Acquired alexithymia following damage to the anterior insula

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Abstract

Alexithymia is a subclinical condition characterized by impaired awareness of one’s emotional states, which has profound effects on mental health and social interaction. Despite the clinical significance of this condition, the neurocognitive impairment(s) that lead to alexithymia remain unclear. Recent theoretical models suggest that impaired anterior insula (AI) functioning might be involved in alexithymia, but conclusive evidence for this hypothesis is lacking. We measured alexithymia levels in a large sample of brain-injured patients \( (N=129) \) and non-brain-injured control participants \( (N=33) \), to determine whether alexithymia can be acquired after pronounced damage to the AI. Alexithymia levels were first analyzed as a function of group, with patients separated into four groups based on AI damage: patients with >15% damage to AI, patients with <15% damage to AI, patients with no damage to AI, and healthy controls. An ANOVA revealed that alexithymia levels varied across groups \( (p=0.009) \), with >15% AI damage causing higher alexithymia relative to all other groups \( (all \, p<0.01) \). Next, a multiple linear regression model was fit with the degree of damage to AI, the degree of damage to a related region (the anterior cingulate cortex, ACC), and the degree of damage to the whole brain as predictor variables, and alexithymia as the dependent variable. Critically, increased AI damage predicted increased alexithymia after controlling for the other two regressors (ACC damage; total lesion volume). Collectively, our results suggest that pronounced AI damage causes increased levels of alexithymia, providing critical evidence that this region supports emotional awareness.
Keywords: Alexithymia, Anterior Insula, Anterior Cingulate Cortex, Human Lesion Study, Interoception.
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Alexithymia is a subclinical condition characterized by impaired awareness of one’s emotional states (Aleman, 2005; Bagby et al., 1994; Nemiah et al., 1976; Taylor et al., 1991). The emotional symptoms of a variety of disorders have been linked to co-occurring alexithymia, such as autism (Bird et al., 2010), eating disorders (Brewer et al., 2015a), and schizophrenia (Fogley et al., 2014), suggesting that intact awareness of one’s own emotions might be critical to mental health. The potential translational value of this work has led to an increasing interest in alexithymia both in the public (Robson, 2015; Serani, 2014) and scientific (Bird and Cook, 2013; Lane et al., 2015) literatures, yet the neurocognitive impairments that lead to alexithymia remain unclear.

Recent theoretical work suggests that alexithymia might be the result of impaired “interoception,” referring to one’s sensitivity to the internal state of one’s own body (Gu et al., 2013; Quattrocki and Friston, 2014; Seth, 2013). According to these theories, interoception involves the continual generation of predictive models of the internal state of the body, which are tested against incoming sensory evidence. Discrepancies between predictions and evidence are referred to as “prediction errors,” which are hierarchically organized, with low-level errors representing basic somatosensory signals, which are used to update increasingly higher-level models. At the highest level of the interoceptive prediction hierarchy, the anterior insula (AI) and anterior cingulate cortex (ACC) are believed to be
involved in representing our conscious emotional experience. In particular, AI has a clearly ascribed functional role in this framework: it generates awareness of one’s emotions by integrating prediction errors across lower levels of the hierarchy (Gu et al., 2013; Seth, 2013).

