Title: Autobiographical memory as a latent vulnerability mechanism following childhood maltreatment: Association with future depression symptoms and prosocial behaviour

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Keywords: Maltreatment, depression, autobiographical memory, prosocial behaviour, conduct problems

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Abstract:

Objectives: Childhood maltreatment is associated with altered neural reactivity during autobiographical memory (ABM) recall and a pattern of over-general memory (OGM). Altered ABM and OGM have been linked with psychopathology and poorer social functioning. The present study investigated the association between altered ABM and subsequent socio-emotional functioning (measured two years later) in a sample of adolescents with (N=20; Maltreatment group, MT) and without (N=17; Non-MT group) documented childhood maltreatment histories.

Method: At baseline, adolescents (aged 12.6±1.45 years) were administered the Autobiographical Memory Test to measure OGM. Participants also recalled specific ABMs in response to emotionally valenced cue words during functional MRI. Adolescents in both groups underwent assessments measuring depressive symptoms and prosocial behaviour at both timepoints. Regression analyses were carried out to predict outcome measures at follow-up controlling for baseline levels.

Results: In the MT group, greater OGM at baseline significantly predicted reduced prosocial behaviour at follow-up and showed a trend level association with elevated depressive symptoms. Patterns of altered ABM-related brain activity did not significantly predict future psycho-social functioning.

Conclusions: The current findings highlight the potential value of OGM as a cognitive mechanism that could be targeted to reduce risk of depression in adolescents with prior histories of maltreatment.

Keywords: Maltreatment, depression, autobiographical memory, prosocial behaviour, conduct problems
1. Introduction

Childhood maltreatment has consistently been associated with significantly increased risk for concurrent and future internalizing and externalizing psychiatric disorders (Gilbert et al., 2009). Recent studies have investigated the potential neurocognitive mechanisms contributing to elevated psychiatric risk in children who have experienced maltreatment (McCrory et al., 2017). These studies have documented altered functioning in a range of neurocognitive domains that have also been shown to characterize individuals presenting with frank psychiatric disorder, including threat processing, reward processing, emotion regulation and most recently autobiographical memory functioning (McCrory et al., 2011; McCrory et al., 2017, Puetz et al., 2016, Johnson, Greenhoot, et al., 2005; Valentino et al., 2009; Lawson et al., 2018). However, there has been a dearth of longitudinal studies that have included measures of brain function or psychological processes (see Dennison et al. (2016) for one exception). In an attempt to address this gap, we here conducted one of the first longitudinal functional imaging studies of children experiencing documented maltreatment. The theory of latent vulnerability postulates that an array of neurocognitive systems adapt to early adverse or neglectful environments in line with the notion of experiential canalization (Blair & Raver, 2012; McCrory & Viding, 2015). Such adaptations are thought to confer short-term functional advantages in atypical early environments but compromise the ability of the individual to negotiate more normative contexts as well as future stressor events. Moreover, such adaptations may also increase the likelihood that future stressful life events will occur (‘stress generation’ (McCrory, Ogle, Gerin & Viding, 2019)). A longitudinal design is necessary to truly interrogate any latent vulnerability construct.

Autobiographical memory (ABM) retrieval refers to the process by which an individual retrieves personally experienced events. The accurate and successful retrieval of autobiographical events is thought to play a critical role in scaffolding our sense of self (self-
coherence (Conway & Pleydell-Pearce, 2000)), the development and maintenance of social bonds (social function) (Alea & Bluck, 2003), and planning of present and prospective actions (directive function) (Pillemer, 2003). These functions are interdependent and it is argued that by using our past experiences to construct representations of others, attribute mental states to others (theory of mind) and subsequently predict other’s future thoughts and actions, ABM is an essential component of adaptive human social behaviour (Alea & Bluck, 2003; Nelson, 1993). This close relationship between autobiographical memory and (pro)social functioning is consistent with the finding that there is substantial overlap in brain areas engaged in social cognition and autobiographical memory processing (Spreng, Mar & Kim, 2009).

