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‘The risks of playing it safe’: a prospective longitudinal study of response to reward in the adolescent offspring of depressed parents

A. Rawal1*, S. Collishaw2, A. Thapar2 and F. Rice1*

1 Department of Clinical, Educational and Health Psychology, University College London, UK
2 Child and Adolescent Psychiatry Section, Department of Psychological Medicine and Neurology, Cardiff University, MRC Centre for Neuropsychiatric Genetics and Genomics, Neuroscience and Mental Health Research Institute, UK

Background. Alterations in reward processing may represent an early vulnerability factor for the development of depressive disorder. Depression in adults is associated with reward hyposensitivity and diminished reward seeking may also be a feature of depression in children and adolescents. We examined the role of reward responding in predicting depressive symptoms, functional impairment and new-onset depressive disorder over time in the adolescent offspring of depressed parents. In addition, we examined group differences in reward responding between currently depressed adolescents, psychiatric and healthy controls, and also cross-sectional associations between reward responding and measures of positive social/environmental functioning.

Method. We conducted a 1-year longitudinal study of adolescents at familial risk for depression (n=197; age range 10–18 years). Reward responding and self-reported social/environmental functioning were assessed at baseline. Clinical interviews determined diagnostic status at baseline and at follow-up. Reports of depressive symptoms and functional impairment were also obtained.

Results. Low reward seeking predicted depressive symptoms and new-onset depressive disorder at the 1-year follow-up in individuals free from depressive disorder at baseline, independently of baseline depressive symptoms. Reduced reward seeking also predicted functional impairment. Adolescents with current depressive disorder were less reward seeking (i.e. bet less at favourable odds) than adolescents free from psychopathology and those with externalizing disorders. Reward seeking showed positive associations with social and environmental functioning (extra-curricular activities, humour, friendships) and was negatively associated with anhedonia. There were no group differences in impulsivity, decision making or psychomotor slowing.

Conclusions. Reward seeking predicts depression severity and onset in adolescents at elevated risk of depression. Adaptive reward responses may be amenable to change through modification of existing preventive psychological interventions.

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Key words: Adolescence, anhedonia depression, impairment, impulsivity, positive affect, prevention, reward, risk, social withdrawal.

Introduction

Low positive affect is a hallmark of depressive disorders and is reflected in key symptoms such as anhedonia, social withdrawal and reduced activity level. Alterations in reward processing may be an important mechanism underlying such disturbances (Naranjo et al. 2001; Eshel & Roiser, 2010). Lowered reward responsiveness may lead to diminished engagement in pleasurable activities and reduced motivation to pursue rewarding outcomes such as social events, sports and interpersonal relationships (Depue & Iacono, 1989; Forbes & Dahl, 2005), suggesting that reward responsiveness may play an important role in the onset and maintenance of depression. Indeed, impaired reward processing has been postulated as a behavioural endophenotype in depression (Hasler et al. 2004, 2009).

Brain structures involved in reward processing, such as the orbitofrontal cortex, amygdala, ventral striatum and medial prefrontal cortex (McClure et al. 2004), function abnormally in depressed adults when anticipating and gaining monetary reward.
(Keedwell et al. 2005; Steele et al. 2007; Pizzagalli et al. 2009; Smoski et al. 2009). On a behavioural level, depressed adults show impairments in changing responses as a function of reward (Henriques & Davidson, 2000; Pizzagalli et al. 2005). Depressed individuals therefore appear hypersensitive to reward and may not develop preferences for behaviours associated with greater reward.

Although lack of reinforcing behaviour is associated with current depression, it is possible that this association may differ for impending depression. A major research aim is therefore to understand whether reward processing influences early vulnerability for depression. Psychiatric disorders frequently begin in adolescence, the incidence of depression is highest during this period and adolescent depression shows substantial continuity over time (Weissman et al. 2006). Early life has been considered a crucial period for the organization of affective systems (Nelson et al. 2009). The reward system undergoes substantial development in adolescence, with an increased sensitivity to, and seeking of, reward (Davey et al. 2008; Forbes & Dahl, 2012). Understanding reward-related aberrations during this period may have important implications for the development of early vulnerability towards depression.

