Cognitive biases predict symptoms of depression, anxiety and wellbeing above and beyond neuroticism in adolescence

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Neuroticism & cognitive vulnerability to depression in adolescence

Introduction

Depression is the leading cause of disability worldwide (WHO, 2017). Research shows that mental health disorders, including depression and anxiety, originate in adolescence (MQ, 2016). Lifetime prevalence in adolescents (13-18-year olds) was reported to be 11% for depression and 32% for anxiety (Avenevoli et al., 2015; Merikangas et al., 2010). Importantly, earlier onset of depression predicts longer episodes, more severe course, poorer recovery and higher recurrence rates (Dunn and Goodyer; 2006). In addition, it is important to consider positive measures of wellbeing and quality of life in combination with psychopathology and distress. Mental health is an integral component of wellbeing; in particular, wellbeing is considered to be greater than the absence of mental illness (World Health Organisation (WHO), 2013). Traditional emphasis on disorder and distress neglects the importance of wellbeing factors and positive measures of quality of life.

Adolescence represents a sensitive neurodevelopmental window for the fostering of lifelong positive mental health (Marco, Macri and Laviola, 2011). Therefore, there is an urgent need to understand risk factors that are specific to the adolescent period. A major risk factor for anxiety and depression is neuroticism (Navardy et al., 2017). Neuroticism has been considered a means of indexing risk and a general risk factor influencing the onset and course of psychological disorders (Klein et al., 2011). Lahey (2009) provided a discussion regarding the powerful predictive value of neuroticism in relation to longevity, psychiatric and physical health disorders.

Evidence supports a biological basis of neuroticism which is considered to be a largely heritable trait, sharing genetic factors underlying risk for internalising disorders (Luciniao et al., 2018, Hettema, et al., 2006). However, there is mixed evidence
regarding the stability of neuroticism over time. Wray et al., (2007) examined genetic and environmental contributions to neuroticism across 22 years in over 20,000 individuals, to demonstrate that genetic correlations between measures over time were very high. However, environmental contributions demonstrated lower stability over time (Wray et al., 2007). Additionally, neuroticism has been shown to reduce following antidepressant administration (Tang et al., 2009), suggesting that the impact of improving mood through treatment may also have a more general impact affecting neuroticism. However, other interventions, such as cognitive, treatments have not demonstrated changes in neuroticism (Tang et al., 2009).

As neuroticism is not universally considered modifiable, it is important to consider mechanisms above and beyond neuroticism that may be more amenable to intervention. Cognitive factors have been targeted by mainstream treatments such as cognitive behavioural therapy (CBT; NICE 2005). However, CBT has demonstrated limited success in adolescents (Goodyer et al., 2017). It may be that CBT is less efficacious due to ongoing neurocognitive development and environmental factors specific to adolescence. As such, the identification of cognitive features that predict depression in this age range may help improve interventions with the potential for lifelong impact.

Negative attributional style and rumination are cognitive processes that have frequently been related to disorders. Meta-analyses consistently demonstrate that rumination is predictive of the onset, severity and course of symptoms of depression (Mor and Winquist, 2002; Nolen-Hoeksema, 2000). Similarly, negative attributional style is considered to create a vulnerability to depression (Lee and Hankin, 2009). In
adolescents, McLaughlin and Nolen-Hoeksema (2011) demonstrated rumination fully mediated associations between anxiety and depression, identifying it as a transdiagnostic risk factor and target for intervention.

Negative cognitive biases of information processing have been proposed as mechanisms underlying the vulnerability, onset and maintenance of depression. Williams et al., (1997) proposes that negatively biased elaborative processing characterises depression. This necessitates increased allocation of cognitive resources to negative materials, resulting in the encoding of negative elaborations to memory, thereby enhancing memory for depression related materials. Previous research has examined cognitive biases in depression in adults and identified the importance of assessing biases across multiple domains of processing, a key limitation of research amongst adolescents (Everaert et al., 2015).

