

Title: Altered Functioning of Reward Circuitry in Youth Offspring of Parents with Bipolar Disorder

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46 **Abstract**

47 **IMPORTANCE**

48 Offspring of parents with BD (BO) are at higher risk of bipolar disorder (BD) than offspring of
49 parents with non-BD psychopathology (NBO), although both groups are at higher risk than offspring of
50 psychiatrically healthy parents (HC) for other affective and psychiatric disorders. Abnormal
51 functioning in reward circuitry has been demonstrated previously in individuals with BD.

52 **OBJECTIVE**

53 To determine whether activation and functional connectivity in reward circuitry circuitry during
54 decision-making differentiated BO, NBO and HC.

55 **DESIGN, SETTING, AND PARTICIPANTS**

56 This study was conducted at the University of Pittsburgh Medical Center. BO (n=29;mean
57 age=13.8 years;14 female), NBO (n=28;mean age=13.9 years;12 female) and HC (n=23;mean
58 age=13.7 years;11 female) were scanned while performing a number guessing reward task. 11 BO and
59 12 NBO had current non-BD psychopathology; 5 BO and 4 NBO were taking psychotropic
60 medications.

61 **MAIN OUTCOMES AND MEASURES**

62 Region-of-interest analyses examined neural activation within prefrontal-ventral striatal reward
63 circuitry and functional connectivity between bilateral ventral striatum (VS) and right ventrolateral
64 prefrontal cortex during a reward paradigm (win, loss, and control conditions).

65 **RESULTS**

66 A 3(Group) x 2(Conditions:Win-Control/Loss-Control) ANOVA revealed a main effect of Group
67 on right frontal pole activation: BO showed significantly greater activation than HC. There was a
68 significant main effect of Group on functional connectivity between bilateral ventral striatum (VS) and
69 right ventrolateral prefrontal cortex (voxel $p<0.001$, cluster $p<0.05$): BO showed significantly greater

70 inverse functional connectivity than other participants. These between-group differences remained after
71 removing youth with psychiatric disorders and psychotropic medications from analyses.

72 CONCLUSIONS AND RELEVANCE

73 This is the first study to demonstrate that reward circuitry functioning distinguishes BO from
74 NBO and HC. The fact that the pattern of findings remained when comparing healthy BO vs. healthy
75 NBO vs. HC suggests that these neuroimaging measures may represent neurobiological markers
76 conferring either risk for, or protection against, BD in youth.

77

78 **Introduction**

79 It is well established that offspring of parents with bipolar disorder (BD) are at increased risk of
80 developing anxiety disorders and BD.¹⁻³ Furthermore, offspring of parents with BD (BO) are at higher
81 risk of BD than offspring of parents with non-BD psychopathology (NBO), although both groups are at
82 higher risk than offspring of psychiatrically healthy parents (HC) for other affective disorders.² Little is
83 known regarding the neurophysiological processes that predispose to, or protect from, risk for BD
84 versus risk for other affective and psychiatric disorders in these offspring, however, given that no study
85 has directly compared neurophysiological processes in BO, NBO and HC. Critically, studies examining
86 these processes have potential to help identify biomarkers denoting which at-risk offspring are most
87 likely to develop which specific psychiatric disorders in the future, and ultimately provide biological
88 targets to guide early interventions for these individuals. Neuroimaging studies are appropriate as a
89 way forward in this research field, as they can determine the extent to which BO and NBO show
90 abnormal functioning in neural circuitries supporting information processing domains known to be
91 aberrant in individuals with established BD. One such information processing domain is reward
92 processing, as an increasing number of studies reported abnormally heightened reward sensitivity in
93 individuals with BD.⁴⁻⁶