One way in which ABM functioning can be measured is by analysing recalled memories for specificity and level of detail (Williams et al., 1996). An ABM style that is more categorical and involves a less detailed recollection of personal events, known as overgeneral memory (OGM), compromises successful processing and regulation of emotions, interferes with self-reflection or self-projection in future scenarios and impedes social functioning by limiting social problem solving skills (Dalgleish et al., 2007; Goddard, Dritschel & Burton, 1996; Raes, Hermans, de Decker, Eelen & Williams, 2003; Rawal & Rice, 2012). According to the affect regulation hypotheses, recalling less vivid and detailed representations of traumatic or negatively valenced memories may serve to minimize negative affect, and thus have an adaptive short-term function (Williams et al., 1996). However, if generalized to everyday memories, such an OGM style may become maladaptive in the longer term by limiting access to prior successful strategies to navigate intrapersonal or interpersonal stress (Raes et al., 2003; Williams et al., 1996). Consistent with this view, OGM has been shown to be associated with, and can predict risk for, the onset of depressive disorder in adolescents (Hipwell, Sapotichne, Klostermann, Battista & Keenan, 2011; Rawal & Rice, 2012), in at-risk adults (Brittlebank,
Scott, Williams & Ferrier, 1993; Gibbs & Rude, 2004) and even non-clinical samples (Anderson, Goddard & Powell, 2010). In addition to the established relationship between OGM and depression, OGM has also consistently been associated with poorer social problem-solving skills, (Goddard, Dritschel & Burton, 1996), which predicts difficulty in the formation and maintenance of social relationships (Pillemer, 1992). However, more recent studies provide a more nuanced picture, such as Gutenbrunner and colleagues (2017), who showed that prospective associations between OGM and psychopathology emerged only among adolescents ‘at risk’, i.e. who reported elevated, and increasing patterns of rumination over time. With regards to the present investigation, it is important to note that amongst individuals at-risk, young people who experienced maltreatment have been shown consistently to present with OGM (e.g., Johnson, Greenhoot, et al; 2005; Valentino et al., 2009; Lawson et al., 2018).

In light of the above findings it is plausible that OGM may represent one candidate autobiographical memory mechanism that serves to index latent vulnerability to psychiatric disorder and poorer social functioning following maltreatment (McCrory & Viding, 2015). We recently investigated OGM behaviourally and autobiographical memory functioning at the neural level using functional MRI in 34 children with maltreatment and 33 matched controls. In line with previous findings, maltreatment status was associated with elevated OGM. At the neural level, maltreatment experience was associated with reduced hippocampal and increased parahippocampal activation during positive (vs. negative) autobiographical memory recall relative to their non-maltreated peers. Conversely, during negative autobiographical memory recall, children with maltreatment experiences exhibited increased activation in the amygdala as well as greater connectivity with other regions implicated in processing salient information. These findings are consistent with the view that altered autobiographical memory processing following maltreatment may index latent vulnerability to
future maladaptive outcomes. It is important to note that the brain activation paradigm was not designed to investigate OGM processes. Rather, in line with other studies, it entailed a task that involved successful recall of detailed events with affective content. Research in clinical and non-clinical adult samples without maltreatment histories have provided the strongest evidence to date supporting a link between overgeneral ABM retrieval (behaviourally) and depression and social functioning (Alea & Bluck, 2003; Dalgleish et al., 2007; Gino & Desai, 2012; Goddard et al., 1996; Williams et al., 1996). Neuroimaging findings indicate that the brain regions associated with altered ABM processing of specific memories in children who have experienced childhood maltreatment, namely the amygdala, the middle temporal gyrus (MTG), parahippocampal gyrus and hippocampus (Young, Bellgowan, Bodurka & Drevets, 2014) are also implicated in altered ABM processing of specific memories in depression (Young et al., 2014; Young, Siegle, Bodurka & Drevets, 2016). These neural differences are consistent with a relative privileging of negative versus positive memories during ABM processing.