**Autobiographical Memory**

Impairment of specific autobiographical memory recall has been robustly associated with adult depression (Williams et al., 1997). Overgeneral memory refers to a memory which lacks specificity and lasts an extended duration (Williams et al., 1997) Sumner et al., (2011) implicated overgeneral memory in the onset of depression, demonstrating that overgeneral memory retrieval was predictive of depressive relapse in adolescents experiencing chronic interpersonal stress. However, studies of autobiographical memory and depression in adolescents, including clinical, community and at-risk samples, have produced mixed findings (Swales et al., 2010; Chan et al., 2007). Overall, there is a lack of consensus regarding the context and mechanisms of autobiographical memory performance and the majority of previous studies were based on adult patient samples.
Self-Referential Memory

Biased self-referential memory has also been implicated in emotional disorders. A study of depressed youth employed a self-referent encoding task to demonstrate memory bias in the recall of negative compared to positive words (Zupan et al., 1987). However, Timbremont et al., (2008) found no differences in memory biases between currently, never and previously depressed youth. A recent review (Platt et al., 2017) concluded that studies using self-referent encoding tasks have produced mixed findings and evidence of negative biases is inconclusive across clinical, at risk and community samples.

Interpretation Bias

Negatively biased interpretation has also been associated with depression in adolescents. Orchard et al., (2016) demonstrated that adolescents with depression make significantly more negative interpretations of ambiguous scenarios than non-depressed patient and community control groups. However, Micco et al., (2014) showed that interpretation bias modification reduced negative biases in depressed and control groups of adolescents, but there was no associated change in anxiety or depression. Again, overall conclusions are limited by the scarcity of studies in the adolescent population.

Biased Processing of Emotional Facial Expressions

Empirical evidence of deficits or biases in emotional processing of facial expressions associated with adolescent depression are particularly inconsistent. Schepman et al., (2011) found no deficits of accuracy amongst depressed adolescents, although the depressed group demonstrated a negative bias at low intensities compared to controls.
Contrastingly, Joormann et al., (2010) demonstrated impaired accuracy of identification and that children at high familial risk for depression required greater emotional intensity of facial stimuli to accurately identify sad facial expressions. These inconsistent results could be due to varying methodology.

While there is substantial, although mixed, research supporting the relationship between cognitive biases and depression in adults, the adolescent population has been less studied and understanding of cognitive biases within this age group is limited. Furthermore, few studies have assessed multiple realms of biases within the same sample, this may account for discrepancy within results. This study aimed to address this limitation by assessing multiple realms of processing within the same sample.

The high prevalence of mental ill-health within the adolescent population (Avenevoli et al., 2015), alongside the conceptualisation of mental health as a spectrum indicates that symptoms are likely present within individuals recruited from community settings. Furthermore, Wang et al., (2005) found a median 8-year delay between symptom onset and contact with services. Therefore, sampling restricted to patient groups risks overlooking individuals who are not help-seeking, those who may be demonstrating pre-clinical or sub-diagnostic symptoms, or those exhibiting resilience factors.

To achieve a representative sample, this study therefore aimed to recruit adolescents from community settings to examine to what extent cognitive processes (attribution style, rumination and cognitive biases) are able to predict three outcome variables (depression, anxiety and wellbeing) within adolescents. Secondly, to consider the contribution of cognitive processes in relation to neuroticism. This study has examined contributions of cognitive bias factors independently from neuroticism in order to distinguish the influence of cognitive factors distinct from underlying personality risk.
We hypothesised that cognitive measures will predict depression, anxiety and wellbeing above that of neuroticism in adolescence, and that there may be distinct contributions of biases to depression, anxiety and wellbeing.
Methodology

Participants
Inconsistencies exist within the research literature as to the age range distinguishing the ‘adolescent’ period. This developmental stage is associated with social and neurological changes as well as related to an increased risk for psychiatric disorders (Caasey et al., 2008; Kessler et al., 2005). In line with previous research this study sought to recruit adolescents aged 12-18 years (mean=14.9, S.D. = 1.52, N=99) from Scottish secondary schools (58.6% female and 70.3% White British; Table 1). A power calculation determined that the current sample size allowed for up to 6 predictors in a regression model with 80% power to detect a medium effect size.