94 Reward circuitry comprises a complex, highly-interconnected network of fronto-subcortical
95 regions.^{7,8} The ventral striatum (VS) supports reward anticipation and prediction error,⁹⁻¹³ the pallidum
96 encodes expected reward value,¹⁴ the amygdala, stimulus-value associations,¹⁵ and the putamen, action-
97 specific value signals¹⁶ and effort costs.¹⁷ Different prefrontal cortical (PFC) regions contribute
98 differently to reward processing and decision-making.¹⁸ The ventromedial prefrontal (vmPFC) and the
99 orbitofrontal (OFC) cortices encode reward values and compare values of different options.¹⁹ The
100 ventrolateral PFC (vlPFC) encodes the value of choice/decision-making options and is important for
101 credit assignment.²⁰ The frontal polar (FP) region encodes the value of a non-chosen option during
102 decision-making,¹⁹ responds in situations of uncertainty,²¹ and maintains possible behavioral choices²²

103 and intentions²³ in memory for future use. The anterior cingulate cortex (ACC) is involved in cost-
104 benefit decision-making^{24,25} and action-reward associations.¹⁸ Neuroimaging studies report abnormal
105 functioning in reward circuitry in individuals with BD during reward and loss expectation and
106 processing, in particular, elevated activity in vmPFC, OFC, vlPFC, FP and striatum,²⁶⁻³² and reduced
107 inverse VS-vlPFC functional connectivity.³³

108 Many components of reward neural circuitry undergo developmental changes during
109 adolescence^{34,35} and have different maturation curves.^{36,37} For example, the later maturation of the
110 prefrontal cortex³⁷ may underlie the difficulty in regulating behaviors in potentially rewarding contexts,
111 and the high prevalence of risky behaviors, in adolescence.³⁸⁻⁴⁰ In parallel, studies report that youth
112 with BD, relative to healthy youth, show impaired reward learning^{41,41} and greater reward-related
113 arousal.^{43,44} Genetic risk for BD (and other affective disorders) may thus influence the course of
114 reward circuitry maturation, leading to heightened reward sensitivity, that ultimately predispose to
115 development of BD.

116 No study to date compared reward circuitry functioning in BO and NBO who have not as yet
117 developed BD, and HC, although one recent study showed elevated vlPFC activity and altered vlPFC-
118 ACC functional connectivity during reward processing in healthy BO vs. HC.⁴⁵ In the present study, we
119 thus aimed to identify the effect of familial genetic risk for BD (BO>NBO>HC) on activation and
120 functional connectivity in reward circuitry by comparing these neural measures in BO vs. NBO vs. HC
121 during a decision-making task.⁴⁶ Choice of regions in reward circuitry was determined from previous
122 neuroimaging findings.⁷⁻²⁵ Given that NBO, like BO, are at significantly higher risk for affective and
123 other psychiatric disorders than HC,² inclusion of NBO allowed us to control for risk for non-BD
124 disorders in BO. BO and NBO included youth with and without current non-BD psychopathology,
125 some of whom were treated with psychotropic medications. This allowed us to determine the effect of
126 genetic risk for BD on reward circuitry functioning in participants without current psychopathology
127 and psychotropic medication.

Based on the above neuroimaging findings in adults and youth with BD, we hypothesized the following:

1. Given previous findings showing abnormally elevated fronto-striatal activity in individuals with BD,²⁶⁻³² we hypothesized that BO (who are at highest risk for BD) would show abnormally elevated activity in this circuitry during reward processing, relative to NBO and HC.
2. Given a key role of the VS in reward processing,⁹⁻¹³ and previous findings showing altered functional connectivity between VS and anteroventral prefrontal cortex in individuals with BD versus HC,³³ we hypothesized that BO, but not NBO, would show significantly altered functional connectivity between these regions during reward processing compared with HC.
3. This differential pattern of reward circuitry functioning between BO and other participants would be present in participants without current psychiatric diagnoses and unmedicated participants.

Methods and Materials

The Bipolar Offspring Study (BIOS) is an ongoing longitudinal study examining psychiatric symptomatology in youth offspring of parents with BD² and functioning in neural circuitries underlying information processing domains implicated in the pathogenesis of BD, including reward circuitry. The study was approved by the Institutional Review Board of the University of Pittsburgh. Prior to study participation, parents/guardians provided written informed consent, and children provided written informed assent. Participants received monetary compensation for their participation.