Several lines of evidence suggest that AI abnormalities might be a neurophysiological factor contributing to alexithymia. Specifically, alexithymia is associated with reduced AI activation when rating the emotional valence of stimuli from the International Affective Pictures System (Silani et al., 2008), and when observing emotional facial expressions (Kano et al., 2003; Reker et al., 2010) and others in pain (Bird et al., 2010; FeldmanHall et al., 2013). In addition to these functional differences, recent structural neuroimaging studies (i.e. voxel-based morphometry) found reduced grey matter volumes of the AI in alexithymic individuals (Borsci et al., 2009; Ihme et al., 2013), and reduced coherence of anterior-insula based anatomical networks (Bernhardt et al., 2014). In accordance with the presence of AI abnormalities in alexithymia, and its role in interoception, a recent behavioral study found a negative association between interoceptive awareness and the emotional symptoms of alexithymia (Herbert et al., 2011). Collectively, such findings have led to the hypothesis that abnormal recruitment of the AI signals impaired integration of interoceptive prediction errors, which might lead to deficient emotional awareness in alexithymia (Brewer et al., 2015b; Quattrocki and Friston, 2014).
Research on the neural basis of alexithymia has traditionally focused on individuals whose alexithymic personality features arose over the course of development, which cannot determine whether any of these neurocognitive features reflect an underlying cause of the condition. In contrast, human lesion studies hold the potential to establish a causal link between particular brain regions and alexithymia. There is some evidence for acquired alexithymia following brain injury (Henry et al., 2006; Williams and Wood, 2013, 2010). However, no studies have analyzed alexithymia as a function of lesion location, making it impossible to link alexithymia to AI damage specifically. Accordingly, we measured alexithymia in a large sample of patients with well-localized penetrating traumatic brain injuries (N=129 total) and closely matched control participants (N=33) as part of the Vietnam Head Injury Study (VHIS; Raymont et al., 2011). We predicted that patients with pronounced AI damage would demonstrate elevated levels of alexithymia, relative to other brain-injured patients and healthy control participants. Furthermore, we hypothesized a continuous relationship, whereby the extent of damage to AI would predict alexithymia severity.
Methods

Participants

Data were from Phase 4 of the VHIS (2008-2012), a prospective study of male Vietnam war veterans with penetrating traumatic brain injuries (TBI), and non-brain-injured combat veterans (Raymont et al., 2011). In total, target measures were collected from 129 patients with TBI and 33 control participants. The TBI and control groups were matched with respect to age ($M_{\text{ctrl}}=63.33$ years, $SD_{\text{ctrl}}=3.80$; $M_{\text{TBI}}=63.29$, $SD_{\text{TBI}}=2.89$; $F_{1.160}=0.01$, $p=0.94$, $\eta^2=0.00$; $BF_{10}=0.21$), education ($M_{\text{ctrl}}=15.06$ years, $SD_{\text{ctrl}}=2.12$; $M_{\text{TBI}}=14.55$, $SD_{\text{TBI}}=2.27$; $F_{1.160}=1.37$, $p=0.24$, $\eta^2=0.01$; $BF_{10}=0.38$), pre-injury intelligence ($M_{\text{ctrl}}=72.91$ percentile, $SD_{\text{ctrl}}=17.06$; $M_{\text{TBI}}=64.37$, $SD_{\text{TBI}}=23.29$; $F_{1.136}=2.68$, $p=0.10$, $\eta^2=0.02$; $BF_{10}=0.76$), and handedness (CTRL group: 26 right-handed, 5 left-handed, 2 ambidextrous; TBI group: 105 right-handed, 22 left-handed, 2 ambidextrous; $X^2=2.24$, $p=0.33$; $BF_{10}=0.06$). An institutional review board at the National Institute of Neurological Disorders and Stroke at the National Institute of Health, Bethesda, MD, USA approved the VHIS Phase 4 protocols, and all participants provided their written informed consent.

Materials

Lesion localization was performed using the ABLe software package (Solomon et al., 2007) in MEDx v.3.44 (Medical Numerics, Germantown, MD, USA), and an AI atlas was created in MRlcron (Rorden, 2007). For statistical
analyses, we performed classical hypothesis tests and Bayesian hypothesis tests, as recent advances in quantitative psychology have criticized the practice of wholly relying on $p$-values for statistical evidence (Kass and Raftery, 1995; Kruschke, 2015; Wagenmakers, 2007). All inferential analyses were carried out using the JASP software package (Love et al., 2015). Figure 2 was constructed using the ggplot2 (Wickham, 2009) function implemented in R v.3.2.1 (R Core Team, 2013).