Only a few studies have examined reward processing in children and adolescents with depression. These studies have revealed attenuated neural activity in reward-related brain regions during reward anticipation and outcome compared to healthy controls (Forbes et al. 2006, 2009) and found that reward-related activity correlated with positive affect in natural environments (Forbes et al. 2009). The two behavioural studies of depressed adolescents to date have shown conflicting findings. In a male sample, Forbes et al. (2007) found that recently depressed boys failed to differentiate between small and large monetary rewards during high-probability reward conditions, thus showing behaviour reflecting diminished reward seeking. By contrast, on a gambling task that involved staking bets on one of two outcomes of varying probability, Kyte et al. (2005) found no difference between depressed adolescents and controls at highly probably outcomes. However, at less probable reward outcomes, depressed adolescents bet more than controls, indicating a less conservative reward-seeking strategy.

Group differences in reward processing do not indicate that the association between reward processing and depression is causal. However, there is some evidence that deficits in reward responding could confer vulnerability to depression (McCabe et al. 2009). Gotlib et al. (2010) found attenuated neural activity during reward processing in adolescent girls free from psychopathology but at familial risk for depression compared to healthy controls. However, they did not examine the relationship between reward processing and subsequent depression. Forbes et al. (2007) showed that choices during trials where both magnitude and probability of reward were high predicted depressive symptoms and the occurrence of depressive and anxiety disorders at 1-year follow-up. These initial findings highlight the role of reward processing as a potential vulnerability factor for adolescent depression.

Parental depression is the most robust risk factor for depression in young people, with around 40% of this group developing depressive disorder by early adulthood (Rice et al. 2002; Weissman et al. 2006). Studying adolescents at elevated risk for depression in a prospective research design provides an opportunity to examine whether behavioural alterations in reward responding are present before the onset of depression and could therefore potentially be targeted in preventive interventions (Gotlib et al. 2010). Moreover, given the heterogeneity in outcome in offspring of depressed parents, this design also allows for a better characterization of risk.

We examined the role of reward processing in a 1-year longitudinal study of adolescents at risk for depression due to a parental history of depression. Specifically, we examined two aspects of reward responding: (1) reward seeking, measured by betting behaviour under a variety of odds when the more likely of two outcomes was chosen, and (2) risk adjustment, measured by the extent to which variation in odds affected betting. In addition to examining differences in overall levels of reward seeking, we were particularly interested in the relationship between depression and reward seeking at highly favourable reward conditions (i.e. when likelihood of reward is high) on the basis of Forbes et al.’s (2007) findings. Adolescents were assessed for psychiatric disorder at baseline and follow-up and only a proportion of the cohort had a current psychiatric disorder during the study. This enabled prospective examination of the role of reward responding in depression in those adolescents without a prior depressive episode. First, we examined reward responding in adolescents with depressive disorder, and in those with no disorder or other psychiatric disorders (externalizing and anxiety disorders). This allowed us to identify whether a particular pattern of reward responding was specific to depression rather than simply a marker of current psychopathology or a general feature in offspring of depressed parents. We expected diminished reward seeking to be characteristic of depression, as externalizing disorders may involve increased reward seeking (Scheres et al. 2007; Gatzke-Kopp et al. 2009) and anhedonia is thought to be less typical of anxiety.
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Method

Participants

The current study was part of an ongoing longitudinal study of parents with recurrent unipolar depression and their biological adolescent offspring: the Early Prediction of Adolescent Depression (EPAD) study (Mars et al. 2012). A history of recurrent depression in the parent was verified using the Schedules for Clinical Assessment in Neuropsychiatry (SCAN; Wing et al. 1990). Exclusion criteria were a diagnosis of bipolar disorder or a history of mania in the index parent, adolescent not living at home, or adolescent IQ <50. There were no diagnostic exclusion criteria for adolescents. One eligible adolescent per household participated. Parents were recruited from primary care in South Wales, UK (78%), from a previous community study of recurrent unipolar depression (19%), and from advertisements in primary care (3%).