Participants

Three groups of participants aged 7-17 years who were not affected with BD took part in this study: youth offspring of parent(s) with BD (BO;n=35), youth offspring of parent(s) with non-BD psychopathology (NBO;n=37) and psychiatrically healthy youth offspring of psychiatrically healthy

152 parents (HC;n=25) without family history of any psychiatric disorders (including first-degree relatives).
153 Twenty-four HC were recruited from the healthy comparison youth group of the Longitudinal
154 Assessment of Manic Symptoms (LAMS) study^{47,48} at the University of Pittsburgh Medical
155 Center/Western Psychiatric Institute and Clinic, a parallel study examining neural circuitry functioning
156 in youth with behavioral and emotional dysregulation. One HC was recruited from BIOS. Most BO and
157 NBO were recruited from BIOS,¹ with the exception of 2 BO and 5 NBO, who were recruited from
158 LAMS.^{47,48}

159 Exclusion criteria for all participants were: systemic medical illness, neurological disorders,
160 head trauma, alcohol or illicit substance use, standard exclusion criteria for MRI research (metal in the
161 head or body, claustrophobia, etc), IQ<70 (using the Weschler Abbreviated Scale of Intelligence⁴⁹),
162 unable to read and write in standard English, and corrected far visual acuity worse than 20/40 on the
163 Snellen visual acuity test. Six BO, 9 NBO, and 2 HC were excluded from data analysis due to inability
164 to complete the scanning session or due to excessive motion in the scanner (translation \geq 4mm in any
165 direction). The total numbers of participants with usable fMRI data were:29 BO, 28 NBO, and 23 HC.
166 Eleven BO and 12 NBO had current non-BD psychopathology. Five BO and 4 NBO were taking one
167 class of psychotropic medications (Table 1). Given ethical concerns with stopping medication for
168 research participation, participants were permitted to use prescribed medication(s) before, and on the
169 day of, scanning.

170 ***Assessment procedures***

171 Parental psychopathology was ascertained by a trained clinician using the Structural Clinical
172 Interview for DSM-IV (SCID-I)⁵⁰ for BIOS youth, and using detailed clinical assessment for LAMS
173 youth. Another trained clinician, blind to the parent's condition, interviewed the parents about their
174 children, and also interviewed their children, using the Kiddie Schedule for Affective Disorders and
175 Schizophrenia-Present and Lifetime version (KSADS-PL).⁵¹ All cases were supervised by a "blind"

176 child psychiatrist who was responsible for the ultimate parental and children diagnoses. Inter-rater
177 reliability for all psychiatric diagnoses ascertained through the KSADS>.8.

178 On the day of scan, parents/guardians of youth participants completed the PGBI-10M (Parent
179 Version, General Behavior Inventory,⁵² to assess the severity of behavioral and emotional dysregulation
180 in their offspring during the last 6 months; only parents of BO and NBO completed this questionnaire);
181 the SCARED-P (Self-Report for Childhood Anxiety Related Emotional Disorders, parent version, to
182 assess offspring anxiety over last 2 weeks);⁵³ the CALS-P (The Children's Affective Liability scale,
183 parent version);⁵⁴ the MFQ-P (Mood and Feelings Questionnaire, parent version, to assess the severity
184 of depression during the last 2 weeks);⁵⁵ and a questionnaire to assess sociodemographic status
185 represented by parental education.⁵⁶ Youth participants completed child report versions of affective
186 symptomatology scales: the CALS-C, SCARED-C, MFQ-C. All participants completed medication
187 forms that documented psychotropic medications used that day, during the last 24 hours, and those used
188 on regular basis; drug/alcohol/pregnancy screens; the Edinburgh Handedness Inventory (EHI);⁵⁷ and the
189 Snellen visual acuity test. Table 1 reports demographic and clinical variables. Table S1 reports
190 demographic and clinical variables for youth without psychopathology and youth untreated with
191 psychotropic medications. Table 2 reports lifetime psychiatric diagnoses in parents.

