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# **Linking Major Depression and The Neural Substrates of Associative Processing**

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### **Abstract**

It has been proposed that mood correlates with the breadth of associative thinking. Here we set to test this hypothesis in healthy and depressed individuals. Generating contextual associations engages a network of cortical regions including the parahippocampal cortex (PHC), retrosplenial complex, and medial prefrontal cortex. The link between mood, associative processing, and its underlying cortical infrastructure provides a promising avenue for elucidating the mechanisms underlying cognitive impairments in Major Depressive Disorder (MDD). Participants included fifteen non-medicated individuals with acute Major Depressive Episode and fifteen healthy matched controls. In an fMRI experiment, participants viewed images of objects that are either strongly or weakly associated with a specific context (e.g., a beach chair vs. a water bottle), while rating the commonality of each object. Analyses were performed to examine brain activation and structural differences between groups. Consistent with our hypothesis, controls showed greater activation of the contextual associations network compared with depressed participants. In addition, PHC structural volume was correlated with ruminative tendency and, volumes of the hippocampal subfields were significantly smaller in depressed participants. Surprisingly, depressed participants showed increased activity in the entorhinal cortex (ERC) compared with controls. We integrate these findings within a mechanistic account linking mood and associative thinking, and suggest directions for the future.

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Disclosures

# Introduction

The notion that our thinking is associative in nature has long been discussed and studied (Barsalou, 1999; Moores, Laiti, & Chelazzi, 2003). Bridging separate concepts through associations is the basis of cognition, via processes such as learning, memory, and the progression of thought. In a series of studies, Bar and colleagues have found evidence for a network of cortical regions that mediate contextual associative processing (Bar & Aminoff, 2003; Bar, 2007; 2009a). This network is active when subjects are presented with visual images of objects that are strongly associated with a specific context (e.g., a bowling pin) relative to objects that are equally common in our environment but are not strongly associated with any specific context (e.g., a fly). The brain regions involved in this contextual processing network include the retrosplenial complex (RSC), the parahippocampal cortex (PHC), and the medial prefrontal cortex (MPFC) (Aminoff, Schacter, & Bar, 2008; Bar, 2003; 2004). It was proposed that the RSC activates the gist of a context frame, while the PHC activates the specific perceptual features and associated items contained within it (Aminoff et al., 2008). The MPFC's suggested function is to generate expectations and predictions about what is going to occur in the immediate environment, based on the information activated within the context frame (Bar, 2004; Bar, Aminoff, Mason, & Fenske, 2007; Herbst, Kveraga, & Bar, Submitted).

It has been hypothesized (Bar & Ullman, 1996; Mandler, 1976; Minsky, 1975; Schank, 1975) that context frames facilitate retrieval of information, whereby top-down information (e.g., context) or bottom-up information (e.g., low spatial frequencies; Oliva & Torralba, 2001; Bar, 2004) about the perceived stimulus can activate associated information within the context frame, increasing expectancy for specific stimuli. Thus, the availability of internally generated, associated information accompanying a stimulus impacts subsequent processing of information.

Mood is another construct that has been documented to influence the encoding and retrieval of associated items. Notably, studies indicate that positive mood broadens the scope of associations and loosens the conceptual relations between associations (Isen & Daubman, 1984; Isen, Johnson, Mertz, & Robinson, 1985). Positive affective states do not only evoke the generation of more unique, broadly related associates in free association tasks, but they also lead to more remote categorical associations (e.g., surfboard = a vehicle) (Isen & Daubman, 1984; Rosch & Mervis, 1975) and promote more creative problem solving, as measured, for example, by facilitated performance on the candle task (Duncker, 1945) and Remote Associates Test (Isen, Daubman, & Nowicki, 1987; Mednick, Mednick, & Jung, 1964; Mednick, Mednick, & Mednick, 1964). Recently, it has been proposed (Bar, 2009a) and later demonstrated that the reverse is also true: increasing the breadth of associations can positively influence mood, indicating that the mood-associative processing relationship may be bidirectional (Mason & Bar, 2012), which could potentially be used as method for alleviating mood. In addition, it was found that mental load directly influences the breadth of associations provided in a free-association task (Baror & Bar, 2016), and as such it was suggested that ruminative thinking can be seen as mental load that accordingly limits associative scope.