**Lesion Localization and Groupings**

Non-contrast axial computerized tomography (CT) scans were acquired during Phase 3 (2003-2006) of the VHIS using a GE Medical Systems Light Speed Plus CT scanner. Six TBI patients in the present sample did not participate in Phase 3, and their CT scans were obtained during Phase 4. On average, patients were scanned 38.18 years after their injury ($SD=7.96$ years). Magnetic resonance imaging (MRI) could not be performed with patients due to the possible presence of metal fragments from shrapnel or bullet wounds, or residual metallic surgical clips or cranioplasties from surgery. Image volumes were composed of 0.4 mm x 0.4 mm voxels and 2.5 mm thick overlapping slices with a 1 mm slice interval. Lesions were manually drawn on CT slices by a trained neuropsychiatrist, and reviewed by a researcher (JG) who was blind to the results of neuropsychological testing. CT images were normalized to a template brain in MNI space. Percent volume loss was calculated by summing the traced lesion areas in each CT slice, multiplying by slice thickness and
dividing by total brain volume. In this study, we identified percent volume loss to two regions of interest: Al damage (target region) was defined based on a 3D insula template (Afif et al., 2013), and ACC damage (comparison region) was defined using the automated anatomical labeling (AAL) atlas (Tzourio-Mazoyer et al., 2002).

Using the percent volume loss data, we wanted to separate participants into groups based on the extent of their damage to Al. Since small lesion volumes are more amenable to recovery of function (Irle, 1987), it was important to set a relatively high volume threshold, above which it was more likely that any acquired neurocognitive impairments would still be present at testing. Past work has demonstrated that >15% damage to a target brain region can be sufficient to induce lasting effects on emotional processes (Koenigs et al., 2008; Tranel et al., 2005). Accordingly, participants were separated into four groups (Figure 1): healthy controls (N=33), patients with 0% damage to Al (N=89), patients with >0% and <15% damage to Al (N=31), and patients with >15% damage to Al (N=9).

Behavioral Measure
Alexithymia was measured using the twenty item Toronto Alexithymia Scale (TAS-20), the gold standard self-report measure used to index alexithymia (Bagby et al., 1994). The TAS-20 was collected during Phase 4 of the VHIS (2008-2012), an average of 44.34 years (SD=7.46 years) after the injury for the
patient group. The TAS-20 contains 20 statements with which participants rate their agreement on a 5-point likert scale (1=strongly disagree, 2=disagree, 3=neutral, 4=agree, 5=strongly agree). The TAS-20 includes a combination of positively- and negatively-scored items (e.g. positive: “I am often confused about what emotion I am feeling”; e.g. negative: “Being in touch with emotions is essential”), and researchers typically calculate a TAS-20 total score (TAS-total), with higher scores indicating higher levels of alexithymia. Furthermore, the TAS-20 can broken down into three subscales: i) difficulty identifying feelings (TAS-DIF; i.e. related to poor interoceptive and emotional awareness; e.g. “When I am upset, I don’t know if I am sad, frightened, or angry,” “I am often puzzled by sensations in my body,” etc.), ii) difficulty describing feelings (TAS-DDF; i.e. related to poor communication of feelings to others; e.g. “it is difficult for me to find the right words for my feelings,” “People often tell me to describe my feelings more,” etc.), and iii) externally-oriented thinking (TAS-EXO; i.e. related to a chronic focus on external events over internal states; e.g. “I prefer to analyze problems rather than just describe them,” “I prefer talking to people about their daily activities rather than their feelings,” etc.; Bagby et al., 1994). Given the hypothesized role of the anterior insula in contributing to emotional awareness, we hypothesized that damage to this region was most likely to influence scores on the TAS-DIF subscale.

Alexithymia covaries with other emotional disturbances, such as anxiety, depression and, of particular relevance to the present sample, post-traumatic
stress disorder (PTSD; Badura, 2003; Grynberg et al., 2010). Therefore, each of the participants completed measures of trait anxiety (Spielberger et al., 1983), depression (Beck, 1996), and PTSD (Keane et al., 1988), and linear regression models were fitted with TAS-total, TAS-DIF, TAS-DDF, and TAS-EXO as dependent variables, and the three control measures as predictors. The standardized residuals from these four models indexed the various dimensions of alexithymia after controlling for individual differences in anxiety, depression, and PTSD. These standardized alexithymia measures were used in all inferential analyses.