192 ***Reward task***

193 Participants were scanned while performing a number guessing reward task⁴⁶ (*Supplemental*
194 *Methods*, Figure 1) that activates fronto-striatal reward circuitry, and has been used previously in
195 neuroimaging studies of adults and youth with mood disorders.^{46,58}

196 ***fMRI data acquisition and analysis***

197 fMRI data were acquired using a Siemens MAGNETOM TrioTim 3T MR system. Acquisition
198 parameters, preprocessing and co-registration procedures are described in *Supplemental Methods*. fMRI
199 data were analyzed using FSL5.0.2(www.fmrib.ox.ac.uk/fsl). Preprocessed data were submitted to a

200 first-level GLM analysis implemented using FEAT (FMRI Expert Analysis Tool,v6.0). The model
201 included four regressors (Win, Loss, and Control trials, and Instructions). The magnitude of activation
202 was examined for each of these conditions and to the Win-Control, Loss-Control, and Win-Loss
203 contrasts. All group-level analyses were conducted using FLAME1 (FMRIB's Local Analysis of Mixed
204 Effects). Significant clusters of activation were determined by thresholding Z-statistic images in the
205 reward circuitry mask using voxel-wise $p < 0.001$ ($z > 3.09$) and a corrected cluster significance threshold
206 of $p < 0.05$.⁵⁹

207 **Hypothesis 1 testing**

208 *Activity in the reward circuitry ROI:*

209 Brain activation in the reward circuitry ROI was analyzed using a 3(Group:
210 BO/NBO/HC)x2(Condition:Win-Control/Loss-Control) ANOVA. The reward circuitry ROI mask was
211 the anatomical mask used in a previous study⁵⁸ that examined reward circuitry function in emotionally
212 dysregulated youth, using the same reward task. The mask included key neural regions implicated in
213 reward processing: bilateral dorsal ACC(BA24/32), medial and lateral FP(BA10), OFC(BA11),
214 vIPFC(BA47), and ventral striatum (VS; spherical ROIs with radius of 8mm centered [$\pm 9, 9, -8$]).

215 **Hypothesis 2 testing**

216 *Functional connectivity between bilateral VS and the reward circuitry mask:*

217 Functional connectivity was examined using psychophysiological interaction (PPI) analysis,⁶⁰
218 in FEAT. The bilateral VS served as a seed region and the reward circuitry mask served as a target
219 region. The PPI first-level analysis model included four psychological regressors (Win, Loss, and
220 Control trials, and Instructions), one physiological regressor—a mean time course extracted from the
221 seed region, and three interaction terms between the physiological and Win, Loss and Control
222 regressors. To parallel the activation analysis, the group-level connectivity analysis was conducted
223 using a 3(Group)x2(Condition) ANOVA.

224

225 **Post-hoc tests**

226 To determine the direction of the between- or within-group effects, post-hoc t-tests of activation
227 and connectivity values (parameter estimates extracted from the significant activation and connectivity
228 clusters) were conducted in R (<http://www.r-project.org/>) and Bonferroni corrected to control for
229 multiple t-tests.

230 **Hypothesis 3 testing**

231 Here, we examined the effect of diagnosis and medications on activation and connectivity in the
232 brain regions identified in the previous analyses. For this purpose, we first extracted activation and
233 connectivity values from the significant clusters. Then, we conducted two 3x2 ANOVAs, using SPSS,
234 on 1) participants without diagnoses; 2) unmedicated participants.

235

236 **Results**

237 **Behavioral data**

238 There were no between-group differences in decision-making reaction time across all trials or
239 across Reward and Loss trials separately.

240 **Neuroimaging**

241 **Hypothesis 1**

242 *Activation*

243 A 3(Group:BO/NBO/HC)x2(Condition:Win-Control/Loss-Control) ANOVA revealed main
244 effects of Group (Figure 2A) and Condition (Table S2), but no Group x Condition interaction, on brain
245 activation. A main effect of Group was found in the right frontal pole (RFP; $n_{vox}=66$, $z_{max}=4.0$,
246 $[24,64,6]$). Follow up t-tests conducted on RFP activation values revealed that activation was
247 significantly greater in BO than HC ($t(50)=3.7$, $p<0.001$), and just missed significance in BO vs. NBO
248 ($t(55)=2.3$, $p=0.02$), using a Bonferroni-corrected statistical threshold of $p=0.05/3$ for between group

249 tests $p=0.017$.

