Increased Amygdalar and Hippocampal Volumes in Young Adults with Social Anxiety

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Abstract

Background: Functional neuroimaging studies have consistently shown abnormal limbic activation patterns in socially anxious individuals, but structural data on the amygdala and hippocampus of these patients are scarce. This study explored the existence of structural differences in the whole brain, amygdala, and hippocampus of subjects with clinical and subthreshold social anxiety compared to healthy controls. We hypothesized that there would be volumetric differences across groups, without predicting their direction (i.e. enlargement or reduction).

Methods: Subjects classified as having social anxiety disorder (n = 12), subthreshold social anxiety (n = 12) and healthy controls (n = 14) underwent structural magnetic resonance imaging scans. The amygdala and hippocampus were defined a priori as regions of interest and volumes were calculated by manual tracing. Whole brain volume was calculated using voxel-based morphometry.

Results: The bilateral amygdala and left hippocampus were enlarged in socially anxious individuals relative to controls. The volume of the right hippocampus was enlarged in subthreshold social anxiety participants relative to controls. No differences were found across groups in respect to total brain volume.

Conclusions: Our results show amygdalar and hippocampal volume alterations in social anxiety, possibly associated with symptom severity. The time course of such alterations and the cellular and molecular bases of limbic plasticity in social anxiety should be further investigated.

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Introduction

Neuroimaging studies have identified a number of brain structures underlying the behavioral manifestations of social anxiety, including mainly the prefrontal and anterior cingulate cortices and limbic and paralimbic structures, with an emphasis on the amygdala [1].

There is reasonable consensus today that amygdala hyper-responsiveness is a core feature of social anxiety. The amygdala has been implicated in the acquisition of conditioned fear [2] and, together with the hippocampus, which is believed to process contextual cues, the amygdala is central to the classification of social stimuli as potentially threatening and to the elaboration of early responses to them.

Neuroimaging has provided consistent evidence concerning the function of limbicareas in social anxiety. Based on the vast literature describing functional alterations underlying social anxiety (for a review, please see Freitas-Ferrari et al., 2010 [1]), recent studies using resting-state magnetic resonance imaging (MRI) have described abnormal connectivity between the amygdala and areas associated with the processing of socially relevant information, including the orbitofrontal, prefrontal, and visual cortices [3,4].

Despite the abundance of functional studies, structural data concerning the same regions are scarce. An early investigation failed to find volumetric differences in the caudate, putamen, thalamus, and whole brain between patients with social anxiety disorder (SAD) and healthy controls [5]. Recently, however, Irle et al. (2010) [6] found decreased amygdalar and hippocampal volumes in men with generalized SAD relative to controls and Liao et al. (2011) [7] reported reduced gray matter volumes in the right hippocampus and inferior temporal gyrus in SAD, which were associated with enhanced resting-state functional connectivity. Finally, in the last and most recent article describing structural changes associated with social anxiety, Syal et al. (2012) [8] described cortical thickness reductions in areas surrounding the fusiform and post-central gyri and, specifically on the right hemisphere, in the frontal, temporal, parietal, and insular cortices. Their volumetric analyses, however, showed no differences...
between patients with SAD and healthy controls in respect to the volume of the amygdala and hippocampus.

The findings of increased activity and decreased volume in the amygdala and hippocampus in SAD are intriguing. Although the relationship between activity and volume of brain structures is not yet clear, some authors suggest that increased metabolic activity is likely to be associated with increased blood flow, which, in turn, might result in subtle volume increases [9]. Along the same line, Supékar et al. (2010) [10] proposed that increased gray matter volume could reflect enhanced synaptic connectivity. If this is true, in agreement with the repeated observations of enhanced amygdala activity in SAD, we could expect amygdala volumes to be enlarged in socially anxious individuals, and not reduced as the only two structural studies available have reported.

Another point to be considered in investigating volume changes in limbic structures is the role of stress. Because of the nature of their fears, socially anxious individuals are subject to increased stress in daily life compared to non-anxious people, and there is evidence that neurons in the amygdala and hippocampus may suffer excitotoxic damage resulting from sustained glucocorticoid activity associated with stress [11,12]. Accordingly, volume reductions in limbic structures would be an expected finding in SAD, at least in patients with a longer disease duration.

With these considerations in mind, we decided to compare the volumes of the amygdala, hippocampus, and whole brain of treatment-naive individuals meeting criteria for social anxiety disorder and subthreshold social anxiety and healthy controls. We hypothesized that amygdalar and hippocampal volumes would be different across the three groups, without predicting, however, the direction of potential differences.

Our data showed that the bilateral amygdala and hippocampus are enlarged in socially anxious subjects compared to controls. Specificities of these findings are described and discussed below.

**Methods**

**Sample**

Subjects were randomly selected among the participants of a previous investigation on the prevalence of SAD involving 2,319 university students assessed with SAD screening instruments [13]. A subgroup of 60 volunteers from the original sample were invited to attend an individual interview with an experienced psychiatrist for diagnostic (or healthy status) confirmation with the Brazilian version [14,15] of the Structured Clinical Interview for the Diagnostic and Statistical Manual of Mental Disorders, 4th edition (DSM-IV)– clinical and non-patient versions (SCID-CV and SCID-NP) [16].