**Results**

**Analyses of Variance**

We began by performing an ANOVA to determine whether pronounced damage to AI causes elevated levels of alexithymia, as indexed using the TAS-total score. The overall ANOVA was significant ($F_{3,158}=3.98, p=0.009, \eta^2=0.07; BF_{10}=2.20$), providing evidence that there were group differences in alexithymia. Planned independent samples $t$-tests and Bayesian $t$-tests suggested that the group with pronounced (i.e. >15%) AI damage was associated with a significant increase in alexithymia relative to healthy control participants ($t_{40}=2.84, p=0.007, d=1.07, BF_{10}=6.38$; **Figure 2a**), patients with no AI damage ($t_{96}=3.43, p<0.001, d=1.20, BF_{10}=31.61$; **Figure 2a**), and patients with some AI damage ($t_{38}=2.60, p=0.01, d=0.98, BF_{10}=4.02$; **Figure 2a**).
Next, we performed ANOVAs on the three TAS-20 subscales to determine whether AI damage was specifically associated with heightened scores on the “difficulty identifying feelings” subscale (TAS-DIF). The TAS-DIF ANOVA provided evidence for significant differences between groups ($F_{3,158}=4.91$, $p=0.003$, $\eta^2=0.08$; $BF_{10}=5.13$; Figure 2b). Specifically, TAS-DIF levels were significantly higher in patients with large AI lesions relative to healthy controls ($t_{40}=4.40$, $p<0.001$, $d=1.65$, $BF_{10}=240.36$; Figure 2b), patients with no damage to AI ($t_{96}=3.29$, $p=0.001$, $d=1.15$, $BF_{10}=21.66$; Figure 2b), and patients with minimal damage to AI ($t_{38}=2.84$, $p=0.007$, $d=1.08$, $BF_{10}=6.38$; Figure 2b). In contrast, results for the “difficulty describing feelings” (TAS-DDF) and “externally-oriented thinking” (TAS-EXO) subscales provided no evidence for a relationship with AI damage. Neither the TAS-DDF ANOVA ($F_{3,158}=2.35$, $p=0.07$, $\eta^2=0.04$; $BF_{10}=0.48$; Figure 2c), nor the TAS-EXO ANOVA ($F_{3,158}=1.58$, $p=0.20$, $\eta^2=0.03$; $BF_{10}=0.27$; Figure 2d), provided credible evidence for a difference between groups.

Therefore, in accordance with our predictions, pronounced AI damage caused increased levels of alexithymia, and this effect was primarily driven by differences in the TAS-DIF subscale.

**Regression Analyses**

Next, we conducted classical and Bayesian linear regression analyses to provide further evidence that damage to AI was specifically associated with alexithymia. Both AI and ACC damage were included in the model. AI and ACC
are among the most commonly co-activated structures in the functional MRI (fMRI) literature on emotional processing (Craig, 2009; Medford and Critchley, 2010), and are often both damaged in lesion patients (Damasio et al., 2013; Khalsa et al., 2009), making it difficult to discern the differential contribution of each region. The initial regression model included the standardized TAS-total score as the dependent variable, and included percent damage to AI, percent damage to ACC, and total percent damage as regressors (Table 2), which were all $T$-scored in an attempt to reduce multicollinearity. Overall, the model explained a significant portion of variance in alexithymia ($R^2=0.10, F_{3,125}=4.39, p=0.006, f^2=0.10, BF_{10}=4.32$; Table 2), with AI damage predicting a significant increase in alexithymia ($\beta=0.02; t_{125}=2.38, p=0.01$, one-tailed, $BF_{10}=8.82$; Table 2). In contrast, neither ACC damage ($p=0.07$, one-tailed, $BF_{10}=0.71$; Table 2) nor total lesion volume ($p=0.33$, one-tailed, $BF_{10}=0.31$; Table 2) contributed significantly to the model. Importantly, none of the regressors demonstrated an issue with multicollinearity (all VIF<2).