Mood disorder patients display characteristic relationships between mood and thought that would be expected from these findings. For instance, patients in hypo-manic or manic states, characterized by elevated mood, often exhibit a loosening of associative links between concepts, which can lead to 'flight of ideas' (Andreasen, 1979; Lake, 2008; Sass & Pienkos, 2013) and increases in creativity (Davis, 2009). In more severe manic states, where elevation of mood is accentuated, the extreme loosening of associations may lead to incoherence of thought (Andreasen 1979; Lake, 2008; Sass & Pienkos, 2013). Conversely, depressed mood and anhedonia – the major features of Major Depressive Disorder (MDD) – are characterized by ruminative thought pattern that is repetitive in nature and revolves around constrained and narrow themes (Andreasen, 1979; Lake, 2008; Nolen-Hoeksema, 2000; Sass & Pienkos, 2013).

Much of the previous experimental research investigating mood and associative thought has employed mood induction techniques in healthy subjects (Ferrer, Grenen, & Taber, 2015), rather than focusing on psychiatric disorders, making the applicability of these findings to clinical populations unclear. Psychiatric disorders are characterized by more long-term and severe levels of emotional distress than could be simulated in a mood induction study. Investigating clinical populations that are characterized by specific affective states allows us to elucidate the neural networks related to changes in associative thinking and, furthermore, whether these networks are related to functional and morphological changes in the identified contextual associative network. By focusing on differences within this network, research could create specific targets for future treatments. For example, Mason & Bar (2012), found that experimentally increasing the breadth of associations may lead to improved mood in healthy individuals. If differences in the contextual associative network are associated with the thought patterns characteristic of mood disorders, regulating this pathway in clinical disorders may be a fruitful therapeutic goal.

Given the proposed link between mood and associativity (Bar, 2009a), we hypothesized that contextual associative processing would be compromised in MDD patients. To test this hypothesis, we used functional Magnetic Resonance Imaging (fMRI) to measure brain activity during contextual associative processing of objects presented visually, and compared brain activation between patients diagnosed with Major Depressive Disorder (MDD) and healthy participants. We predicted that MDD patients would display reduced activity in the contextual associative network when processing contextual information, compared with healthy controls. Building on previous studies demonstrating that morphological differences between MDD patients and healthy controls are often related to behavioral differences (e.g. Guo, Gatchel, & Sahay, 2015), we also predicted that regions connected to contextual associative processing would show differences in grey matter volume. Further, because of its explicit connection to both MDD and associative thinking, we also predicted that ruminative tendency would be associated with differences of functional activation within the contextual associations network, as well as with differences in grey matter volume. Lastly, we predicted that our depressed participants would display selective decreases in hippocampal subfield volumes in accordance with previous studies showing reduced hippocampal subfields volumes in MDD, specifically the dentate gyrus (DG), and data showing direct links between associations and the hippocampal complex (Mayes, Montaldi, & Migo, 2007; Travis et al., 2015; Treadway et al., 2015).

The DG subfield of the hippocampus is specifically responsible for transforming similar memories and patterns into distinct mental representations (pattern separation) (Guo et al., 2015), which may play a role in connecting associated contextual representations. Therefore, we predicted that we would find decreased DG volume in MDD patients if they displayed our predicted activation differences in the contextual associative network. Furthermore, the DG has been linked with adult neurogenesis (Cameron & Gould, 1994), which has been suggested to be reduced in mood disorders (Treadway et al., 2015) and to be up-regulated with remission of symptoms mediated by SSRIs or psychotherapy (Guo et al., 2015). Although the link between associations, DG and depression-related reduction in neurogenesis is speculative, showing such a link empirically here will provide critical support for this hypothesis.