Measures
All of the following standardized measures have been used in previous research with adolescents and considered valid and reliable. Their reliability in this sample has been confirmed with high Cronbach’s alphas, as reported below. Depressive symptoms were assessed using the Mood and Feelings Questionnaire (Child Version; MFQ; Angold and Costello; 1987; α=.91). Anxiety symptoms were assessed using the Spence Children’s Anxiety Scale (SCAS; Spence; 1998; α=.89). Wellbeing was assessed using the BBC Well-being Scale (BBC; Kinderman, Schwannauer, Pontin and Tai; 2011; α=.95). Neuroticism was measured with the 12-item short version of Eysenck Personality Questionnaire-Neuroticism subscale (EPQ-N; Eysenck et al., 1985; α=.83). The Ruminative Response Styles scale (RRS; Treynor et al., 2003; α=.87) measured rumination; higher scores indicate more ruminative thinking. Dysfunctional attribution style was measured with the Dysfunctional Attitudes Scale:
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24 Item Version (DAS-24; Power et al., 1994; \( \alpha = .80 \)) with lower scores indicating more dysfunctional thoughts.

**Self-Reference Recall Task (Kelvin et al. 1999)**

This task assessed biases of self-referential memory recall and consisted of 12 positive and 12 negative adjectives. This task has been previously employed in research literature and significant differences in this task were demonstrated in participants before and after negative mood induction (Kelvin et al., 1999). Participants indicated the extent to which each word described them on a four-point scale (1='not at all like me', 4='very much like me'). To counter primacy and recency effects, three neutral adjectives were included at the beginning and end, which were excluded from analysis. Ratings were recoded as either: not self-referent ('not me'), or self-referent ('me'). Participants were subsequently asked to recall descriptors in a free recall task. A proportional score was calculated to account for the overall number of words recalled. Self-referent and non-self-referent conditions were calculated separately; positive scores reflect a positive bias whereas negative scores indicate a negative bias (Connolly, Abramson and Alloy, 2016).

**Autobiographical Memory Test (AMT; Williams and Broadbent, 1986)**

Participants were presented visually and orally with five positive (relaxed, lucky, excited, relieved, loved) and five negative (hopeless, failure, sad, angry, lonely) cue words, in a randomised order, and asked to recall a specific memory. This was defined as “a memory of a particular event that occurred on a particular day which could be from a long time ago or very recently and could be something very important or
something very ordinary". Participants were given 60 seconds to produce a memory, which they then verbally described under no time condition. This task was audio recorded and later coded for specificity. Participants were given two practice trials, where they were prompted to recall a specific event if their memory did not fit the criteria of a specific memory. Practice items were excluded from analysis. Responses to positive and negative cues were recorded as either specific (e.g. “My friend took me to the Manchester derby on my birthday.”) or overgeneral (e.g. “During the summer holidays when there was no one around.”). The total number of specific and overgeneral memories was calculated. This task has been extensively used within research and has previously demonstrated good psychometric properties (Griffith, Kleim, Sumner and Ehlers, 2012).

*Ambiguous Scenarios Task for Depression in Adolescents: (AST; Orchard, Pass and Reynolds; 2016)*

Participants were presented with 20 hypothetical ambiguous scenarios designed to assess interpretation bias in relation to depression, asked to consider each as happening to them and imagine the outcome. For example: “You see a man running down the street and think about why he is running”. Participants were asked to write down their imagined outcome, and rate its pleasantness on a 9-point scale (1=‘extremely unpleasant’, 9=‘extremely pleasant’). Orchard et al., (2016) demonstrated good psychometric properties of this task. Only written descriptions were included in the current analysis due to lower internal reliability of the rating scale in this sample (α=.68). Written descriptions were coded as positive, negative, neutral or mixed. Overall bias scores were calculated by subtracting number of negative
responses from number of positive responses. Therefore, positive scores reflect a positive bias whereas negative scores indicate a negative bias.

Facial Expression Recognition Task (Chan et al., personal communication)

This computer task, based on a task employed by Chan et al., (2007) and adapted by Chan et al. (personal communication), presented five emotional expressions: anger, disgust, fear, happy and sad. Stimuli were morphed using Morpheus Photo Morpher v3.17 software, from 0-100% intensity at incremental increases of 10%, with 0% reflecting a neutral expression and 100% reflecting full intensity of expression. In total 220 stimuli were presented in a random order across five blocks. Each face was presented for 500ms against a black background preceded by a fixation cross of 100ms. Participants were asked to identify the emotion displayed by a key press. Prior to the experimental trials participants completed six practice trials, which were excluded from analyses. Mean accuracy of identification was computed across each emotion. An accuracy score was calculated for each emotion.