Akin to our ANOVA procedures, we also conducted separate linear regression analyses for each of the TAS-20 subscales. A significant amount of variance on the TAS-DIF subscale was accounted for by our regression model ($R^2=0.09, F_{3,125}=4.25, p=0.007, f^2=0.10, BF_{10}=3.67$; Table 2). Increases in AI damage predicted increases in TAS-DIF ($\beta=0.02; t_{125}=1.86, p=0.03$, one-tailed, $BF_{10}=4.04$; Table 2), whereas the ACC damage ($p=0.05$, one-tailed, $BF_{10}=1.00$; Table 2) and total volume loss variables ($p=0.44$, one-tailed, $BF_{10}=0.39$; Table 2)
did not contribute significantly to the model. In contrast, the models did not account for a significant proportion of variance in the TAS-DDF ($R^2=0.04$, $F_{3,125}=1.86, p=0.14, f^2=0.04, BF_{10}=0.19$; Table 2) and TAS-EXO ($R^2=0.04$, $F_{3,125}=1.76, p=0.16, f^2=0.04, BF_{10}=0.17$; Table 2) subscales.

**Control Analyses**

To ensure that our key ANOVA results were specific to alexithymia and not some unanticipated factor, we compared the groups on a variety of demographic and control measures (Table 1). The only measures that showed slight differences between groups were pre- and post-injury intelligence, and we entered these variables into an analysis of covariance similar to our original ANOVA. Again, there was a significant difference between the groups ($F_{3,156}=3.08, p=0.03, \eta^2=0.06$), suggesting that group differences in intelligence could not account for our findings.

Lastly, since the TAS-20 asks participants to perform a metacognitive appraisal (e.g. “I often don’t know why I am angry”), this might be problematic if they are not aware of their own interoceptive or emotional difficulties (Lane et al., 2015). Therefore, we wanted to ensure that TAS-20 performance was negatively associated with a measure of emotional awareness that did not rely on metacognition. Accordingly, a subsample of our participants ($N=135$) completed the Mayer-Salovey-Caruso Emotional Intelligence Scale (MSCEIT; Mayer et al., 2002), a self-report measure that asks participants to interpret emotional stimuli.
rather than performing an appraisal of their own emotions. Crucially, the “perceiving emotions” and “understanding emotions” subscales of the MSCEIT were negatively associated with raw TAS-total scores (perceiving emotions: $r=-0.18$, $p=0.02$, $BF_{10}=1.05$; understanding emotions: $r=-0.24$, $p=0.003$, $BF_{10}=4.66$), supporting the construct validity of our alexithymia scale.

**Discussion**

In the present study, we examined whether damage to AI can cause acquired alexithymia. Recent computational models suggest that interoceptive signals are integrated by AI to produce a cohesive emotional awareness state (Gu et al., 2013; Seth, 2013) and, by extension, that impaired AI processing might be involved in producing alexithymia (Brewer et al., 2015b; Quattrocki and Friston, 2014). The present data represent compelling evidence in support of this hypothesis, demonstrating that pronounced damage to AI causes impaired awareness of emotional states in the self. The present study also helps to distinguish between AI and ACC involvement in emotional cognition. AI and ACC are commonly co-activated in the fMRI literature (Craig, 2009; Medford and Critchley, 2010), and are both compromised in most AI lesion studies (Damasio et al., 2013; Khalsa et al., 2009), making it difficult to demonstrate clear dissociations between their functions. Here, using a multiple linear regression approach, we present preliminary evidence for a rare dissociation between the
two regions: AI damage is associated with elevated levels of alexithymia whereas ACC damage is not.