## **Methods & Materials**

# Subjects

Fifteen adults diagnosed with MDD and fifteen healthy control participants were included in this study. Depressed subjects were recruited through the Depression Clinical and Research Program (DCRP) of Massachusetts General Hospital and through an advertisement in the volunteer section of Craigslist. All subjects went through a phone screening procedure and whoever appeared to meet the inclusion and exclusion criteria of the phone screening was scheduled for a full psychiatric SCID interview by a trained psychiatrist (EVH). All subjects provided informed consent in writing. The protocol was approved by the research ethics committee of Massachusetts General Hospital. All the included depressed participants met criteria for a DSM-IV diagnosis of current Major Depressive Episode (MDE) based on the Structured Clinical Interview for DSM-IV Disorders. Depressed individuals with psychotic features or participants taking psychotropic medications at the time or 4 weeks prior to the study or who met criteria for a current, co-morbid diagnosis of any Axis I disorder, with the exception of social anxiety disorder, were not included in the study. Participants were evaluated using a clinician rated Hamilton rating scale of Depression (HRSD), a 17-item scale for evaluating the severity of depressive symptoms, as well as the Quick Inventory of Depressive Symptoms- Self-Rated questionnaire (QIDS-SR) and the Ruminative Response Style (RRS) questionnaire (Treynor, Gonzalez, & Nolen-Hoeksema, 2003). The RRS is a 22-item, self-report measure of self-focused rumination about depressive mood, its causes, and consequences.

A group of healthy control participants comparable in age, sex, and education was recruited for this study using an advertisement in the volunteer section of Craigslist. Participants were excluded if they had any lifetime Axis I Disorder or were suffering from any medical condition, acute or chronic. The Demographic data collected is depicted in Table I. After complete description of the study to the subjects, written informed consent was obtained.

For the functional analysis we removed runs in which movement exceeded 2 mm at any direction. 4 runs from 3 subjects were excluded in the depressed group while 2 runs from 2 subjects were excluded in the healthy control group.

# Task Design

The paradigm (see Figure 1) employed a rapid event-related design. Target images were color photographs (256×256 pixels) of objects (69 strongly contextually associated, 69 weakly contextually associated) in isolation presented on a white background. Each target image was presented briefly (150ms), and immediately followed by a colorful mask presented for 100ms. A red fixation cross then appeared signaling the start of the response period and turned black after 1500ms, signaling the end of the response period. The black fixation-cross remained on the screen for the duration of the inter-trial interval (ITI), which ranged from 200ms to 9250ms (to allow a jittered ITI in multiples of TR length (2200 ms) and jittered stimulus presentation from the start of each TR). The fMRI session consisted of 138 unique trials, pseudo-randomly ordered across the three functional runs, in addition to 28 practice trials that were completed prior to the main task (practice trials presented images not included in the main task). Each target image was only presented once within each session.

Subjects were instructed to rate how common each object was on a three-point scale ranging from not common to more common (for example a car would be rated common whereas boxing gloves would be rated less common). This task was chosen because it requires high-level object recognition without focusing attention explicitly on perceptual features or on the associative qualities of the object, an assumption we later verified by comparing ratings of commonality across image conditions. Responses were provided on a three-button MR-compatible response box. The order of the three-point scale was counterbalanced across subjects to prevent confounds between rating and motor mapping, and subjects practiced the appropriate mapping to proficiency before the practice trials began. Stimulus presentation and response collection was performed using Psychtoolbox (http://www.psychtoolbox.org) running on Matlab software (http://www.mathworks.com), controlled by a MacBook Pro laptop with a monitor resolution of 1024×768 and refresh rate of 60 Hz.

Images of strong and weak associative objects were compiled from a set previously normed and used in studies of contextual processing ((Bar & Aminoff, 2003); available at https://faculty.biu.ac.il/~barlab/context\_localizer.html).

# Image acquisition

Images were acquired using a Siemens 3T Trio Tim MR magnet and a 32-channel RF head coil. We acquired functional image volumes as T2\*-weighted echo-planar images (EPIs) with the following parameters: 36 interleaved slices, 2200ms TR, 28ms TE, 2.5mm thickness, 0.75mm gap, 64×64 matrix, 200mm FOV (resulting in an in-plane voxel size of  $3.125 \times 3.125 \times 2.5$ mm). Our fMRI sequence and slice prescription was optimized for reducing signal loss and distortion in the orbitofrontal cortex (based on recommendations of Deichmann, Gottfried, Hutton, & Turner, 2003; Deichmann, Josephs, Hutton, Corfield, & Turner, 2002; Weiskopf, Hutton, Josephs, & Deichmann, 2006), including the use of a modified z-shim pre-pulse moment and 30° tilt of our slice prescription counterclockwise of the AC/PC line along the sagittal plane (Deichmann et al., 2003, 2002; Weiskopf et al., 2006). As a consequence of the limited slice prescription used in order to achieve optimal MPFC signal, the most dorsal portions of posterior parietal cortex were not captured in the

scan volume for a majority of subjects. Each subject performed 3 functional runs, each consisting of 97 TRs. Each run included 11s of fixation at the beginning (to allow for the fMRI signal to reach steady-state), and the corresponding 5 EPI volumes were discarded from further analysis. Each session included the acquisition of two high-resolution T1-weighted Multi-Echo MPRAGE (MEMPRAGE) anatomical images (1mm isotropic voxels), which were later averaged together.