Procedure

This study was approved by University Research Ethics Committee and local educational authorities. Written consent from participants and parents for those under 16 was obtained. Participants completed measures of depression, anxiety, wellbeing, neuroticism, rumination, and dysfunctional attitudes online. Tasks assessing memory, interpretation and facial expression recognition tasks were conducted during a face-to-face interview with a trained psychologist either individually or in small groups. Self-report and cognitive tasks were conducted within the same week.
Statistical Analyses

Analyses were primarily conducted in IBM SPSS version 22 (SPSS Inc., USA). Three participants did not complete the facial recognition task. Due to a procedural error, item 13 from the BBC Well-being Scale was missing at random and therefore excluded and mean rather than total scores were employed for further analysis. The reliability of this scale remained high ($\alpha=.95$).

Assumptions of normality were met. Despite variables being correlated collinearity and tolerance statistics were within accepted limits (VIF=1.50 and tolerance =0.61; Field, 2009).

False discovery rate (FDR) correction for multiple comparisons was applied to control for familywise error rate (Benjamini and Hochberg, 1995).

Analysis comprised two stages. Initially, to identify salient predictors, backwards-elimination regressions were employed based on a criterion of $F>=.100$. Subsequently, surviving variables were entered in a hierarchical regression to examine their contribution in relation to neuroticism.

The impact of gender and age were assessed using t-test and Pearson’s correlation, if tests demonstrated significant effects these variables were included in regression models.
Each regression model consisted of two steps to predict each dependent variable (depression, anxiety and wellbeing) in turn: 1. Neuroticism, 2. Cognitive variables surviving backwards removal regression. When age and gender were included within regression models, these did not impact any results and as such have been excluded as variables in order to preserve statistical power. Gender was included in models of anxiety as univariate tests indicated significant relationships between gender and anxiety.
Results

See Table 1 for demographic information and descriptive statistics.

Age and Gender Differences

Female participants reported higher levels of anxiety than male participants: \( t(97)=2.83, p=0.006 \). No significant gender differences were demonstrated for depression (\( t(97)=-1.22, p=0.27 \)) or wellbeing (\( t(96)=1.85, p=0.07 \)). There were no significant correlation of age with depression (\( r=0.13, p=0.20 \)), anxiety (\( r=0.06, p=0.55 \)) or wellbeing (\( r=-0.13, p=0.21 \)).

Correlations

Pearson’s correlation results, controlling for age and gender, are presented in Table 2 and Figure 1.

Pearson’s correlations, controlling for age and gender, indicate a significant relationship between depression and anxiety and a negative relationship between depression and wellbeing. Neuroticism was positively correlated with depressive symptoms and anxiety, while negatively correlated with wellbeing. See Table 2 and
Figure 1 for full results. Briefly, correlations demonstrated the expected pattern with higher depression and anxiety symptoms as well as lower wellbeing generally correlated with greater negative biases, reduced positive biases or more maladaptive processing. However, mean accuracy bias scores of facial emotion recognition and autobiographical memory bias scores were not significantly correlated with any other variables.

[insert figure 1 here]

Backwards-Elimination Regression

Variables surviving the regression model of depression were: rumination, dysfunctional attitudes and non-self-referential recall bias. Those surviving the regression model of anxiety were: ambiguous scenarios bias and rumination. The ambiguous scenarios task bias, self-referential recall bias, dysfunctional attitudes and accuracy of anger identification were included in the wellbeing regression model (Supplementary Material).

Hierarchical Multiple Regression

See Table 3.

Depression

In predicting depression, neuroticism was a significant predictor ($r^2 = .40 \ p<0.001$, $\beta=0.35$). The addition of cognitive variables: rumination ($\beta=0.26$), dysfunctional attitudes ($\beta=-0.19$) and self-referential ‘not me’ recall scores ($\beta=0.17$), resulted in an
increase of explained variance ($\Delta r^2 = .12$, $p < .005$; $r^2 = .52$, $F(4,94)=25.49$, $p < .001$). All included variables demonstrated significant beta values.