Psychopathology data were available at baseline (when adolescents completed the reward task) and at follow-up (average = 12.5 months). Full psychopathology data were available for 277 adolescents at baseline and 251 adolescents at follow-up, 216 of whom also had reward task data. Non-completion of the task was due to: shortage of equipment (28), time limitations (17), participant refusal (7), other reasons (9), for example a fractured arm. For 19 participants, computer failure caused a loss of reward task data. Thus, 197 participants had complete reward task data and these did not differ on key study variables from those for whom reward data were unavailable: age (t = 0.97), gender (χ² = 0.64); depressive symptoms (t = 0.45), rates of depressive disorders (χ² = 0.23), anxiety disorders (χ² = 0.34), disruptive disorders (χ² = 0.29), or attention deficit hyperactivity disorder (ADHD; χ² = 0.31); all p values > 0.33.

Measures

Psychiatric symptoms and disorder

Adolescent psychiatric symptoms and disorders (depressive disorders, anxiety disorders, eating disorders, conduct disorder, oppositional defiant disorder, ADHD, bipolar disorder and psychosis) were assessed on two occasions using the Child and Adolescent Psychiatric Assessment (CAPA; Angold et al. 1995). The CAPA is a semi-structured interview that provides a detailed assessment of adolescent psychopathology over the preceding 3 months. Interviews were conducted separately with the parent and adolescent. Inter-rater reliability was excellent (κ = 0.9 for adolescent depression). All cases meeting DSM-IV diagnostic criteria and subthreshold cases were reviewed by two child psychiatrists and diagnoses were agreed by clinical consensus. A disorder was considered to be present if a diagnosis was made based on interview of either the parent or the adolescent (Angold & Costello, 1995).

The severity of depressive symptomatology over the preceding 3 months was assessed with the 34-item version of the Mood and Feelings Questionnaire (MFQ; Costello & Angold, 1988) at baseline and follow-up (score range: 0–68). The MFQ correlates highly with other measures of depressive symptoms and clinical interviews of depression (Angold et al. 1995). Parents and adolescents completed the MFQ. If either informant endorsed a symptom it was counted as present. Evidence indicates that parents and adolescents offer complementary information (Costello & Angold, 1988) and that combining child and parent ratings improves sensitivity in detecting depressive mood compared to the use of either score alone (Angold et al. 1995; Daviss et al. 2006). The MFQ showed high internal reliability (Cronbach’s α = 0.96 at baseline and 0.95 at follow-up). An anhedonia score was calculated from items regarding loss of pleasure, loss of interest and loss of energy.

Functional impairment

Functional impairment was assessed at baseline and follow-up with the impact supplement of the Strengths and Difficulties Questionnaire (SDQ; Goodman, 1999). Parent and adolescent reports were combined, whereby if either informant endorsed a problem it was counted as present. The SDQ indexes the extent to which emotional and behavioural difficulties cause distress and social impairment and
predicts psychiatric service use (Goodman, 1999; Ford et al. 2008). Cronbach’s α = 0.78 at baseline and 0.77 at follow-up.

Peer relationship quality
This was assessed at baseline using 10 items that assess friendship quality (e.g. ‘Children in my class are friendly to me’). Cronbach’s α = 0.87. This measure was devised for the study and was negatively correlated with the SDQ peer problems scale (r = -0.69), indicating convergent validity.

Extra-curricular activities
A four-item checklist was used to assess frequency of exercise, sport and participation in clubs, groups or classes at baseline. Cronbach’s α = 0.63.