According to the results of the assessment, 40 participants were invited to take part in the study and assigned to three groups: social anxiety disorder (SAD, n = 13), subthreshold social anxiety (SSA, n = 13), and healthy non-anxious subjects (NSA, n = 14). Participants were classified as having social anxiety disorder when they fulfilled DSM-IV criteria, and subthreshold social anxiety when unreasonable fear of a social situation was present but without associated avoidance or impairment, as proposed by Crum and Pratt (2001) [17]. Data from three subjects (two from the SAD and one from the SSA group) were not included in the analysis because their MRI scans were deemed inadequate by the radiology staff; therefore, the final SAD group had 11 participants and the SSA group had 12. The three groups were matched according to age, sex, education, and socioeconomic status and all subjects were right-handed.

We did not include participants with organic brain syndromes or relevant general medical conditions identified during the interview and clinical examination, epilepsy, psychiatric disorders other than SAD (SAD and SSA groups), history of drug abuse (except nicotine) or who were pregnant at the time of assessment.

Participants in the three groups were not using psychotropic medications and had never received pharmacological or psychotherapeutic treatment for any psychiatric disorder.

**Ethics statement**

The study protocol was approved by the Ethics Committee of the Ribeirão Preto Medical School University Hospital (Process HCRP #11194) and all volunteers gave their signed informed consent to participate.

**MRI data acquisition**

MRI scans were conducted by the radiology staff of the Ribeirão Preto Medical School University Hospital using a 1.5 T Siemens Magneton Vision (Erlangen, Germany) unit with a 25 cm head gradient coil and a circularly polarized coil. To minimize the effects of head movements, subjects were positioned at the scanner by the same staff member using the orbitomeatal line as a reference.

A T1-gradient echo volumetric sequence was acquired in the sagittal plane for the multplanar reformatting and for ROI and voxel-based morphometry (VBM) analyses (TR = 9.7 ms, TE = 4 ms, flip angle = 12°, FOV = 256 mm with continuous 1-millimeter slices in a total of 160 slices per block, matrix = 256x256, NEX = 1). All images were examined by a neuroradiologist and considered adequate.

**Image processing**

The images were initially processed with the software ANA-LYZE AVW 7.0 [18]. The hippocampus and the amygdala were manually traced according to detailed directions proposed by Schumann et al. (2004) [19]. The images of the T1-weighted sequence were converted into 0.5 mm³ voxels and reoriented according to the hippocampal axis (horizontal axis parallel to a line crossing the rostral and caudal poles of the hippocampus). Manual tracing was done over oblique coronal slices and complementary checked on the sagittal and axial planes. Random repeated measures were made from 10 subjects and yielded an intraclass inter-rater reliability coefficient >0.96 for the bilateral hippocampus and >0.95 for the bilateral amygdala.

To compensate for possible differences in brain volume across subjects, the volumes of the amygdala and hippocampus were corrected by dividing the volumes of these structures by the volume of the whole brain of each participant.

Total brain volume was measured through VBM analysis using the VBM Toolbox of the software Statistical Parametric Mapping 5 (SPM5 - dbm.neuro.uni-jena.de/vbm). The sum of all voxels within the segmented images was similar to the total volume of the corresponding partition. The total volume of the brain was then calculated through the sum of gray and white matter volumes.

**Statistical analysis**

Clinical and demographic data were analyzed using Student’s t test for continuous variables and chi-square tests for nominal variables. Volumetric data were compared using analysis of variance (ANOVA) followed by Duncan's post hoc tests when there were differences across the three groups.
Results

We found no statistically significant differences among the SAD, SSA, and NSA groups in terms of their socio-demographic characteristics, as shown in Table 1.

In respect to the volumetric analysis, the volumes of the whole brain \((p = 0.76)\) and of gray \((p = 0.84)\) and white matter \((p = 0.79)\) were equivalent across groups; there were, however, specific differences in the amygdala and hippocampus. Post hoc tests revealed volume increases in the bilateral amygdala \((rAMG: F_{2,34} = 8.66; p = 0.001; lAMG: F_{2,34} = 11.33; p < 0.001)\) and left hippocampus \((F_{2,34} = 10.20; p < 0.001)\) of SAD and SSA participants compared to healthy controls. The volume of the right hippocampus was significantly increased in the SSA group compared to controls \((F_{2,34} = 4.50; p < 0.02)\). The mean volumes of all brain structures examined in the three groups are presented in Table 2. There were no differences between the SAD group and the other two groups regarding right hippocampal volume. Figure 1 provides a graphic representation of these differences.

Discussion

This study investigated the existence of structural abnormalities in the amygdala and hippocampus of treatment-naive socially anxious individuals in relation to healthy controls using MRI. As hypothesized, we found volumetric alterations in subjects with clinical and sub-clinical social anxiety compared to volunteers with no social anxiety. Specifically, socially anxious participants had increased bilateral amygdala and left hippocampus volumes. The right hippocampus was also enlarged in the group with sub-clinical social anxiety compared to controls.