However, the degree to which a tendency to recall autobiographical memories in an overgeneral way predicts future depression and reduced prosocial functioning has not yet been investigated in a maltreated sample. That is, OGM as a latent vulnerability factor following maltreatment experience has not been assessed to date within a longitudinal framework. In addition, the degree to which atypical neural processing associated with intact ABM recall predicts future vulnerability to depression and reduced prosocial functioning has also not been investigated. The primary aim of the current study was to address these outstanding questions in the literature and conduct one of the first longitudinal functional imaging studies of children experiencing documented maltreatment. Based on the extant evidence demonstrating that OGM predicts future depressive symptoms in at risk samples (Hipwell et al., 2011; Rawal & Rice, 2012; Guttenberg et al., 2018), we hypothesized that a similar pattern would be observed in
adolescents with maltreatment histories. We also hypothesised that OGM would predict future prosocial functioning, in the light of the established association of OGM social functioning in non-clinical samples (Alea & Bluck, 2003; Dalgleish et al., 2007; Gino & Desai, 2012; Goddard et al., 1996; Williams et al., 1996). In addition, we tested the hypothesis that the brain areas associated with atypical neural activation during ABM recall of specific valenced memories (MTG, hippocampus and amygdala) were related to higher levels of depression and lower levels of prosocial behaviour.
2. Method

2.1 Participants

Participants had been part of an earlier study on autobiographical memory processing in children with and without maltreatment experiences. At baseline, $n=77$ children ($n=34$ Maltreatment group; $n=33$ Non-Maltreatment group) underwent the Autobiographical Memory Test (see Blinded 3 for study details). From the initial sample of $n=77$, we retained a sample of $n=39$ ($n=19$ MT; $n=20$ NMT; mean age of 14.83±1.53 years; range 10.15 – 14.92 years in the control sample to 9.95 – 14.82 years in the MT sample) who had complete and usable data from the baseline assessment and complete data at follow-up for the measurement of depressive symptoms and prosocial behaviour (see below) and returned for a second assessment two years later. The main reason for drop-out at follow-up was the difficulty in maintaining and re-establishing contact with the children in the maltreatment group due to placement and social-worker changes. Children in the follow-up sample ($n=53$ before exclusions due to e.g. missing data) and those who did not return for follow-up ($n=9$ MT and $n=5$ NMT, $p=.26$) were comparable in terms of age ($p=.68$), gender ($p=.55$), IQ ($p=.50$), ethnicity ($p=.22$), SES ($p=.37$), maltreatment severity ($p=.24$) as well as OGM: ($p=.48$) and depressive symptoms T1 ($p=.16$) or prosocial behaviour ($p=.16$). Exclusion criteria were: pervasive developmental disorder, neurological abnormalities, standard magnetic resonance imaging (MRI) contraindications and IQ <70. Groups were closely matched on key demographics such as age, gender, SES, at baseline (see Blinded 3 for details) and remained matched on these parameters at follow-up except for ethnicity (see Table A).
The average time interval between baseline and follow-up was 27.18 ± 4.82 months, and between group analyses showed that groups were comparable on time interval between baseline and follow-up ($p>0.05$).

Children’s legal guardians gave written informed consent, while verbal and written assent was obtained from the children themselves. Verbal consent was witnessed and formally recorded.

The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. All procedures involving human subjects/patients were approved by UCL Research Ethics Committee (0895/002).

### 2.2 Measures

#### 2.2.1 Maltreatment history. Children in the maltreatment group had experienced a level of maltreatment requiring Social Services referral at baseline. Severity of neglect, emotional, sexual and physical abuse as well as intimate partner violence, were rated by the child’s social worker or adoptive parent on the basis of Social Service’s file information. Ratings from zero (not present) to four (severe) were applied to each category (Kaufman, Jones, Stieglitz, Vitulano & Mannarino, 1994). The majority of children had been exposed to neglect, emotional abuse and intimate partner violence. Additionally, children completed the Childhood Trauma Questionnaire (CTQ) (Bernstein & Fink, 1998). For details regarding maltreatment type and onset, duration and severity at baseline assessment see online Table DS1 in Blinded.

