      | 468 | 45% |
| Female                    | 574 | 55% |
| Race                      |     |     |
| Caucasian                 | 473 | 45% |
| Non-Caucasian             | 569 | 55% |
| Maternal Level of Education |    |     |
| 12 years or less          | 395 | 38% |
| Greater than 12 years     | 634 | 61% |
| Missing                   | 13  | 1%  |
| Lifetime Prevalence *     |     |     |
| Typically Developing      | 284 | 27% |
| ADHD                      | 168 | 16% |
| Agoraphobia               | 72  | 7%  |
| Anorexia                  | 14  | 1%  |
| Bulimia                   | 4   | 3%  |
| Conduct Disorder          | 101 | 10% |
| Generalized Anxiety Disorder | 22 | 2%  |
| Major Depression          | 171 | 16% |
| Mania                     | 11  | 1%  |
| Obsessive-Compulsive Disorder | 36 | 3%  |
| Oppositional Defiant Disorder | 386 | 37% |
| Panic                     | 13  | 1%  |
| Psychosis-spectrum        | 351 | 34% |
| PTSD                      | 147 | 14% |
| Separation Anxiety        | 42  | 4%  |
| Social Phobia             | 275 | 26% |
| Specific Phobia           | 322 | 31% |

Note.

* Due to comorbidity, individual participants may be present in more than one category.