#### fMRI Analysis

Structural and functional imaging analyses were performed using the Freesurfer and FS-FAST analysis tools and processing stream developed at the Martinos Center for Biomedical Imaging (http://surfer.nmr.mgh.harvard.edu) and R statistical analysis software version 4.12.1 (R Development Core Team, 2011). Cortical reconstruction and volumetric segmentation of the T1-weighted scans were performed with Freesurfer 5.3 (http://surfer.nmr.mgh.harvard.edu/fswiki). Functional imaging data was preprocessed using the Freesurfer Functional Analysis Stream (FSFAST), version 5.1 (http://surfer.nmr.mgh.harvard.edu/fswiki/FsFast). The images were motion corrected (AFNI 3dvolreg; http://afni.nimh.nih.gov), slice timing corrected to the middle of the TR of a volume, intensity normalized, and registered to the same-subject anatomical volume. Each 4D functional dataset was resampled to "common space" using the surface-based intersubject registration created during the cortical reconstruction of the cortical mantle that occurred as a portion of the Freesurfer processing pipeline (this brings the left and right cortical hemispheres into the average space of Freesurfer; fsaverage). Finally, 5 mm FWHM smoothing was applied to each surface (2D surface-based smoothing)

A first-level general linear model (GLM) was set up consisting of two conditions/regressors of strong and weak context object processing. In addition to the task regressors and their temporal derivatives, estimated motion correction parameters and a set of polynomials (up to second degree) were included in the GLM as nuisance regressors. The first five volumes were removed from each fMRI run to allow for signal stabilization. The estimated hemodynamic response was defined by a gamma function of 2.25-s hemodynamic delay and 1.25-s dispersion. Data were then tested for statistical significance for each individual (first level), and contrast maps were constructed comparing the BOLD estimates for each condition. For second level analysis, both within-group (strong vs. weak context objects processing) and between-group (Depressed vs healthy control participants) whole-brain analyses were conducted.

A within group analysis in the strong versus weak objects contrast was done for the healthy control group and the depressed group using a voxel-wise threshold of p<0.001, extent-thresholded to achieve a whole-brain cluster-wise corrected for multiple comparisons at p<0.01. This was done in order to replicate previous findings of activation of the context network. Further, a between-group comparison (Depressed vs controls) of the strong versus weak contrast was performed using a voxel-wise threshold of p<0.05 and cluster wise threshold of p<0.05. Correction for multiple comparisons was done for the left hemisphere cortical surface. Lastly, analyses testing the associations between percent signal change

functional activation and rumination scores as measured by the RRS questionnaire were also performed.

#### Structural Analysis

To investigate relationships between brain structure and MDD status, we conducted analysis of covariance (ANCOVA) with the volume of the region of interest (ROI) as the outcome variable and MDD status as the grouping variable. We included total brain volume, age, sex, and years of education as covariates. Furthermore, to investigate relationships between brain structure and ruminative thinking, we performed linear regressions with the volume of the ROI as the outcome variable, the participant's score on the RRS survey as the explanatory variable, and total brain volume and MDD status as a covariate. Regions of interest included coordinates displaying activational differences in the functional analyses, regions that have been previously connected to the context network, and the hippocampal subfields. Significance levels are adjusted for multiple comparisons where applicable. Specifically, p<0.001 for hippocampal subfields and p<0.01 for non-hippocampal subfield group comparisons. One depressed subject was excluded from structural analyses because of issues with auto-segmentation and volume calculation.

#### Results

#### fMRI results

In the healthy control group (n=15), the contrast between strong and weak context objects (contextual processing), using whole brain analysis, showed cluster activations with voxelwise threshold set to p<0.001 (Figure 2) and clusterwise threshold set to p<0.01in the RSC [comprised of the precuneus (-7.5, -53.4, 20.5), fusiform (-34.2 - 42.1 - 10.9), isthmus cingulate (-4.7, -33.2, 31.4)], PHC (-30.7, -40.2, -5.8), and MPFC (-7.1, 50.7, -8.1) (Figure 2a). These results replicate previous findings in healthy participants during contextual processing (Bar & Aminoff, 2003; Bar, Aminoff, & Schacter, 2008).