**Anxiety**

In the first block gender alone significantly predicted anxiety ($r^2 = .08$, $p < .05$), as did the addition of neuroticism ($r^2 = .54$, $p < .001$, $\beta = .55$). Inclusion of cognitive variables (ambiguous scenarios task bias ($\beta = -0.13$) and rumination $\beta = 0.21$) demonstrated a small but significant increase of explained variance ($\Delta r^2 = .05$ and $r^2 = .59$; $F(3,95)=44.46$, $p < .001$). Only neuroticism and rumination demonstrated significant beta values.

**Wellbeing**

In the first block of the model of wellbeing, neuroticism significantly predicted wellbeing ($r^2 = .36$, $\beta = -0.32$, $p < .001$). Inclusion of cognitive variables: ambiguous scenarios task bias ($\beta = 0.19$), self-referential recall ‘me’ bias ($\beta = 0.18$), self-referential recall ‘not-me’ bias ($\beta = -0.16$), dysfunctional attitudes ($\beta = 0.17$), and recognition of angry facial expressions ($\beta = -0.14$), demonstrated a significant $r^2$ change ($\Delta r^2 = .19$, $p < .001$; $r^2 = .55$, $F(6,89)=17.74$, $p < .001$). Only neuroticism, ambiguous scenarios task bias and self-referential recall ‘me’ bias demonstrated significant beta values.

[insert table 3]

**Discussion**
Rumination, dysfunctional attitudes, and negative biases in ambiguous scenarios interpretation and self-referential memory, significantly predicted depression in adolescents above and beyond neuroticism. Neuroticism predicted around 40% of the variance of depression symptoms, 54% of anxiety symptoms and 35% of wellbeing, in line with previous research (e.g. Kotov et al., 2010 and Bartels et al., 2013). As hypothesised, adding cognitive variables significantly increased the explained variance of depression by 12%, anxiety by 5% and wellbeing by 19%. This supports previous findings that have associated negative cognitive biases with psychological outcomes. Orchard (2016) demonstrated that biased interpretation was most negative for adolescents with a depression diagnosis and most positive for non-depressed controls. The current study highlights the salience of interpretation bias across psychological distress and wellbeing.

In an undergraduate student sample, Rude et al., (2002) demonstrated that including a measure of interpretation under a cognitive load condition increased predictions of depression scores by 11% for male participants and 0% for female participants. This increase is in line with the current findings; however, we did not identify such gender disparity. Rude et al. (2002) suggest that such divergence may be due to gender bias in self-report or that despite decreased negative biases in women, their subsequent behaviour or processing (such as rumination) increases their vulnerability to experience depression.

This study demonstrates, similar to findings of adult populations, neuroticism to be a key predictor of mental health outcomes (Lahey, 2009). Neuroticism was the strongest
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predictor of each mood variable; each standard deviation increase of MFQ was associated with a 0.55 increase of neuroticism. This magnitude is similar to findings of a meta-analysis of 33 population based samples (Malouff, Thorsteinsson and Schutte, 2005). The predictive power of neuroticism cannot be overlooked as it indicates underlying biopsychological components of depressive disorders worthy of further neurobiological study to assess its expression and mechanism of action.

Like Zupan et al., (1987), this study indicates that greater depressive symptomology and anxiety was strongly correlated with a bias towards recalling negative self-referenced words. However, for non-self-referential words, higher levels of depression and anxiety were related to recalling more positive adjectives. This suggests that symptoms of depression and anxiety are not associated with global negative biases, but a specific negative bias in relation to the self. This is consistent with findings of more negative self-perceptions in young people at risk for depression (Chan et al., 2007). Previous research has identified a self-positivity bias within healthy individuals; whereby, individuals are more likely to overestimate their own success compared to the success of others. However, depressed individuals have been shown to lack such positivity bias (Alloy and Ahrens, 1987). Self-concept and self-referent bias may be of particular significance within adolescent populations. Adolescence is a sensitive period were self-concept is developing (Marcia, 1980), biases towards negative self-descriptors at this age group may impair the development of adaptive self-concepts.