250 **Hypothesis 2**

251 ***Connectivity***

252 The PPI analyses conducted with the bilateral VS as a seed region and the reward circuitry mask
253 as a target region revealed a main effect of Group (Figure 2B), but no main effect of Condition or
254 Group x Condition interaction, on functional connectivity. A main effect of Group was found in the
255 right vLPFC($n_{vox}=96$, $z_{max}=4.7$, $[40,46,-10]$). Follow up t-tests revealed that functional connectivity
256 between bilateral VS and right vLPFC was significantly more negative in BO than NBO ($t(55)=-3.3$,
257 $p=0.002$) and in BO than HC ($t(50)=-6.2$, $p<0.001$), using a Bonferroni-corrected threshold of $p=0.017$,
258 as above.

259

260 **Hypothesis 3**

261 ***Activation and connectivity***

262 The results of RFP activation and bilateral VS-right vLPFC connectivity analyses in
263 unmedicated participants and those without psychopathology were consistent with the results of the
264 full-sample analyses testing Hypotheses 1-2. There was a significant effect of Group on RFP activation
265 (participants without psychopathology: $F(2,54)=11.1$, $p<0.001$; unmedicated participants: $F(2,68)=7.3$,
266 $p=0.001$; Figure S1), and on bilateral VS-right vLPFC functional connectivity (participants without
267 psychopathology: $F(2,54)=14.4$, $p<0.001$; unmedicated participants: $F(2,68)=16.1$, $p<0.001$; Figure
268 S2). The results of the post-hoc comparisons have paralleled main findings from *Hypotheses 1-2*, and
269 are in *Supplemental Results*.

270

271 **Exploratory analyses**

272 Across all participants, RFP activation positively correlated with CALS-P ($r=0.34$, $p=0.002$)
273 and MFQ-P ($r=0.23$, $p=0.047$). In BO, RFP activation positively correlated with CALS-P ($r=0.37$,

274 $p<0.05$).

275 **Discussion**

276 The goal of the present study was to determine the extent to which alterations in reward
277 circuitry function characterized at-risk youth offspring of parents with BD (BO) relative to offspring of
278 non-bipolar parents (NBO) and healthy youth offspring of psychiatrically healthy parents (HC). Main
279 findings supported all three hypotheses, that activation and functional connectivity in the reward
280 circuitry are associated with genetic risk for BD. RFP activation for decision (Win and Loss) trials,
281 relative to non-decision (Control) trials, was significantly greater in BO than in HC. In contrast,
282 bilateral VS-right vIPFC functional connectivity was significantly more inverse in BO than in NBO and
283 HC. These patterns of activation and functional connectivity remained for unmedicated participants and
284 those without psychopathology, supporting our third hypothesis.

285 The FP is involved in decision-making and prospective memory by supporting the maintenance
286 of delayed intentions and representations^{22,23} and integrating outcomes of previous trials⁶¹ for potential
287 use in future trials. The magnitude of FP activation positively correlated with amount of uncertainty
288 remaining between multiple choices,²¹ and tracked unchosen options.¹⁹ Abnormally elevated RFP
289 activation during decision-making trials in BO suggests that they may have experienced abnormal
290 levels of uncertainty during these trials, and maintained non-chosen option-outcome contingencies in
291 memory to predict (i.e., make more certain) response-outcome mapping for future trials. Furthermore,
292 similar patterns of significantly elevated RFP activation during decision-making trials were present in
293 unmedicated BO and NBO vs. HC, and in BO and NBO without psychopathology vs. HC. A recent
294 study demonstrated a similar pattern of significantly greater activation in the right frontal cortex
295 [$x=11, y=55, z=14$] during reward anticipation in adolescents with BD vs. HC.³¹ Taken together, these
296 findings suggest that abnormally elevated RFP activation may represent a vulnerability marker for
297 future development of affective disorders.