Alexithymia has traditionally been studied with respect to its comorbidity with neurodevelopmental disorders such as autism spectrum disorders (Bird and Cook, 2013) and schizophrenia (Fogley et al., 2014), but recent studies have suggested that alexithymia can be acquired following brain injury (Henry et al., 2006; Williams and Wood, 2013, 2010). The present findings extend this work by showing that alexithymia can be acquired following injuries to the AI specifically, with more pronounced lesions being associated with greater impairment. This finding provides causal support for the view that abnormal AI recruitment may be a key neurophysiological feature of developmental alexithymia (Brewer et al., 2015b; Quattrocki and Friston, 2014).

The present evidence for acquired alexithymia following AI damage is in agreement with a considerable neuroimaging literature (Craig, 2009), and several human lesion studies implicating AI in interoception and emotional awareness (Gu et al., 2013; Jones et al., 2010). For example, AI grey matter volumes (Borsci et al., 2009; Ihme et al., 2013) and functional activations are reduced in alexithymic individuals (Bird et al., 2010; Kano et al., 2003; Reker et al., 2010; Silani et al., 2008). Within the broader alexithymic construct, the present findings suggest that AI is specifically implicated in emotional awareness (i.e. TAS-DIF subscale). This conclusion is in line with other human lesion studies, which have
found AI damage to be associated with reduced sensitivity to aversive somatosensory stimuli (Greenspan et al., 1999), and a muted experience of disgust (Calder et al., 2000). These converging lines of evidence support the arguments that AI produces an emotional awareness state by integrating ascending interoceptive signals (Gu et al., 2013; Seth, 2013), and that this system is disrupted in alexithymia (cf. Brewer et al., 2015b; Herbert et al., 2011; Quattrocki and Friston, 2014).

In contrast to the present findings, recent case studies have documented relatively intact performance on tasks related to conscious awareness, emotion, and interoception in patients with AI lesions (Damasio et al., 2013; Feinstein et al., in press; Khalsa et al., 2009; Philippi et al., 2012). However, these studies are based on two patients (Patient B and Patient R), raising the possibility that these individuals have developed compensatory strategies for coping with impaired emotional awareness that were not deployed by the present sample of AI patients. Additionally, none of the previous AI lesion studies measured alexithymia using the TAS-20, and it is possible that the TAS-20 measures dimensions of emotional awareness that require a functioning AI, whereas the other behavioral tasks used in previous studies did not. By showing significant increases in alexithymia in a group of patients with AI damage ($N=9$), and a specific association between the extent of AI damage and increased alexithymia, our data provide evidence that AI is indeed necessary for the dimensions of emotional awareness that are disrupted in alexithymia.
The present findings also appear to contradict a Positron Emission Tomography study by Lane and colleagues (1998), which found that ACC activity during emotional experience was positively associated with scores on the “Levels of Emotional Awareness Scale” (LEAS; Lane et al., 1990) in healthy volunteers (Lane et al., 1998). In contrast, the present findings suggest that emotional awareness is not disrupted in brain-injured patients as a function of ACC damage, beyond the variance accounted for by AI damage. The reason(s) for this discrepancy are unclear, but it seems likely that the LEAS and TAS-20 reflect different aspects of emotional awareness that differentially recruit the ACC and AI, respectively. Future functional neuroimaging studies will be required to further dissociate the emotional awareness operations of AI and ACC.

One potential limitation of the present study was that we relied on an alexithymia measure (the TAS-20) that required participants to have insight into their own emotional awareness difficulties. Fortunately, the negative association between our alexithymia data and the more objective MSCEIT supports the construct validity of the TAS-20. In future AI lesion-symptom mapping studies, researchers should aim to provide conceptual replications of the present findings using more objective measures of emotional awareness. An additional caveat to our findings is that our sample was limited to older male combat veterans studied approximately 45 years post-injury.
We believe that our findings, notwithstanding the caveats articulated above, provide crucial evidence for the importance of AI in emotional awareness. Future studies should work to clarify how emotional states are linked with the ongoing cognitive and social context within conscious awareness – an important step in determining the personal relevance of daily experiences.
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# Table 1. Demographics from the three groups in the initial ANOVA analyses.