In the group of depressed patients, the contrast of contextual processing using whole brain analysis, showed cluster activations with voxel wise threshold set to p<0.001 and cluster wise threshold of p<0.01 for the medial frontal gyrus (MFG) of the dorsolateral prefrontal cortex (DLPFC) (-20.9, 57.1, 5.4) and the ERC (-20.5, -18.1, -20.6) (Figure 2b).

A group comparison using a voxelwise threshold of p<0.05 and cluster wise threshold of p<0.05 corrected for multiple comparisons for left hemisphere cortical surface (Depressed vs. healthy control participants) in the contrast of strong vs. weak context objects revealed differential activity in 3 regions when correcting: the MFG, PHC and ERC (figure 3). A cluster in the left PHC (-18.3, -34.2, -9.6) that showed greater activation (P<0.05) in the healthy control group compared with the depressed group and cluster in the ERC (-19.8, -14.5, -23.6) and MFG (-33.6, 36.0, 23.4) that showed greater activity (p<0.05) in the depressed group compared with the healthy control group (Figure 3a). Percent signal change metrics from functional clusters of differential activity were extracted for each subject in each condition (Figure 3b).

Analyses testing the association between rumination and the contrast of contextual processing failed to find any significant associations. Lastly, the behavioral component of our task, reaction time of commonality ratings and the commonality ratings, did not significantly differ between healthy controls and depressed patients.

## **Structural Results**

A group comparison (Total, n=29; MDD subjects, n=14) did not provide evidence of differences in volumes of the coordinates showing activation differences (ERC: F=0.02, p>0.10; PHC: F=0.13, P>0.10; MFG: F=0.30, P>0.10). However, there was significant association between rumination score and the gray matter volume of the coordinates located in the PHC (Fig. 4; t-value: 2.93; p-value: 0.007). After adjusting for total brain volume and MDD status, an increase of one point on the RRS-RSQ scale is associated with an increase of 5.6mm<sup>3</sup> in PHC volume (SE: 1.92 mm<sup>3</sup>). Auto-segmented ERC volumes in the left hemisphere were found to be significantly smaller in depressed patients compared with matched controls (ERC: F=8.349, p=0.0008).

The total volume of the right hippocampus was significantly reduced in the MDD group (F=10.090, p=0.00421). Within the right hippocampus, the CA1 (F= 15.153, p=0.00073; Effect Size = 1.25(0.5, 1.99), CA2-3 (F=14.297, p=0.00097; Effect Size = 1.2(0.47, 1.94), and CA4-DG (15.637, p=0.00063; Effect Size = 1.25(0.51, 1.99)) were significantly reduced in the MDD patients compared to controls. The subiculum (F=12.013, p=0.0021) is trending towards significant volume reduction in MDD patients compared to controls. Neither the total left hippocampus nor its subfields showed a volume reduction significant or trending at p=0.001 (results summarized in Supplemental Information).

## **Discussion**

Previous studies indicate a direct relationship between mood and associative thinking, whereby negative mood constricts and positive mood broadens the breadth of associations. With this framework in mind, we hypothesized that contextual associative processing might be compromised in depression. To test this, we compared brain activity and gray matter volume of regions involved in the processing of contextual associations between depressed participants and a healthy control group. Consistent with our hypothesis, depressed patients displayed a pattern of activation and morphological characteristics that differed from the typical "context network" displayed in healthy participants.

Compared with healthy controls, depressed participants showed several activational differences in MTL and PFC regions: 1) A decreased differential activation in the PHC, 2) A greater differential activation in the ERC, and 3) A greater differential activation in the MFG (Figure 2). We did not find statistically significant group differences in other areas of the context network such as the RSC and the MPFC. Alongside these results, our morphological analyses revealed several findings: 1) PHC volume was positively associated with ruminative tendency, 2) Decreased left ERC volume in depressed subjects, and 3) Decreased volume in specific hippocampal subfields (CA1, CA2-3, CA4-DG) in depressed subjects.