#### 2.2.2 Cognitive ability. Cognitive ability was assessed at baseline using two subscales of the Wechsler Abbreviated Scales of Intelligence (Wechsler, 1999), the word reading subscale of
the Wide Range Achievement Test (WRAT 4) (Wilkinson & Robertson, 2006), and two subtests of the CogState battery pertaining to visual and verbal episodic memory (see Blinded for details).

2.2.3 Psychiatric symptomatology. The depression symptom subscale of the Child and Adolescent Symptom Inventory (CASI) (Lavigne, Cromley, Sprafkin & Gadow, 2009) and the Strengths and Difficulties Questionnaire subscale assessing prosocial behaviour (SDQ, parent-rated) (Goodman, 1997) were used to characterize depression symptoms and social functioning. We have selected this common, well standardised measure of prosocial behaviour as one index of social functioning. These subscales were also chosen for analyses due to the established relationship between OGM and depression and social functioning (Dalgleish et al., 2007; Goddard et al., 1996; Raes et al., 2003; Rawal & Rice, 2012) and altered neural ABM functioning in patients with depression and at risk for depression (Young et al., 2014; Young et al., 2016). The hyperactivity subscale of the SDQ (parent-rated) (Goodman, 1997) as well as the Inattentiveness and Hyperactivity subscales of the CASI (Lavigne et al., 2009) were included in order to rule out the possibility that OGM and atypical ABM functioning at the neural level were associated with increased risk of psychopathology more generally (see 3.1.3.).

2.2.4 Autobiographical Memory Test (AMT) and fMRI Stimulus Generation. The AMT (Williams & Broadbent, 1986), a standard measure of OGM, was administered to all participants in a session 1-3 weeks before scanning at baseline. Participants generated specific memories in response to ten positive (e.g. achieve, caring) and ten negative cue words (e.g. mistake, lonely). OGMs were defined as ‘memories that did not contain at least one specific detail that identifies an event as a distinct episode’ (Valentino, Toth & Cicchetti, 2009; Williams & Broadbent, 1986). Correlational analyses showed that positive and negative OGM’s were correlated in each group (MT: \( r_{OGMpos\_neg} = .50, p = .004 \); Non-MT: \( r_{OGMpos\_neg} = .65 \).
Correlations between .30 and .70 are considered as ‘moderate’ and we therefore collated performance across conditions into one overall measure of OGM (OGM Total) to reduce the amount of comparisons. Stimuli for the fMRI task included specific memories generated from the AMT and specific memories generated from supplemental valenced cue words, administered to ensure a full set of twenty specific memories were generated (10 positive ABMs, 10 negative ABMs) for the imaging component, in line with prior fMRI studies (Summerfield, Hassabis & Maguire, 2009). These memories were rephrased by the participants in 3-4 words to prompt recall during the scanning session. In line with prior studies in adults (Summerfield et al., 2009), a control task was employed in which participants were presented with pictures and brief descriptions of ten everyday objects. Participants were given 20 seconds to memorise each object. As with the ABMs, key features were captured in brief sentence cues for presentation during fMRI. The memories were coded by a trained member of the research team in the lab (blinded) and double rated by a second, independent member of staff on 10% of the memories. Inter-rater agreement was high (average agreement $k>.82$).