***: Significant at the $p<0.001$ level; $^t$: Trending at $p<0.1$. Age and education are listed in the number of years. Pre-injury intelligence is a percentile score on the Armed-Forces Qualification Test (AFQT-7A), which was collected at induction to the military and is highly correlated with the Wechsler Adult Intelligence Scale-III (WAIS-III; Grafman et al., 1988). Post-injury intelligence was measured using percentile scores on the WAIS-III (Wechsler, 1997) at the time of evaluation. Missing values for both the pre-injury intelligence and post-injury intelligence scores were filled using a mean replacement method. Lastly, given their potential association with alexithymia, we also analyzed raw scores from the Beck Depression Inventory-II (Beck, 1996), the State-Trait Anxiety Inventory (Spielberger et al., 1983), and the Mississippi PTSD scale (Keane et al., 1988).

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<thead>
<tr>
<th></th>
<th>Controls</th>
<th>0% AI</th>
<th>&lt;15% AI</th>
<th>&gt;15% AI</th>
<th>Inferential Tests</th>
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<td><strong>Demographics:</strong></td>
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<tr>
<td><strong>Age</strong></td>
<td>63.33 (3.80)</td>
<td>63.38 (3.15)</td>
<td>62.97 (2.17)</td>
<td>63.44 (2.45)</td>
<td>$F_{3,158}=0.15, p=0.93, \eta^2=0.003; BF_{10}=0.06$</td>
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<tr>
<td><strong>Education</strong></td>
<td>15.06 (2.12)</td>
<td>14.60 (2.29)</td>
<td>14.68 (2.01)</td>
<td>13.67 (2.92)</td>
<td>$F_{3,158}=0.96, p=0.41, \eta^2=0.02; BF_{10}=0.12$</td>
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<tr>
<td><strong>Handedness (R,L,A)</strong></td>
<td>(26,5,2)</td>
<td>(69,18,2)</td>
<td>(28,3,0)</td>
<td>(8,1,0)</td>
<td>$\chi^2=5.08, p=0.53; BF_{10}=2.65e-4$</td>
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<tr>
<td><strong>Control Measures:</strong></td>
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<tr>
<td><strong>Pre-injury Intelligence</strong> $^t$</td>
<td>70.52 (14.24)</td>
<td>63.28 (23.20)</td>
<td>70.61 (16.66)</td>
<td>55.67 (24.37)</td>
<td>$F_{3,158}=2.32, p=0.08, \eta^2=0.04; BF_{10}=0.56$</td>
</tr>
<tr>
<td><strong>Post-injury Intelligence</strong>*</td>
<td>75.28 (21.45)</td>
<td>66.68 (26.37)</td>
<td>61.39 (25.11)</td>
<td>28.73 (31.31)</td>
<td>$F_{3,158}=8.19, p&lt;0.001, \eta^2=0.14; BF_{10}=185.55$</td>
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<tr>
<td><strong>Depression</strong></td>
<td>10.64 (8.74)</td>
<td>7.96 (7.67)</td>
<td>7.55 (9.44)</td>
<td>6.11 (6.68)</td>
<td>$F_{3,158}=1.24, p=0.30, \eta^2=0.02; BF_{10}=0.18$</td>
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<tr>
<td><strong>Anxiety</strong></td>
<td>37.55 (12.45)</td>
<td>33.65 (10.14)</td>
<td>33.26 (11.54)</td>
<td>31.11 (7.37)</td>
<td>$F_{3,158}=1.46, p=0.23, \eta^2=0.03; BF_{10}=0.23$</td>
</tr>
<tr>
<td><strong>PTSD</strong></td>
<td>83.67 (22.88)</td>
<td>79.79 (22.70)</td>
<td>76.26 (18.68)</td>
<td>74.33 (19.58)</td>
<td>$F_{3,158}=0.80, p=0.50, \eta^2=0.02; BF_{10}=0.11$</td>
</tr>
</tbody>
</table>
Table 2. Results from the regression analysis. *: Regressor is significant at the $p<0.05$ (one-tailed) and $BF_{10}>3$ level. **: Regressor is significant at the $p<0.01$ level.