According to previous accounts of contextual processing in healthy participants (Bar, 2003, 2007; Bar et al., 2008), the PHC plays a crucial role in object recognition by using information about the current context frame (e.g., bowling alley) to generate expected associated stimuli (e.g., bowling ball, bowling pin) (Bar, 2004; Bar & Ullman, 1996). Moreover, in regards to this framework, it has been shown that the RSC (Aminoff et al., 2008; Bar, 2007; Bar & Aminoff, 2003) contains contextual representations that may be more abstract and prototypical (i.e., context frames), while the PHC seems more directly related to the specific appearance and physical properties of a context and its associated elements (and thus might contain more episodic versions of context frames). Thus, our finding of impaired PHC activity in depressed subjects provides evidence for a possible impairment in the generation of contextual associates, which is in line with our overarching hypothesis.

It is also interesting that PHC volume was positively correlated with ruminative tendency as measured with the RRS questionnaire. Whether these volume differences can be interpreted as increased effort by this region to retrieve contextual associations is unclear. If so, then the possibility that this activation and morphological impairment may give rise to ruminative thought pattern is highly intriguing and is also in accord with our general hypothesis connecting negative mood and rumination to constricted associative thought.

The greater ERC activation in our MDD patients compared with controls was not predicted in advance. Nevertheless, when viewed alongside the decreased ERC volume in our MDD patients, this finding corroborates and enriches existing morphological data that point to ERC pathology in depression. For example, previous studies have found decreases in left ERC cortical thickness in depressed participants (Tu et al., 2012), reductions in left ERC volume in females with treatment resistant depression (Furtado, Maller, & Fitzgerald, 2008), an inverse correlation between the bilateral hippocampal-entorhinal volume and years since onset of depression in geriatric depressed subjects (Bell-McGinty et al., 2002), and an association between late onset depression and smaller ERC volume in subjects with atherosclerotic disease (Gerritsen et al., 2011).

A possible account of our findings may be that the increased ERC activation during contextual processing is related to the decreased activity in PHC. The ERC seems to act as a relay station between the prefrontal cortex and the hippocampus and has a role in novelty detection (Witter, Wouterlood, Naber, & Van Haeften, 2000). It is believed to function as a buffer that holds multimodal sensory information and compares it with internal representations to detect "familiarity" versus "novelty" (Arnold, 1999; Lörincz & Buzsáki, 2000), while the PHC had been proposed to make contextual associations available. The task instructions were to rate the commonality of the object in the environment, requiring a comparison of the object presented to similar representations in memory that are common to our environment. Perhaps in depression the lack of availability of such information fed to the hippocampus by the PHC induces this increased, compensatory activity in the ERC. These inter-relations between MTL structures should naturally be further investigated in order to validate these proposals. Our finding of increased ERC activity during contextual processing in depression and decreased activity in the PHC opens interesting new avenues both for

research and for potential therapy approaches. To our knowledge, this is the first such demonstration of functional abnormalities in the ERC in depression.

Elucidating the role of the ERC in the pathophysiology of depression is increasingly important because this region is among the first to show volume loss with preclinical Alzheimer's disease (AD) (Dickerson et al., 2001) and the ERC is a potential target area for intervention by brain stimulation techniques to improve memory and learning (Suthana et al., 2012). The link between AD and depression has been suggested by studies showing lifetime depression as a risk factor for AD (Dotson, Beydoun, & Zonderman, 2010; Green et al., 2003), that reversible memory impairment during depressive episode is specifically a risk factor for dementia (Sáez-Fonseca, Lee, & Walker, 2007), and that exposure to abnormal levels of psychosocial stress is a common pathway leading to both neuro-pathologies (Aznar & Knudsen, 2011). Given our findings and literature reviewed here, there likely exist interesting connections between the ERC and some characteristic hallmarks of MDD such as anhedonia and negative recall bias. However, we did not specifically focus upon these traits when preparing our instruments and task design, so connecting these specific traits to our findings here may be overly speculative. These questions, while intriguing, will need to be addressed by future studies with a nuanced focus on these traits.