Humour
Humour plays an important role in social interaction (Berns, 2004). The Multidimensional Sense of Humour Scale (Dowling et al. 2003) assessed humour appreciation and creation (e.g. ‘I like a good joke’, ‘I can make other people laugh’) at baseline. Cronbach’s α = 0.95.

Pubertal status
We assessed pubertal status at baseline using a self-report questionnaire (Petersen et al. 1988) that shows good validity in comparison to physician ratings (Brooks-Gunn et al. 1987). Adolescents rated the extent to which their bodies had changed (from ‘not at all’ to ‘a lot’) on indices of pubertal development (e.g. height, facial and body hair) and reported whether each of these aspects of pubertal development was completed [e.g. ‘Are you as tall as an adult (have you finished growing)? ’]. Participants indicating no change and no completion were defined as pre-pubertal, those indicating some change/completion as pubertal, and those indicating completed development on all indices as post-pubertal.

Full-scale IQ (FSIQ)
We assessed FSIQ on one occasion using 10 subscales of the Wechsler Intelligence Scale for Children – Fourth Edition (WISC-IV; Wechsler, 2004).

Reward task
We used the Cambridge Gambling Task (CGT), a well-characterized reward task associated with neural substrates of reward processing (Clark et al. 2008; CANTAB, www.camcog.com). On each trial, 10 coloured boxes (blue or red) are presented on screen and the ratio of blue to red boxes varies from 9:1 to 1:9, in pseudo-random order. In total, five possible probabilities occur in the task (9:1, 8:2, 7:3, 6:4, 5:5). Initially, the participant must decide under which colour (blue or red) a token has been hidden (Fig. 1, left; the numbers of red and blue boxes reflect the probability that the token is associated with a particular colour). This yields two indices of decision making: the proportion of times the more likely outcome is chosen (quality of decision making) and deliberation time. In the second phase of each trial, the participant must bet a proportion of their points on the chosen colour. Possible bets of varying magnitude are offered in a sequence (5, 25, 50, 75, 95% of points), in 2.5-s increments. In half the blocks, bets are presented in ascending order, in the other half in descending order (the order of condition was counterbalanced across participants). Subtraction of bets on ascending trials from descending trials measures impulsivity (indexed by low bets in the ascend condition coupled with high bets in the descend condition). Participants place their bet by touching an answer box on the screen (Fig. 1, right). The hidden token’s location is subsequently revealed. The amount of the bet is then added to (if correct) or subtracted...
from (if incorrect) the total score. This second phase of the task yields two measures of reward responding: (1) reward seeking, which is conceptualized as motivation to risk already accumulated points to acquire further reward (measured by the proportion of points gambled on trials where the more likely outcome is selected); and (2) risk adjustment, which assesses the linear effect of probability on betting behaviour. As the ratio of blue to red boxes varies, this measures the extent to which participants adjust reward-seeking behaviour to changing context. Risk adjustment is calculated as: 

\[ \frac{2a + b - c - 2d}{\text{average bet}} \]

where \( a \) represents the mean bet in the 9:1 ratio, \( b \) represents the mean bet at the 8:2 ratio, and so on (Clark et al. 2011).

Participants began the task with 100 points. They were told: ‘The idea is to build up as many points as you can. Try not to let your score get as low as 1 point because then you will lose the game.’ Participants completed four practice trials, followed by eight blocks of nine trials. At the start of each block, the total was reset to 100 points. Analysis of betting behaviour was limited to trials where the more likely outcome was selected (i.e. the colour in the majority) to maintain independence of betting behaviour and decision making (Clark et al. 2008). Trials where the ratio of boxes was equal (5:5) were included in the task but excluded from analysis.

**Procedure**

Assessments were conducted in families’ homes. Parents and adolescents aged >16 years provided written informed consent, younger participants provided written assent. Ethical review and approval were provided by the Multi-Centre Research Ethics Committee for Wales.