### 2.3 MRI Scanning Session

There were three memory conditions: Positive ABM recall, Negative ABM recall and Object recall. The fMRI task consisted of 2 runs of 15 trials, each comprising 5 positive ABMs, 5 negative ABMs and 5 object memories presented in a pseudo-randomized order, which prevented the consecutive presentation of a stimulus category. Each trial began with a sentence cue and elaborate phase lasting 20 seconds. Participants indicated successful or unsuccessful memory retrieval (silent) via a self-paced button press. The text cue then disappeared and the word “Elaborate” was presented for 8 seconds during which time the participants recalled the
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memory or object. The elaboration period was followed by an audio tone (1 sec), then a 5-point Likert scale rating recall difficulty, vividness and remoteness of memory. The two rating periods lasted 4.5 seconds each and were followed by a 2 second inter-trial interval (ITI). Post-scanning, ABM’s were rated on emotional salience, remoteness and agency. These ratings were used to ensure the phases were comparable across participants. Analyses of vividness, recall difficulty and remoteness did not yield significant differences between the groups, confirming that cue phrases and corresponding memories were comparable between groups (all Ps > .19). Average recall success of cued ABMs was very high, with no significant differences in number of non-retrieved memories between the groups (MT: 1.75 ± 1.52 memories, Non-MT: 1.97 ±1.17 memories, p=0.62).

2.4 fMRI Data Acquisition

Data were acquired on a 1.5 tesla Siemens (Siemens Medical Systems, Erlangen, Germany) Avanto MRI scanner with a 32-channel head coil during 2 runs of approximately 9 minutes each. During each run, a total of 181 T2* weighted echo-planar (EPI) volumes were acquired, covering the whole brain with the following acquisition parameters: slice thickness: 2mm; TR: 85ms; TE: 50ms; FOV: 192 mm x 192 mm2; 35 slices per volume, gap between slices: 1mm; flip angle: 90°). A high-resolution, three-dimensional T1-weighted structural scan was acquired with a magnetization prepared rapid gradient echo sequence. Imaging parameters were: 176 slices; slice thickness = 1 mm; gap between slices = 0.5 mm; echo time = 2730 msec; repetition time = 3.57 msec; field of view = 256 mm x 256mm2; matrix size = 256 x 256; voxel size = 1 x 1 x 1 mm resolution.
2.5 Analyses

2.5.1. Over-general memory (OGM) as a behavioural predictor

Hierarchical linear regression models were conducted to predict follow-up depressive symptoms (CASI) and prosocial behaviour (SDQ) using the baseline total OGM score from the autobiographical memory test as the independent variable, controlling for baseline depression symptoms and baseline prosocial behaviour by entering baseline symptoms into the model as a covariate in SPSS version 24 (IBM Corp. 2012). We conducted an a priori power calculation based on our planned sample size and desired power (80% at $\alpha_p = .05$) to show that with 20 and 17 subjects respectively in each group we had 80% power to detect a ‘medium’ effect size of $d = 0.52$ at $\alpha = .05$ (one-tailed) in any of our behavioural measures, an effect size commensurate with effect sizes typically reported in this field, indicating sufficient power.

2.5.2 Brain-behaviour relationships

2.5.2.1 Pre-processing

Data analyses were conducted using the software package SPM8 (www.fil.ion.ucl.ac.uk/spm/software/spm8) implemented in Matlab 2015a. After discarding the first 3 volumes of each run to allow for T1 equilibration effects, data were realigned; initially within each run and then across the two runs to the first image of the first run. Data were normalized into MNI space using deformation fields from T1 scan segmentation at a voxel size of 3x3x3mm. The resulting images were smoothed with a 6mm Gaussian filter and high-pass filtered at 128Hz. Fixed-effects statistics for each individual were calculated by convolving
boxcar functions modelling the 4 conditions (Positive ABM, Negative ABM, Object recall and Rating) with a canonical hemodynamic response function (HRF).

Additionally, the six motion parameters were added to the model as regressors of no interest. In order to further minimize movement-related artefacts, images corrupted due to head motion greater than 1.5mm were removed and replaced by interpolations of adjacent images (<7% of each participant’s data). For these participants, we included an additional regressor of no interest to model the interpolated scans. Individual participants’ SPMs containing the parameter estimates of the 4 conditions were then entered as fixed-effects factors into a repeated measures mixed-effects ANOVA containing a ‘subject’ factor for random effects for group analyses.