The finding that the MFG is differentially more engaged during contextual processing in depressed patients raises the question as to its functional significance during this task. The MFG, corresponding to BA 46d of DLPFC, has been associated with cognitive control in numerous studies (Koechlin, Ody, & Kouneiher, 2003; Kouneiher, Charron, & Koechlin, 2009) and abnormal activity in the left DLPFC has been previously documented in imaging studies of depression compared with healthy individual. For example, measures of activity observed during rest, as measured with PET or fMRI, point toward a reduction of baseline activity in DLPFC (Mayberg, Lewis, Regenold, & Wagner, 1994; Rogers et al., 2004; Videbech et al., 2002). Additionally, behavioral studies of cognitive control tasks (e.g., working memory) typically associated with DLPFC activity show impairments in depression (Abas, Sahakian, & Levy, 1990; Austin et al., 1999; Beats, Sahakian, & Levy, 1996; Degl'Innocenti, Ågren, & Bäckman, 1998; Elliott et al., 1996; Grant, Thase, & Sweeney, 2001; Harvey et al., 2005; Sweeney, Kmiec, & Kupfer, 2000), suggesting that depressed patients have abnormally functioning DLPFC activity during tasks demanding cognitive control. While some neuroimaging studies of brain function in depressed patients during cognitive tasks have reported contradicting results (Harvey et al., 2005; Wagner et al., 2006; Walter, Wolf, Spitzer, & Vasic, 2007), most point to DLPFC differential hyperactivity in depressed subjects.

In the present study, we observed a similar higher activity of DLPFC, specifically when subjects were engaged in contextual processing, albeit an overall reduced activity in DLPFC as seen by percent change from baseline (Table 2). Most models of brain abnormalities in depression emphasize the importance of the limbic-DLPFC reciprocal interactions where limbic activity predominates DLPFC activity in depression (Mayberg et al., 1994). Whether this predominance is a result of a failed top-down inhibition by DLPFC or a reciprocal process originating in limbic areas and resulting in DLPFC hypo-activity is still not clear.

Our results displaying selective decreases in the volumes of the CA 1-3 and DG subfields of the hippocampus are particularly intriguing. Reduction in hippocampal volume is one of the most replicated neurobiological changes found in MDD (McKinnon, Yucel, Nazarov, & MacQueen, 2009), but comparatively few studies have investigated whether these volumetric reductions affect the hippocampus universally or if they are specific to certain subfields (Adam Samuels, Leonardo, & Hen, 2015). Hippocampal subfields are differentially activated for specific tasks and may be sensitive to specific environmental stimuli (Fanselow & Dong, 2010; Thompson et al., 2008; Zhao et al., 2001), so elucidating volumetric reductions of individual subfields may provide insight into the pathogenic mechanisms of MDD (Small, Schobel, Buxton, Witter, & Barnes, 2011). The CA1-3 and DG subfields are responsible for pattern completion and pattern separation, respectively (Guo et al., 2015), and they each express large numbers of receptors for stress-related hormones (e.g. glucocorticoids) (Small et al., 2011). Since MDD is characterized by increased, chronic exposure to stress, the experience of MDD may lead to impairments in the structure, and likely the function, of these subfields. If pattern completion and separation are impaired, individuals may have limited ability to integrate new, positive experiences into their thought patterns or form new, broader associations to break ruminative thought patterns. The first portion of this pathway has been supported by recent research displaying longitudinal decreases in the DG with increased depressive episodes (Treadway et al., 2015), but the second portion connecting the DG, CA, and associative thinking is more hypothetical still.

#### Limitations

The present study has several limitations. First, the sample size (15 participants in each group) is smaller than would be ideal. However, the MDD subjects are all non-medicated, removing the confounding effects of treatment on brain activation and morphology, which is a common limitation in studies of MDD. Second, the task used to assess contextual processing had no behavioral component apart from the commonality ratings of the objects and the reaction time for those ratings. Hence, the amount of engagement in the task cannot be measured. However, given our previous experience with this task, there is little doubt about participants' actual engagement and adherence. Furthermore, this limitation should be regarded keeping in mind the wide acceptance of the measurement of resting state activity and the conclusions drawn from such tasks about default network activity. In this sense, the task used in this study is much more constrained in measuring associative processing.

#### **Conclusions**

In summary, our fMRI results show impaired brain activity during processing of contextual information in acute non-medicated depressed individuals when compared with controls. Furthermore, we found that these individuals also have volumetric differences in regions connected to this contextual processing. Overall, these results suggest that the MTL structures of the PHC and the ERC are differentially affected by MDD, potentially causing ERC activity to increase in an attempt to increase recollection of contextual memory.

Our study implicates a connection between mood and contextual processing wherein the characteristic traits of MDD thought patterns, specifically ruminative tendency and narrow associative scope, are connected to measurable changes in both the activity and the structure

of the context network. Combined with existing psychiatric knowledge of the symptomatology of MDD, our findings point at potentially important targets both for the research and treatment of depressive disorders.