298 The VS supports reward anticipation, reward evaluation and prediction error.⁹⁻¹³ The vIPFC is

299 implicated in learning the value of different options and formation of associations between visual
300 stimuli and reward values.¹⁸ Functional interaction between prefrontal cortex and VS influences guided
301 behavior and may be modulated by the environment.⁶² In HC, an increase in bilateral VS activation was
302 associated with increase in right vIPFC activation during decision-making vs. non-decision-making
303 trials, highlighting a positive relationship between encoding stimulus-outcome associations and reward
304 evaluation. In BO, an inverse relationship between these regions suggests that learning stimulus-
305 outcome associations may impede reward evaluation, and vice versa.

306 Given that BO not only differed from HC, but also differed from NBO on the direction of VS-
307 vIPFC functional connectivity, and that this pattern remained significant even for unmedicated
308 participants and those without psychopathology, inverse bilateral VS-right vIPFC functional
309 connectivity during decision-making trials may reflect a vulnerability marker for BD, rather than for
310 affective disorders in general. Because the between-group differences in functional connectivity were
311 independent of the fact that BO, NBO and HC did not differ significantly in magnitude of activation in
312 these regions, our findings may provide further support for dysfunctional coupling between the vIPFC
313 and subcortical regions during emotionally-salient processing as a pathophysiological process
314 underlying vulnerability to BD.^{63,64}

315 Exploratory analyses showed that greater RFP activation was associated with higher mood
316 dysregulation scores (CALS-P) and higher MFQ across all participants, and with higher CALS-P in
317 BO. Higher RFP activation during decision-making trials may thus be a precursor for development of
318 mood dysregulation and depression. Future studies need to examine these exploratory findings.

319 The fact that findings remained significant even after approximately 40% of participants were
320 removed for comparisons of healthy BO vs. healthy NBO vs. HC suggests that the pattern of findings
321 was robust, at least with regard to the general BO and NBO populations. Given that only a small
322 number of youth were taking medications, there was insufficient statistical power to directly compare
323 unmedicated BO vs. medicated BO vs. unmedicated NBO vs. medicated NBO. Further studies should

compare larger samples of medicated and unmedicated BO and NBO, and BO and NBO with and without current diagnoses. Additionally, future studies can also directly compare youth with established BD, and genetically and symptomatically at-risk youth, to determine the degree of similarity between neural measures of risk status and neural measures of BD.

In summary, our findings demonstrate, for the first time, that youth offspring, as yet unaffected with BD, of parents with BD exhibit altered patterns of frontal activation and vIPFC-striatal functional connectivity, that distinguish these youth from youth offspring of parents unaffected with BD. These activation and connectivity differences remained significant for participants without current psychopathology and medication history and may represent neurobiological markers conferring either risk for, or protection against, BD in youth. Future, longitudinal follow-up studies in youth at-risk for BD should aim to distinguish between these two possibilities, by determining the extent to which abnormal patterns of reward circuitry functioning predict, or protect against, development of BD, and development/worsening of affective pathology in general.

353

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564 **Figure Captions**

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568 **Figure 1.** An example of a Win trial in the reward task. A green arrow shows that a subject correctly
569 guessed that the number was greater than 5.

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571 **Figure 2.** A main effect of Group on activation in the right frontal pole (RFP [24 64 6]; shown in red)
572 and functional connectivity between the bilateral VS (centered 8 mm around [\pm 9 9 -8]; shown in
573 green) and the right vIPFC ([40 46 -10]; shown in blue) for decision-making trials (i.e., Win and
574 Loss trials) vs. non-decision-making trials (i.e., Control trials). The reward circuitry ROI mask is
575 shown in yellow. “au” stands for arbitrary units. “*” indicates significant t-test results. BO – offspring
576 of parents with bipolar disorder, NBO – offspring of parents with psychiatric disorders other than
577 bipolar disorder, HC – healthy offspring of psychiatrically healthy parents.