**Statistical analysis**

Reward task data were transformed to approximate normality (latency data logarithmically, proportion data arcsine transformed; Howell, 1997). Data presented in the text and figures correspond to untransformed means. Repeated-measures analysis of variance (ANOVA) was used to examine diagnostic group differences (no disorder, depressed, anxiety and externalizing) on reward measures. The ratio of coloured boxes (9:1, 8:2, 7:3 and 6:4) and the condition that bets were presented in (ascending or descending) were within-subject factors. Pearson’s \( r \) and linear regression were used to examine associations between continuous variables. Logistic regression was used to examine reward responding as a predictor of new-onset depression. The main predictor variables were overall reward seeking and reward seeking at high probability ratios (9:1 and 8:2).

**Descriptive characteristics**

At baseline, participants were classified as having a depressive disorder (\( n = 19 \)) if they received a diagnosis of major depressive disorder, dysthymia, minor depression, or depression not otherwise specified. Minor depression (\( n = 2 \)) was defined as 2 weeks of low mood in addition to one other symptom and associated incapacity. Participants were classified as having an anxiety disorder (\( n = 15 \)) if they received a diagnosis of generalized anxiety disorder, separation anxiety, social phobia, panic disorder, agoraphobia, or obsessive-compulsive disorder (but no diagnosis of depression). Externalizing disorders (\( n = 24 \)) included diagnoses of oppositional defiant disorder, conduct disorder, disruptive disorder or ADHD (but no diagnosis of depression). Adolescents were assigned to the ‘no disorder’ group if they were free from psychopathology (\( n = 136 \)). Three participants had other psychiatric disorders (eating disorders, adjustment disorder) and were excluded from analyses. The final sample consisted of 194 adolescents [108 females, 86 males; mean age = 13.63 years, S.D. = 2.06, range 10–18; mean IQ = 97.29, S.D. = 12.13, range 69 (\( n = 1 \)) to 131 (\( n = 1 \)) at baseline, of whom 187 (96%) provided psychopathology data at follow-up. Table 1 presents demographic characteristics according to diagnostic status at baseline.

**Results**

**Preliminary analyses**

Repeated-measures ANOVAs showed no group differences in deliberation time, quality of decision making or delay aversion/impulsivity on the CGT (Table 2). There were within-subject effects of ratio on deliberation time, quality of decision making and delay aversion (\( F's > 3.78, p's < 0.02 \)), showing that participants deliberated the least at 9:1, chose the more likely outcome more often at higher probabilities, and were less impulsive at higher probabilities. This did not differ by group.

**Reward responding and depressive disorder**

Overall reward seeking differed by diagnostic status (\( F_{3,190} = 4.44, p = 0.01 \)). The depressive disorder group bet less than the no-disorder and the externalizing group. The externalizing group bet more than the anxiety group (Table 2). These effects were qualified by a group × ratio interaction (\( F_{9,114} = 2.13, p = 0.03 \)). Follow-up univariate analyses showed that the
depressive disorder group was less reward seeking than the no-disorder group and the externalizing group at the high probability ratios of 9:1 and 8:2 ($F_{3,190} > 4.25, p < 0.01$; Fig. 2). At 8:2, the difference between the depressive and anxiety group was significant at trend level ($p = 0.06$). At 7:3, the depressive disorder group bet significantly less than the externalizing group ($p = 0.03$) and the no-disorder group at trend level ($p = 0.07$). The interaction between group and ratio remained significant when depressive symptoms and antisocial behaviour symptoms were included as covariates and when cases with ADHD were excluded. These results were not influenced by condition (ascending or descending), pubertal status or gender ($F's < 1.57, p's > 0.11$). Eight depressed adolescents had a co-morbid anxiety disorder. The pattern of results was the same when these were excluded from analysis (data available from A.R.). There were no group differences in risk adjustment ($F_{3,190} = 52, p = 0.67$), as indicated by the gradient of reward seeking across probability ratios (Fig. 2).