2.5.2.2 fMRI Analyses

Contrast estimates were extracted from the peak voxels of clusters showing significant group differences in the contrast positive ABM recall vs. negative ABM recall in the previous study at baseline. These regional clusters were (i) the right middle temporal gyrus, (ii) right hippocampus and (iii) right amygdala and were extracted with MarsBaR Toolbox (Brett et al., 2004). Except for the amygdala, all parameter estimates were retrieved from whole brain analyses that were corrected using cluster-size thresholding (Monte-Carlo Simulation via 3D ClusterSim) (Ward, 2000) of voxel-wise \( p < 0.005, ke=75 \), which corresponds to \( p=0.05 \), family-wise error (FWE). The amygdala was retrieved via ROI analyses (anatomically defined using the Automated Anatomical Labelling (AAL) atlas as implemented on WFU-PickAtlas software) and small volume corrected for multiple comparisons at \( p<0.05 \). The data from the subset of participants who came back for a second assessment and are included in the present study mirrored the data from the larger set of participants included at the first timepoint with
regard to the hippocampus ($p=.006$), MTG ($p=.005$), but failed to reach significance in the amygdala ($p=.21$).

As with OGM score, hierarchical linear regression models were conducted to predict follow-up depressive symptoms (CASI) and prosocial behaviour (SDQ) using the baseline ABM-related brain activity in the: (i) right hippocampus, (ii) right MTG and (iii) right amygdala as independent variables. All regression models controlled for baseline depression symptoms and baseline prosocial behaviour by entering baseline symptoms into the model as a covariate. Analyses were conducted using SPSS version 24 (IBM Corp. 2012).

3. Results

Table A presents demographic data at baseline and follow up.

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3.1. Over-general memory (OGM) as a behavioural predictor of depressive symptoms and prosocial behaviour

3.1.1. OGM and depressive symptoms
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Relationships between OGM at baseline and depressive symptoms at follow-up were assessed by means of a linear regression model using the depressive symptom score at follow-up as the dependent variable and OGM as the predictor while controlling for levels of depressive symptoms at baseline. Baseline OGM was not found to be associated with elevated depressive symptoms at follow-up in the MT group ($\beta=.65$, $p=.15$; total model: $R^2=.22$, $F(2,16)=.81$, $p=.51$), when correcting for severity of maltreatment and baseline levels of depressive symptoms. In the NMT group, the only significant predictor of depressive symptoms at follow-up was level of depressive symptoms at baseline ($\beta=-.54$, $p=.012$; Total Model: $R^2=.38$, $F(2,19)=5.41$, $p=.015$), whilst OGM was non-significant ($\beta=.23$, $p=.25$).

3.1.2. OGM and prosocial behaviour

Hierarchical linear regression analyses were conducted in each group to investigate the effect of OGM at baseline on indices of prosocial behaviour (derived from the SDQ subscale). In the MT group, OGM significantly predicted prosocial behaviour ($R^2=.26$, $F(2,19)=6.2$, $p=.023$), controlling for severity of maltreatment and prosocial behaviour at baseline. Higher OGM was significantly associated with reduced prosocial behaviour measured at follow-up ($\beta=-.520$, $p=.023$; Bonferroni corrected). No relationship between OGM and prosocial behaviour was found in the non-MT group (OGM: $R^2=.01$, $F(2,16)=.04$, $p=.96$).

Analyses of the slopes of prosocial behaviour and depressive symptoms yielded no significant differences between the groups ($t_{prosocial}(33)=.19$; $t_{depressive}(26)=.60$).