Regression models predicting each outcome variable included different predictors, signaling salience of distinct cognitive bias in depression, anxiety and wellbeing. Specifically, dysfunctional attitudes emerged as a significant predictor of depression
and wellbeing. Robinson and Alloy (2003) found that dysfunctional attitudes and components of rumination interacted to prospectively predict onset, number and duration of depressive episodes, in an undergraduate sample. In the present study, dysfunctional attitudes were found to be more strongly related to depression and wellbeing than anxiety. This is expected as the dysfunctional attitudes scale was developed to capture thinking styles associated with depression (Power et al., 1994). Rumination is a significant research focus and has been frequently related to emotional disorders (Young and Dietrich, 2015). Consistent with this, our findings demonstrate that rumination predicted depressive and anxiety symptoms but was less strongly associated with wellbeing.

No significant relationships between autobiographical memory recall and depression, anxiety or wellbeing were demonstrated. Previous findings have been inconsistent. Chan et al., (2007) demonstrated no significant differences in this task between individuals at high vs. low risk for depression by virtue of neuroticism. Further, Swales et al. (2010) found group differences in specificity of autobiographical memory when comparing clinical groups to controls, but that this was due to individuals within clinical groups recalling the same suicide related memories in response to multiple cue words. However, autobiographical memory impairment in depressed groups is a robust finding with large effect sizes within adult samples (Williams et al., 1997). The above mixed findings may indicate that differences in autobiographical memory are related to symptom severity; such bias may be a scar effect rather than an antecedent risk; or that, contrary to adult research, overgeneral memory is not a reliable cognitive marker of adolescent depression.
This study demonstrated that accuracy in identification of angry faces predicted wellbeing. This indicates that individuals with lower wellbeing were better able to recognise anger, suggesting higher sensitivity towards negative facial expressions. This is consistent with previous research which found that recognition accuracy of negative emotions to be associated with depressive relapse (Bouhuys et al., 1999). This finding supports the protective value of wellbeing in that, the association with a positivity bias in facial emotion identification may foster positive social interactions. However, the effect size of our finding was small and no other significant relationships between psychological outcomes and facial expressions were demonstrated. Previous research demonstrated inconsistencies which may indicate that facial emotion recognition bias is not a robust marker of mood disorders. Alternatively, there may be a task failure to detect subtle differences in interpretation. Picci and Scherf (2016) showed that adolescents were significantly better at identifying faces of their own age and hypothesised a ‘dip’ in facial recognition whereby there is a recalibration of the face-processing system away from caregivers towards peers. It is possible that the use of adult faces has interfered with an effect of emotion. Review of adult studies of facial emotion interpretation indicates evidence supporting an increased tendency to interpret facial expressions more negatively (Bourke, Douglas and Porter, 2010). However, like within adolescent samples, there is discrepancy with some evidence indicating a global deficit of facial emotional processing, while some studies support a mood-congruent bias towards negative emotions in depressed groups.

A number (18%) of participants reached scores associated with clinical depression on the MFQ (Daviss et al., 2006), consistent with prevalence estimates within this age group (Avenevoli et al., 2015). This indicates the representative nature of the sample.
A key strength of this study has been the assessment of wellbeing. The mental health spectrum ranges from highly disabling disorder to positive states of wellbeing. Components of wellbeing are recognised protective factors against disorders (NHS Scotland, 2016). Our results highlight the importance of cognitive biases for subjective wellbeing. Aiming to enhance positivity bias in order to boost wellbeing, potentially protecting against depression or anxiety in preventative or early intervention strategies, may be an avenue of future research. Future research may investigate the potential to develop interventions that address specific biases, relevant to individuals’ experiences and symptoms, particularly in light of the importance of cognitive contributions (rumination, dysfunctional attitudes, self-referent processing and interpretation biases). Similarly, development of interventions addressing cognitive bias in order to enhance wellbeing, which is of importance for quality of life, has been associated with favourable life outcomes, including longevity (Sadler et al., 2011).