**Reward responding and environmental/social functioning**

Table 3 shows correlations between reward and environmental/social functioning measures. Overall reward seeking was correlated with humour and friendship quality. Both reward seeking at 9:1 and risk adjustment correlated with engagement in extracurricular activities. Reward seeking at 8:2 also correlated with humour scores. Humour and friendship quality were substantially correlated, whereas extracurricular activities showed a small correlation with humour and were not correlated with friendship quality. Associations between reward responding and environmental/social functioning were next examined using multiple regression to adjust for covariation between variables. Both reward seeking at 9:1 ($\beta = 0.15, p = 0.05$) and risk adjustment ($\beta = 0.17, p = 0.02$) were still associated with extra-curricular activities. Reward seeking at 8:2 was associated with humour at trend level ($\beta = 0.11, p = 0.09$). Overall reward seeking was no longer associated with humour and friendship ($\beta's < 0.10, p's > 0.15$) when adjusting for correlated social functioning variables. Anhedonia was negatively correlated with reward seeking and measures of environmental/social functioning (Table 3). Measures of environmental/social functioning were negatively correlated with depression severity ($r's < -0.19, p's < 0.01$).

**Reward responding and impending depressive symptoms, functional impairment and new-onset depressive disorder**

Only adolescents free from depressive disorder at baseline were included in the analyses. Both overall reward seeking ($R^2 = 0.04, \beta = -0.22, p = 0.01$) and reward seeking at 9:1 ($R^2 = 0.05, \beta = -0.23, p < 0.01$) were associated with severity of depressive symptoms at follow-up in adolescents free from depressive disorder at baseline. Depressive symptoms at baseline and follow-up were significantly correlated ($r = 0.66, p < 0.001$). When controlling for baseline depressive
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Table 2. Full-scale IQ, psychopathology symptoms and Cambridge Gambling Task (CGT) characteristics by diagnostic status

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No disorder (n = 136)</th>
<th>Depressive disorder (n = 19)</th>
<th>Group comparison</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anxiety disorder (no depression)</strong></td>
<td>98.48</td>
<td>97.27</td>
<td>&lt;0.10</td>
</tr>
<tr>
<td><strong>Externalizing disorder (no depression)</strong></td>
<td>12.19</td>
<td>11.00</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Depressive disorder</strong></td>
<td>94.88</td>
<td>92.37</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>CGT reward seeking</strong></td>
<td>12.61</td>
<td>11.70</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Reward seeking (%)</strong></td>
<td>12.2</td>
<td>11.35</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Deliberation time (ms)</strong></td>
<td>59.22</td>
<td>50.23</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Quality of decision making (%)</strong></td>
<td>2519.36</td>
<td>2292.12</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Delay aversion (%)</strong></td>
<td>91</td>
<td>84.3</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

**Notes:** Standard deviation is presented. Reward seeking represents the proportion of points bet on trials where the more likely outcome was chosen. Risk adjustment reflects the tendency to stake higher bets on favorable compared to unfavorable trials. Antisocial behavior symptoms were derived from the conduct disorder and oppositional defiant disorder sections of the Child and Adolescent Psychiatric Assessment (CAPA). Severity of depressive symptoms and pubertal development, both overall reward seeking (ΔR² = 0.03, β = −0.17, p = 0.01) and reward seeking at 8:2 (ΔR² = 0.02, β = −0.15, p = 0.02) were still associated with depressive symptoms at follow-up. Reward seeking at 8:2 was also associated with depressive symptoms at follow-up (ΔR² = 0.02, β = −0.13, p = 0.03). There were no significant interactions between reward seeking and pubertal development (p’s > 0.24; however, small cell sizes limited analysis). Additionally, both overall reward seeking (ΔR² = 0.02, β = −0.14, p = 0.02) and reward seeking at 9:1 (ΔR² = 0.02, β = −0.15, p = 0.01) were associated with functional impairment at follow-up after controlling for baseline functional impairment. Risk adjustment was not associated with depressive symptoms or functional impairment (β’s < 0.11, p’s > 0.09).