3.1.3. OGM and symptoms of attention deficit hyperactivity disorder (control analyses)
In order to test if the association between OGM and socio-emotional functioning (depressive symptoms and prosocial behaviour) is specific we ran control analyses testing the association between OGM and symptoms typically associated with maltreatment experiences, i.e. hyperactivity (SDQ subscale) and inattention (CASI-subscale). OGM did not predict symptoms of attention-deficit-hyperactivity disorder (Hyperactivity: $R^2=.01$, $F(2,19)=.32$, $p=.72$; Inattention: $R^2=.11$, $F(2,16)=.77$, $p=.55$).

### 3.2. Atypical brain activity associated with specific ABM recall as a predictor of depressive symptoms and prosocial behaviour

#### 3.2.1. Hippocampus / MTG / amygdala activity during ABM recall (positive versus negative contrast) and depressive symptoms

No significant relationship was found between activity in either hippocampus, MTG or amygdala during positive versus negative ABM recall at baseline and depressive symptoms at follow-up in either group (all $P$s>.56).

#### 3.2.2. Hippocampus / MTG / amygdala activity during ABM recall (positive versus negative contrast) and prosocial behaviour

Controlling for prosocial behaviour at baseline, we did not find a significant relationship between activity in the hippocampus, MTG or amygdala during positive versus negative ABM recall at baseline and prosocial behaviour at follow-up in either group (all $P$s>.26).
4. Discussion

Here we investigated whether levels of OGM and neural indices of autobiographical memory functioning predicted future socio-affective functioning in adolescents with documented experiences of maltreatment. This is the first study to examine autobiographical memory functioning as a candidate mechanism for understanding future psychiatric vulnerability in individuals who have experienced childhood maltreatment. Greater levels of OGM at baseline were found to significantly predict lower levels of prosocial behaviour two years later in children who had experienced maltreatment. A similar pattern was also evident in relation to future depression symptoms but only at trend level. Analyses indicated that altered patterns of neural functioning associated with maltreatment experience (hippocampus, MTG, amygdala) were not related to either depressive symptoms or prosocial behaviour two years later, at least using the measures employed in this study.

Autobiographical memory plays a crucial role in social functioning (Alea & Bluck, 2003) and may serve to strengthen social relationships through the process of sharing past experiences with one another (Bauer, Stennes & Haight, 2003). Specifically, a pattern of overgeneral autobiographical memory may lead to impaired social problem-solving (Gino & Desai, 2012; Goddard et al., 1996). Prior studies have shown that individuals with maltreatment experience are more likely to present with social difficulties, particularly increased risk of re-victimization (such as bullying) (Benedini, Fagan & Gibson, 2016; Widom, Czaja & Dutton, 2008) and reduced levels of social support in adulthood (Sperry & Widom, 2013) Here, we found that adolescents with greater overgeneral memory presented with less prosocial behaviour two years later, suggesting that OGM may be one factor that impacts the ability to engage in adaptive social behaviour.
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Given this impact on social functioning, overgeneral autobiographical may be understood to increase latent vulnerability to future psychiatric disorder in two ways. First, it might have a ‘direct’ effect impacting on online social processing, increasing the likelihood of stressful events occurring. If an individual is less able to draw on past experience to mitigate potential problems through adaptive social problem solving, then these problems may be more likely to escalate into a crisis (e.g. relationship breakdown), a phenomenon that has been referred to as ‘stress generation’ (McCrory et al., 2019). Second, it might have an ‘indirect’ effect, cumulatively impacting an individual’s social ecology, as poorer adaptive social problem solving may reduce the likelihood that they can effectively elicit, cultivate and sustain protective social relationships. If an individual is less able to build a stable network that provides social support they will be more susceptible and vulnerable to stress.