Only two previous studies examined the same brain structures in social anxiety. While Syal et al. (2012) [8] found no volumetric differences in the amygdala and hippocampus of subjects with SAD and controls, Irle et al. (2011) [6] reported precisely the opposite to our findings; that is, volume reductions in the amygdala and hippocampus of subjects with generalized SAD relative to controls, which led us to take a closer look at the processes of atrophy/hypertrophy of limbic structures possibly associated with anxiety.

Research on stress-induced brain plasticity has shown that chronic immobilization stress increases anxiety-like behavior in rats and that this is accompanied by dendritic hypertrophy in the basolateral amygdala and dendritic atrophy in hippocampal area CA3 [20]. In the molecular level, dendritic architecture is mediated by BDNF, the expression of which has been shown to decrease in area CA3 and increase in the basolateral amygdala as a result of chronic immobilization stress, mirroring the structural changes described and following the same temporal profiles of reversal after stress cessation [21].

Considering that social anxiety implies chronic stress, it is possible that the same mechanisms apply to the brain of socially anxious humans, whose amygdala could be enlarged as a result of...
brain-derived neurotrophic factor (BDNF) over-expression leading to dendritic hypertrophy. Even though this interpretation might be correct, our finding of unilateral hippocampal enlargement in subjects with subthreshold social anxiety relative to controls remains unexplained.

A closer analysis of the samples enrolled here and in the study by Irle et al. (2010) [6] might help reduce the discrepancy between findings and shed light into limbic plasticity processes underlying social anxiety. While the mean age of our participants was around 20 years in the three groups, participants in the study by Irle et al. (2010) [6] were on average 10 years older. This is of particular importance because prolonged stress has been linked to brain atrophy - especially in the hippocampus – believed to result from chronic exposure to glucocorticoids [12], whose receptor density is high in the amygdala and hippocampus [22]. Interestingly, the study by Syal et al. (2012) [8] describing cortical thinning in the brain of socially anxious subjects also involved a sample whose mean age was very similar to - and somewhat higher than – the sample studied by Irle et al. (2010) [6].

Although glucocorticoid excitotoxicity is a strong candidate to explain volume reductions in limbic structures in individuals suffering from social anxiety disorder for long periods, it is not the only one. One typical feature of SAD is the high rate of co-occurring psychiatric conditions, the most common of which is depression [23]. It seems beyond doubt that depression is linked to hippocampal atrophy [24,25] and a relatively recent meta-analysis has shown that unmedicated depression is linked with amygdalar atrophy [26].

Taken together, evidence from neuroimaging, cellular, and molecular studies show that our findings and those of Irle et al. (2010) [6] are not necessarily discrepant and can actually be combined in a more comprehensive hypothesis postulating that limbic - and especially amygdalar - plasticity in SAD is biphasic, with volume increases in early stages followed by atrophy resulting from excitotoxic processes in the long run.

The possibility also exists that the volume of limbic structures may be affected by disorder severity instead of or in complex interaction with age, and it could be the case that progressive volume reductions actually result from increased disorder severity. Unfortunately, we did not include measures of disorder severity in our study and were thus unable to investigate this interaction.

Our study has other limitations that should be taken into consideration in interpreting our findings. Although we recruited only subjects who had not been diagnosed with SAD before and thus had never received any psychological or pharmacological treatment for the disorder; we did not specify the SAD subtype (specific or generalized) of our socially anxious participants or how many of them had indicators of depression. In respect to our methods, ROI-based analyses have been reported to increase the risk of false positive and negative results [1] that cannot be ruled out and future studies should consider the inclusion of whole-brain VBM or other automated techniques to check for alterations in non-specific regions.

**Conclusion**

We found structural abnormalities in the amygdala and hippocampus of socially anxious individuals compared to healthy controls. Specifically, subjects with clinical and subthreshold social anxiety had increased bilateral amygdala and left hippocampus volume.

Combined with results from the only other investigation available on the morphology of the amygdala and hippocampus in social anxiety, our findings suggest that limbic plasticity underlying social anxiety has a biphasic pattern characterized by increased amygdalar and hippocampal volume in the early stages of the disorder followed by sustained volume reductions over time.

Further longitudinal structural neuroimaging research is warranted to test this hypothesis and establish the direction of morphological alterations in SAD and their time course and possible interactions with disorder severity. In the molecular level, investigators should look at processes responsible for the atrophy/hypertrophy of limbic structures, with an emphasis on the role of BDNF and the relationship between altered activation and volume.

**Author Contributions**

Conceived and designed the experiments: APJ JASC JECH JPMS RAB. Performed the experiments: FLO JPMS MHNC. Analyzed the data: ALDD APJ MHNC NTA. Contributed reagents/materials/analysis tools: APJ JASC RAB. Wrote the paper: ALDD FLO JECHE JPMS NTA.
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