Given the small number of cases with new-onset depression at follow-up (n = 4, all female), secondary analysis examined whether reward seeking was associated with new-onset depressive disorder. Baseline reward seeking at ratios 9:1 (Nagelkerke R² = 0.15, B = −0.48, s.e. = 0.23, p = 0.03) and 8:2 (Nagelkerke R² = 0.15, B = −0.60, s.e. = 0.28, p = 0.03) were associated with new-onset depressive disorder at follow-up. Thus, adolescents with new-onset depression at 9:1 and 8:2 (mean = 0.50, S.D. = 0.13, v. mean = 0.71, S.D. = 0.17; p = 0.02, d = 1.38) and at baseline (mean = 0.48, S.D. = 0.14, v. mean = 0.66, S.D. = 0.15; p = 0.02, d = 1.24). With baseline depressive symptoms in the model, reward seeking at 8:2 remained significantly associated with new-onset depressive disorder (Nagelkerke R² = 0.07, B = 0.06, S.E. = 0.03, p = 0.10 for depressive symptoms; Nagelkerke ΔR² = 0.12, B = −0.51, S.E. = 0.27, p = 0.05 for reward seeking), and reward seeking at 9:1 was significantly associated with new-onset depressive disorder at trend level (Nagelkerke ΔR² = 0.11, B = −0.41, S.E. = 0.23, p = 0.07). One adolescent with new-onset depression had an anxiety disorder at baseline. Excluding this participant did not alter the results. Adolescents with new-onset depression at follow-up were not significantly less risk adjusting at baseline (mean = 0.64, S.D. = 1.05 v. mean = 1.01, S.D. = 0.71; p = 0.31).

To test for the specificity of the reward seeking and new-onset depression association, we examined the association between reward seeking and new-onset anxiety or externalizing disorders when excluding those with baseline anxiety or externalizing disorders respectively. The analysis showed that reward seeking at baseline was not associated with new onset of anxiety disorders (n = 16; B = −1.80, S.E. = 1.54, p = 0.24) or externalizing disorders (n = 7; B = 1.68, S.E. = 2.32, p = 0.47) at follow-up.
Discussion

We examined the association of reward responding with adolescent depression, measures of environmental/social functioning and impairment. Adolescents with current depressive disorder were less reward seeking than adolescents without psychopathology for trials where a positive outcome was very likely (at ratios 9:1 and 8:2). These findings are consistent with Forbes et al. (2007) and with previous reports of depressed adults (Henriques & Davidson, 2000; Pizzagalli et al. 2005). Moreover, the response profile at ratio 8:2 was specific to depressive disorder. The current results did not differ when gender and pubertal status were entered as between-subject factors and diagnostic groups did not differ in age and IQ. Crucially, differences in reward seeking were apparent in the absence of group differences in the quality of decision making, impulsivity and deliberation time. Thus, deficits in reward seeking seem to represent a feature of adolescent depression that is not secondary to psychomotor impairment (a key symptom of adult depression). Depressed adolescents seem no worse at making decisions about, or identifying possibilities for, reward but are less likely to engage in reward-seeking behaviour and this seems unlikely to be attributable to impulsivity.

Diminished reward seeking under high-probability reward conditions may translate to low levels of positive environmental engagement (e.g. social relationships, education, activities), which over time is likely to impact on fundamental aspects of adolescent and adult life. Our results show that reward seeking at highly favourable ratios and risk adjustment (i.e. adjusting betting behaviour in line with the likelihood of reward) were correlated with indices of social/environmental functioning (e.g. humour, extracurricular activities) and anhedonia. The relationship between reward processes, physical activity and social functioning is consistent with imaging studies. Humour, interaction with friends and exercise engage neural substrates of the reward network (Mobbs et al. 2003; Brene et al. 2007; Guroglu et al. 2008). Changes in depressive symptoms do not fully explain changes in social functioning/activity level (Denninger et al. 2011). Thus, a key question for future research is whether alterations in reward processes mediate impairments in positive affective functioning in depression (e.g. social withdrawal).