# **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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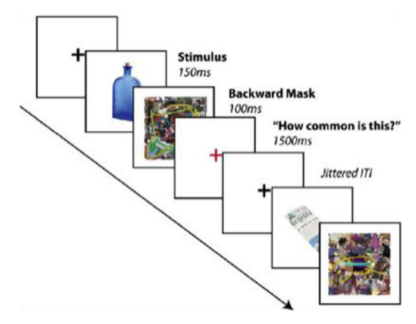
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**Figure 1.**Task timeline. Subjects viewed objects presented briefly and in isolation (150ms), followed by a colorful backward mask (100ms) and then rated how common the object was (1.5s response period). In between trials subjects viewed a black fixation cross for variable durations.

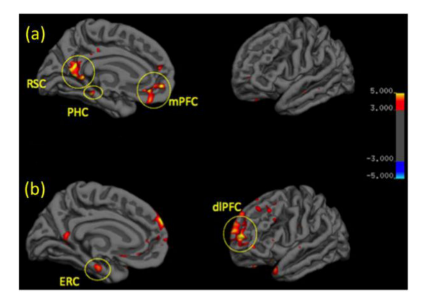


Figure 2. Regions significantly associated with contextual processing in (a) Healthy Controls and (b) Depressed patients (threshold p<0.001). RSC - retrosplenial cortex; PHC - parahippocampal cortex; mPFC - medial prefrontal cortex; ERC - entorhinal cortex; dlPFC - dorsolateral prefrontal cortex.

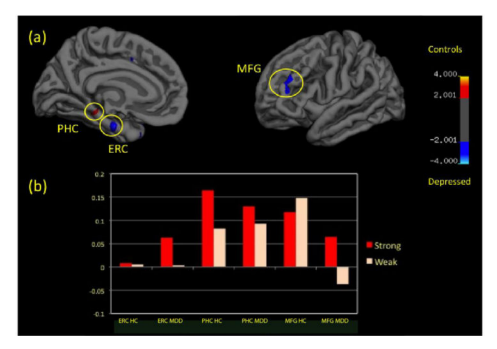
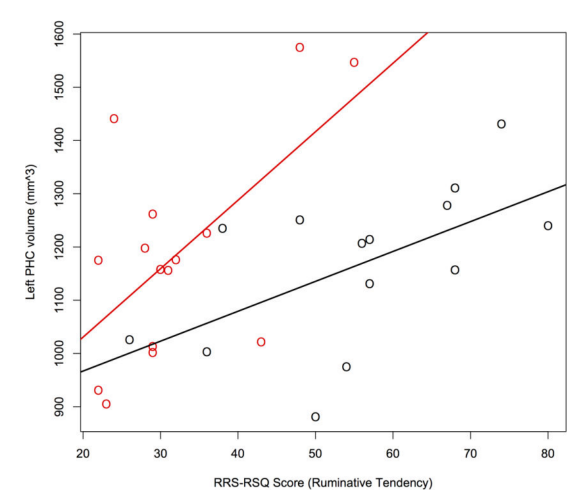


Figure 3. Compared to Healthy Controls (HC), Depressed Patients (MDD) display reduced parahippocampal cortex (PHC) but increased entorhinal cortex (ERC) and medial frontal gyrus (MFG) differential activity during contextual processing (p<0.01). (a) Displays the distribution of this differential activity and (b) displays the levels of activation (percent signal change) in these regions during the Strong (red) and Weak (white) contexts.



**Figure 4.** PHC volume is significantly associated with ruminative tendency (RRS-RSQ score) after adjusting for Total Brain Volume and MDD status ( $\beta$  = 5.6 (±1.92); p=0.007). (Red = healthy controls, Black = Depressed patients)

Table 1

# Sample demographics

	Depressed subjects	Healthy controls	
Age [years (SD)]	35.73 (9.87)	37 (14.13)	P=0.78
Sex (M/F)	7/8	8/7	/
Education [years (SD)]	14.9 (2.2)	16.26 (2.78)	P=0.14
QIDS score (SD)	22.1 (6.6)	2.7 (2.6)	
RSQ score (SD)	58.9 (13)	29.1 (6)	
HRSD (SD)	21.8 (4.7)	1.5 (1.2)	