Our analyses were inconclusive with regard to the overgeneral memory as a latent vulnerability factor for depression in adolescents with maltreatment histories. A relationship between OGM and depression has been observed in samples of depressed children and adults (Brittlebank et al., 1993; Peeters, Wessel, Merckelbach & Boon-Vermeeren, 2002; Vrielynck, Deplus & Philippot, 2007). However, more recent research provides a more nuanced picture and show that prospective associations between OGM and psychopathology emerged only among adolescents ‘at risk’, i.e. who reported elevated, and increasing patterns of rumination over time, whilst overgeneral memory did not predict psychopathology in adolescents without rumination (Gutenbrunner et al., 2018). Furthermore, some studies seem to suggest that the relationship between OGM and depressive symptoms is valence dependent, sometimes emerging to negative cue words compared to positive (Woody et al., 2015 children of mothers with major depressive disorder) and sometimes emerging to positive but not negative cues (Hipwell et al. 2011). The finding that OGM is evident even in individuals where depression
Autobiographical memory and latent vulnerability has remitted and predicts future depression in at-risk individuals suggests that OGM is not simply a state-like marker of depression, but a trait-like marker that is implicated in the pathogenesis of the disorder (Medsker, Forno, Simha, Juan & Sciences, 2016; Rawal & Rice, 2012). The nature of this association remains unclear. One possibility is that an increase in depression symptomatology is driven by poorer social problem solving that can result from OGM (McCrorry et al., 2017). OGM may also increase vulnerability to depression by limiting access to strategies to reduce negative affect that have previously been successful (Williams et al., 1996). Studies with larger samples and longer follow-up periods are needed to conclusively evaluate the potential role of OGM as a latent vulnerability factor for depression in children with experiences of childhood maltreatment.

Finally, in contrast to the significant findings observed for the behavioural index of OGM, no such relationship was found between the neural correlates of ABM and subsequent psychological and behavioural functioning. It is important to note that the brain activation was associated with a task that required retrieval of specific memories, and therefore did not index the neural correlates of OGM. Rather, it elucidated differences in how everyday positive and negative memories engaged a neural network of regions implicated in memory specification and salience. There are several possible explanations for this finding. First, while fMRI may be effective in indexing reliable group differences, different fMRI paradigms are likely to vary in how robust they are for reliably indexing individual differences (Nord, Valton, Wood & Roiser, 2017) and it may be that the signal generated by each individual in this task might not be sufficiently reliable to predict psychopathology. A second related possibility is that what was measured by fMRI (which related to affective content of successfully retrieved detailed memories) did not capture the aspect of information processing that indexes risk of reduced prosocial functioning and increased depression.
A number of limitations of the current study should be noted. First, due to the attrition over the two years in our sample, our sample was potentially not large enough to detect more subtle effects, such as group differences in the amygdala or sex differences. Previous research indicates that there may be some differential mental health outcomes for boys and girls exposed to early adversity and also that autobiographical memory processes can be different between the sexes (Reese, Haden & Fivush, 1996). It is important to note that power limitations arising from the smaller sample potentially influenced our ability to detect an effect for OGM and neural ABM processing on the development of depression. However, albeit smaller, our follow-up sample was comparable in terms of demographics, psychopathology and brain activation (except amygdala) to the full baseline sample. Second, a longer follow-up after the mean age indexed here (14 years old) would increase the likelihood of symptom development and sample size, which would increase power to detect associations with the development of mental health problems. Finally, in this study we used prosocial behaviour as a proxy measure for social functioning. In order to characterise more fully the impact of OGM on social behaviour it will be helpful for future studies to include other measures, including, for example, a measure of social problem solving and Theory of Mind.

The current study is the first to investigate whether alterations in neurocognitive processing shown to be associated with maltreatment experience predict future psychosocial functioning and mental health. We have shown that altered autobiographical memory processing in the form of increased OGM predicts reduced prosocial behaviour later in development, and may also predate risk for depression, in individuals with maltreatment histories. OGM may therefore represent a potential marker of latent vulnerability to poorer psychosocial functioning and mental health. The current findings highlight the potential importance of OGM as a potential therapeutic target for a preventative approach that would aim
to reduce risk of decreased pro-social behaviour and depression emerging in adolescents with prior histories of maltreatment. Such an approach would need to be complemented by a careful understanding of how OGM memory impacts social functioning in ways that potentiate risk.
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