There are some limitations of this study; primarily, predictions are based on regression analyses of cross-sectional data, limiting the ability to conclude causality. Similarly, age has been demonstrated to exert non-linear development which has not been accounted for within this study. Self-report questionnaires have been employed rather than clinical interviews. The latter was deemed less feasible due to issues of anonymity, confidentiality and the non-clinical nature of the sample. Further, although the impact of neuroticism has been analysed as distinct to cognitive factors, it is possible that neuroticism itself is influenced by mood state and biases, which may impact interpretations of the current results. Within this study, including neuroticism in initial blocks has allowed for the explanation of unique variance by cognitive biases. In future work a larger sample may be employed to allow for full mediation analysis to examine such effects.
To examine a wide range of mood states, participants were recruited from community settings, therefore results are less generalizable to clinical groups but are representative of typical adolescents. Recruiting adolescents is notoriously difficult. While the hierarchical regression models were sufficiently powered, initial identification of salient variables was underpowered and as such, variables with smaller effect sizes may not have been identified. Finally, our regression models explained approximately 50% of the variance of depression, anxiety and wellbeing, indicating that there are important factors that have not been accounted for here. For example, stressful life events and general cognitive performance would be valuable factors to include in future studies.

**Conclusion**

This study assessed a relatively wide range of cognitive biases within a single sample using validated and standardized measures in combination with experimental paradigms to demonstrate that cognitive biases accounted for variability in depression, anxiety and wellbeing over and above that of neuroticism. Results stress the importance of cognitive factors in symptoms of depression, anxiety and wellbeing. Contributions of cognitive mechanisms, identified here, are a feasible target for behavioural and cognitive modification and improvement of interventions, potentially targeting specific biases and to enhance wellbeing as a protective factor are worthy of further study.
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Table 1: *Descriptive Statistics*

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<th>Measure</th>
<th>Mean</th>
<th>Std. Deviation</th>
<th>Minimum</th>
<th>Maximum</th>
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<td>1.52</td>
<td>13</td>
<td>18</td>
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<tr>
<td>MFQ (Depression)</td>
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<td>12.88</td>
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<td>SCAS (Anxiety)</td>
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<td>BBC (Wellbeing)</td>
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<td>Neuroticism</td>
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<td>RRS (Rumination)</td>
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<td>AMT: Autobiographical Memory Task (Overgeneral)</td>
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<td>SRR: Self-Referential Recall 'Me'</td>
<td>0.33</td>
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<tr>
<td>SRR: Self-Referential Recall 'Not Me'</td>
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<td>Facial Expression Recognition: Anger</td>
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### Table 2: Pearson’s Correlation Coefficient (r) between outcome variables and cognitive variables, controlling for age and gender

<table>
<thead>
<tr>
<th>Cognitive Variable</th>
<th>MFQ</th>
<th>SCAS</th>
<th>BBC</th>
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Note: ** indicates significance at p<0.001; * indicates significance at p>0.05. MFQ refers to the Mood and Feelings Questionnaire, SCAS refers to Spence Child Anxiety Scale, BBC refers to the BBC Well-being Scale.
Table 3: Hierarchical Regression Models of Mood Variables

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Note: **indicates significance at p<0.001; *indicates significance at p>0.05. MFQ refers to the Mood and Feelings Questionnaire, SCAS refers to Spence Child Anxiety Scale, BBC refers to the BBC Well-being Scale, DAS refers to the Dysfunctional Attitudes Scale, RRS refers to the Ruminative Response Scale, AST refers to the Ambiguous Scenarios Scale, SRR refers to the Self-Referential Recall task. Anger Accuracy refers to the proportional accuracy of the facial emotional expression questionnaire.
Correlation Map: Strength of Association

Figure 1: *Strength of Association Heat Map*

*Note:* Figure depicts Pearson correlation between variables. For simplicity, Facial Emotion Recognition Bias is represented by a mean across emotions. Diagonal stripes indicate negative correlation. MFQ refers to the Mood and Feelings Questionnaire, SCAS refers to Spence Child Anxiety Scale, BBC refers to the BBC Well-being Scale, DAS refers to the Dysfunctional Attitudes Scale, RRS refers to the Ruminative Response Scale, AST refers to the Ambiguous Scenarios Scale, SRR refers to the Self-Referential Recall task. ABMT refers to the Autobiographical Memory Test.