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588 **Tables**

589 Table 1

590 Demographic and clinical variables

	BO n=29	NBO n=28	HC n=23	Statistics	p-value
Number of youth without psychiatric diagnoses	18 (62%)	16 (57%)	23 (100%)	BO vs. NBO $\chi^2(2)<1$	ns
Number of youth untreated with psychotropic medications	24(83%)	24(86%)	23(100%)	BO vs. NBO $\chi^2(2)<1$	ns
Age at scan	13.81(2.45)	13.93(2.38)	13.74(1.80)	F(2,77)<1	ns
Gender (female)	14	12	11	$\chi^2(2)<1$	ns
Handedness (right hand)	26	26	21	Yates' $\chi^2(2)<1$	ns
IQ (WASI)	103.21(14.51)	102.86(14.33)	105.78(13.79)	F(2,77)<1	ns
SES based on parental education	5.48(0.95)	5.54(0.96)	5.30(1.02)	F(2,77)<1	ns
Medications					
antidepressants	na	sertraline HCl: n=1	na		
antipsychotics	Risperidone: n=1 Quetiapine Fumarate: n=1	na	na		
mood stabilizers	na	na	na		
stimulants	amphetamine, dextroamphetamine mixed salts: n=1	methylphenidate: n=1 amphetamine, dextroamphetamine mixed salts: n=2	na		
non-stimulants	Atomoxetine HCl: n=2	na	na		
benzodiazepines	na	na	na		
Youth offspring current psychiatric diagnoses					
Number of youth with more than 1 diagnosis	6	7	na		
MDD/DDNOS	3	2	na		
Attention deficit	6	5	na		

hyperactivity disorder

Anxiety disorders	2	2	na
Oppositional Defiant Disorder	1	2	na
Phobias	2	2	na
Tourette's Disorder	1	0	na
Obsessive compulsive disorder	0	2	na
Eating disorder	2	0	na

Symptom Assessment Scale Scores administered on the day of scan

SCARED Parent Total	9.45(6.86)	10.85(11.70)	4.17(4.32)	F(2,76)=4.35	0.02
SCARED Child Total	11.66(8.61)	10.79(13.80)	9.33(11.42)	F(2,77)<1	ns
MFQ Parent	5.90(8.97)	5.42(9.09)	1.57(2.09)	F(2,75)=2.4	ns
MFQ Child	8.86(10.73)	10.18(10.97)	5.09(10.57)	F(2,77)=1.5	ns
CALS Parent Total	7.97(10.26)	5.07(7.65)	1.78(2.59)	F(2,76)=4.04	0.02
CALS Child Total	10.52(12.22)	8.79(11.28)	5.96(13.39)	F(2,77)<1	ns

591 *Note:* Standard deviations (SD) are reported in parentheses.

592 Table 2

593 Lifetime psychiatric diagnoses in parents

	BO n=29	NBO n=28	Statistics	p-value
BD-I	23	0	$\chi^2(1)=37.2$	p<0.001
BD-II	6	0	$\chi^2(1)=6.5$	p=0.01
BD-NOS	0	0		
Major depressive disorder/Depressive disorder NOS	1	20	$\chi^2(1)=28.3$	p<0.001
Generalized Anxiety Disorder/Anxiety disorders NOS	16	8	$\chi^2(1)=4.1$	p=0.04
Phobias	21	14	$\chi^2(1)<3.0$	ns

	BO n=29	NBO n=28	Statistics	p-value
Alcohol/Drug abuse/dependence	23	13	$\chi^2(1)=6.6$	p=0.01
Post-traumatic stress disorder	12	4	$\chi^2(1)=5.2$	p=0.02
Panic disorder	16	6	$\chi^2(1)=6.8$	p<0.01
Eating disorder	4	1	$\chi^2(1)=1.8$	ns
Obsessive-compulsive disorder	10	0	$\chi^2(1)=11.7$	p<0.001
Attention deficit hyperactivity disorder	4	2	$\chi^2(1)<1$	ns

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**Guess
Number**

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+

2000 ms
Instructions

3000 ms
Subject response
(e.g., >5)

500 ms
Feedback

500 ms
Feedback and
subject response

3000 ms
ITI