Our main aim was to test whether altered reward responding may constitute a risk factor for the onset of depression (Kraemer et al. 2001). Our findings showed that reward seeking at highly favourable ratios was associated with depressive symptoms and new onset of depressive disorder at follow-up in adolescents free from depressive disorder at baseline. These findings remained significant when we controlled for pubertal development and depressive symptoms at baseline. Adolescents with new-onset depressive disorder were less reward seeking than adolescents who remained free from depression. Moreover, reward seeking predicted functional...
impairment at follow-up in adolescents free from depressive disorder at baseline. These findings illustrate that hypo-responsivity to reward is associated with the development of depression in adolescents at familial risk for affective disorder and that reward seeking predicts depression onset above and beyond baseline depressive symptoms. Reward seeking did not predict the onset of externalizing or anxiety disorders, suggesting that lowered reward responding represents a specific behavioural vulnerability marker for depression.

The limitations of this study merit consideration. The generalizability of the findings is limited by the small number of new-onset cases with depressive disorder at follow-up. Nevertheless, we chose a conservative approach for the longitudinal data analysis and excluded individuals with current depression. Moreover, the results converged with those from cross-sectional analysis of depressive disorder and longitudinal analysis of depressive symptoms. Although there were some missing data for the reward task, which reduced our sample size, missing data were not associated with psychopathology. One adolescent was receiving antidepressants at the time of reward task completion. Excluding this individual from analyses did not alter the results. The follow-up interval was approximately 1 year and it is not known how reward-seeking behaviour is related to depression over longer time periods. Reward processes in new-onset depression compared to recurrence require consideration as these may involve different processes (Kendler et al. 2000). The present sample comprised only adolescents at familial risk for depression. We were therefore unable to determine a potential influence of parental depression on adolescent reward seeking and whether diminished reward seeking predicts depression over time in the absence of familial risk. Replication in population-based studies is needed to assess the generalizability of these findings. Given the assessment time frame of the CAPA (the preceding 3 months), it is possible that some episodes of disorder may have been missed. However, this would probably have made analyses more conservative. Finally, reward seeking requires consideration within the context of other phases of reward processing (e.g. anticipation, outcome), processes of negative affect and regulatory strategies (Somerville et al. 2010). An understanding of their interplay should further advance risk characterization for depression and other psychiatric disorders.

In conclusion, the current findings show an association between abnormalities in reward processing and depressive disorder in a high-risk sample of adolescent males and females. The findings are novel in that they suggest that diminished reward seeking at highly favourable reward conditions is specific to current depressive disorder and is associated with future depressive symptoms, functional impairment and the onset of depressive disorder in adolescents without a diagnosis of depression at baseline. These findings illustrate that similar impairments in reward processing characterize both current and impending depression. Behavioural alterations in reward processing were also associated with social behaviour and engagement in everyday life. Thus, this is a potential mechanism through which reduced reward seeking confers risk for the development and maintenance of depressive disorder.

Several interventions based on cognitive behavioural therapy (CBT) have been found to prevent the onset of adolescent depression in a range of high-risk groups (Garber et al. 2009; Merry, 2009; Stice et al. 2009). Initial evidence suggests it may be possible to alter reward processing with psychological therapies (Dichter et al. 2009; Geschwind et al. 2011). Incorporating strategies to boost effective reward responding into such preventive interventions may be worthwhile.

<table>
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Anhedonia was derived from Mood and Feelings Questionnaire (MFQ) items associated with anhedonic symptoms. *p < 0.05, **p < 0.01.
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Declaration of Interest

None.

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Geschwind N, Peeters F, Drukker M, van Os J, Wichers M (2011). Mindfulness training increases momentary positive emotions and reward experience in adults vulnerable to...