<table>
<thead>
<tr>
<th>Regressor</th>
<th>$\beta$</th>
<th>$t$-Test</th>
<th>$BF_{10}$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TAS-DIF (“Difficulty Identifying Feelings”)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T$-AI*</td>
<td>0.02</td>
<td>$t=1.86$, $p=0.03$</td>
<td>4.04</td>
</tr>
<tr>
<td>$T$-ACC</td>
<td>0.01</td>
<td>$t=1.64$, $p=0.05$</td>
<td>1.00</td>
</tr>
<tr>
<td>$T$-Volume Loss</td>
<td>0.005</td>
<td>$T=0.14$, $p=0.44$</td>
<td>0.39</td>
</tr>
<tr>
<td>Overall Model**: $R^2=0.09$; $F_{3,125}=4.25$, $p=0.007$, $f^2=0.10$; $BF_{10}=3.67$</td>
<td></td>
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</tr>
<tr>
<td><strong>TAS-DDF (“Difficulty Describing Feelings”)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T$-AI</td>
<td>0.02</td>
<td>$t=1.78$, $p=0.04$</td>
<td>1.79</td>
</tr>
<tr>
<td>$T$-ACC</td>
<td>0.004</td>
<td>$t=0.50$, $p=0.31$</td>
<td>0.31</td>
</tr>
<tr>
<td>$T$-Volume Loss</td>
<td>-0.01</td>
<td>$t=-0.31$, $p=0.38$</td>
<td>0.33</td>
</tr>
<tr>
<td>Overall Model: $R^2=0.04$; $F_{3,125}=1.86$, $p=0.14$, $f^2=0.04$; $BF_{10}=0.19$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>TAS-EXO (“Externally-Oriented Thinking”)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T$-AI</td>
<td>0.02</td>
<td>$t=1.70$, $p=0.04$</td>
<td>0.94</td>
</tr>
<tr>
<td>$T$-ACC</td>
<td>0.009</td>
<td>$t=1.11$, $p=0.13$</td>
<td>0.50</td>
</tr>
<tr>
<td>$T$-Volume Loss</td>
<td>-0.03</td>
<td>$t=-0.84$, $p=0.20$</td>
<td>0.30</td>
</tr>
<tr>
<td>Overall Model: $R^2=0.04$; $F_{3,125}=1.76$, $p=0.16$, $f^2=0.04$; $BF_{10}=0.17$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>TAS-total</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$T$-AI**</td>
<td>0.02</td>
<td>$t=2.38$, $p=0.009$</td>
<td>8.82</td>
</tr>
<tr>
<td>$T$-ACC</td>
<td>0.01</td>
<td>$t=1.50$, $p=0.07$</td>
<td>0.71</td>
</tr>
<tr>
<td>$T$-Volume Loss</td>
<td>-0.01</td>
<td>$t=-0.44$, $p=0.33$</td>
<td>0.31</td>
</tr>
<tr>
<td>Overall Model**: $R^2=0.10$; $F_{3,125}=4.39$, $p=0.006$, $f^2=0.10$; $BF_{10}=4.32$</td>
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</tr>
</tbody>
</table>
Figures

Figure 1. Representative axial slices depicting (A) the template used to define the insula (pink circles approximately marking the anterior sector), and (B) overlay density maps depicting the preserved anterior insula (AI) in the no damage group ($N=89$), some slight damage to AI in the $<15\%$ group ($N=31$), and pronounced damage in the $>15\%$ group ($N=9$). Images are in radiological space (i.e. right is left).
Figure 2. (A) Patients with pronounced damage to AI demonstrated significant increases in alexithymia (TAS-total) relative to all other patient groups as well as healthy controls. (B) Scores on the “difficulty identifying feelings” (DIF) subscale mirrored the TAS-total results, whereas (C-D) neither the “difficulty describing feelings” (DDF) nor the “externally-oriented thinking” (EXO) subscales varied significantly as a function of group. ***: p<0.001, **: p<0.01; ns: ANOVA p>0.05.