Cingulo-insular structural alterations associated with psychogenic symptoms, childhood abuse and PTSD in functional neurological disorders

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ABSTRACT

Objective Adverse early-life events are predisposing factors for functional neurological disorder (FND) and post-traumatic stress disorder (PTSD). Cingulo-insular regions are implicated in the biology of both conditions and are sites of stress-mediated neuroplasticity. We hypothesised that functional neurological symptoms and the magnitude of childhood abuse would be associated with overlapping anterior cingulate cortex (ACC) and insular volumetric reductions, and that FND and PTSD symptoms would map onto distinct cingulo-insular areas.

Methods This within-group voxel-based morphometry study probes volumetric associations with self-report measures of functional neurological symptoms, adverse life events and PTSD symptoms in 23 mixed-gender FND patients. Separate secondary analyses were also performed in the subset of 18 women with FND to account for gender-specific effects.

Results Across the entire cohort, there were no statistically significant volumetric associations with self-report measures of functional neurological symptom severity or childhood abuse. In women with FND, however, parallel inverse associations were observed between left anterior insular volume and functional neurological symptoms as measured by the Patient Health Questionnaire-15 and the Screening for Somatoform Symptoms Conversion Disorder subscale. Similar inverse relationships were also appreciated between childhood abuse burden and left anterior insular volume. Across all subjects, PTSD symptom severity was inversely associated with dorsal ACC volume, and the magnitude of lifetime adverse events was inversely associated with left hippocampal volume.

Conclusions This study reveals distinct cingulo-insular alterations for FND and PTSD symptoms and may advance our understanding of FND. Potential biological convergence between stress-related neuroplasticity, functional neurological symptoms and reduced insular volume was identified.

INTRODUCTION

While functional neurological disorder (FND) (conversion disorder) is common and imposes significant healthcare costs,1–3 the neuropathophysiology of this condition remains poorly understood. Functional neurological symptoms comprise 16% of outpatient neurology referrals, second only to headache.4 Patients with FND present with medically unexplained sensory and motor symptoms including fatigue, pain, non-dermatomal sensory deficits, limb weakness, gait difficulties, non-epileptic seizures and abnormal movements. In the USA, an estimated $256 billion is spent annually in healthcare for medically unexplained illness.5 This public health problem is heightened by observations that FND is a neglected condition at the intersection between neurology and psychiatry. While neurologists are frustrated in caring for this population and lack a neurobiological understanding of FND,6 psychiatrists are similarly uncomfortable treating patients whose chief complaint is physical.

Adverse life events are a risk factor for FND, and the magnitude of early-life maltreatment is associated with illness severity.7 Studies suggest that between a quarter and three-quarters of individuals with FND endorse early-life maltreatment.8–11 Childhood abuse has been linked to increased FND severity, and individuals reporting several traumatic childhood events more often exhibit a multiplicity of FND symptoms.7 In those with non-epileptic seizures, past sexual abuse is linked to earlier age of onset, more severe convulsions, intrusive traumatic recollections and disability.12 Furthermore, the development of additional medically unexplained symptoms in FND is predicted by antecedent trauma.11 These collective observations suggest that previously experienced adverse life events, particularly childhood abuse, are important predisposing factors for the development of functional neurological symptoms.

Childhood abuse is also linked to increased risk of developing affective disorders, and those with these conditions and past abuse exhibit greater illness severity, increased psychiatric comorbidities and less favourable prognoses.13 Associations between FND and other trauma-related disorders including post-traumatic stress disorder (PTSD) are well documented.10,14 Importantly, studies also suggest that gender may modulate the development of psychopathology following childhood abuse.15 It is not yet known, however, if adverse life events affect developmentally vulnerable neural circuits in a disorder-specific manner or if aberrant neuroplastic changes following adverse life events facilitate a general increased predilection for psychopathology.
While FND is understudied compared to other neuropsychiatric disorders, early neuroimaging studies suggest an important role for the cingulo-insular (salience) network in the pathophysiology of FND.36–38 The anterior cingulate cortex (ACC) and the insula are two paralimbic convergent sites for the multimodal integration of affective, viscerosomatic/nociceptive and cognitive processing.20–22 Compared with healthy controls, patients with non-epileptic seizures display structural alterations in the dorsal ACC23 and insula.24 Individuals with somatic symptom disorder with predominant pain compared with controls exhibit decreased cingulate and insular grey matter volumes.25 In functional neuroimaging studies, FND patients show heightened insular and ACC activations during motor tasks compared to controls.26–28 Increased cingulate gyrus activations during affectively valenced face processing have also been observed in FND.28 These convergent findings implicate cingulo-insular regions in the biology of FND, yet limited research has studied structural brain-symptom severity and brain-disease risk relationships in this population.

Neuroimaging studies also link cingulo-insular alterations to the pathophysiology of PTSD. MRI studies show grey matter reductions in the ACC and hippocampus in patients with PTSD compared with controls,29 30 as well as inverse associations between ACC volume and PTSD severity.31 To a lesser extent, grey matter reductions in the insula have also been characterised in PTSD.32 While heightened insular activations may occur across anxiety disorders, reduced top-down rostral ACC and increased amygdala activations are well described in PTSD.30

In this voxel-based morphometry (VBM) MRI study, we used a within-group design to dimensionally probe grey matter associations with self-report measures of functional neurological symptoms, adverse life events and PTSD symptoms in 23 patients with FND. Given the increased incidence of FND in women and observations that gender may modulate the development of psychopathology following childhood abuse,15 secondary analyses were also performed separately in the subgroup of 18 women to account for gender-specific effects. As detailed in several conceptual models by our group,16–18 we hypothesised that functional neurological symptom severity and the magnitude of childhood abuse would be associated with overlapping ACC and insular volumetric reductions. We also hypothesised that FND and PTSD symptom severity would map onto distinct cingulo-insular brain areas.

METHODS

Participants and psychometric assessments

Twenty-three subjects with FND (mean age 41.6±11.6 years; 18 women and 5 men; mean illness duration 3.7±4.5 years) were recruited from an integrated behavioural neurology-neuropsychiatry FND clinic at the Massachusetts General Hospital.34 All patients met diagnostic criteria for at least one FND subtype including clinically established functional movement disorders34 (n=12), documented (n=6) or clinically established (n=1) non-epileptic seizures35 and/or exhibited positive examination findings for functional weakness (n=11).36 Seven out of 23 subjects also exhibited non-dermatomal sensory deficits, and seven distinct individuals had mixed motor FND. Diagnoses were based on a clinical evaluation by a dual-trained and board-certified neurologist and psychiatrist (DLP). Exclusion criteria included: any significant major neurological disorder resulting in specific MRI abnormalities (ie, stroke, severe traumatic brain injury), epileptic seizures, other movement disorders, poorly controlled major medical illnesses with known central nervous system consequences, ongoing alcohol dependence or illicit substance misuse, a history of mania or psychosis, and/or active suicidality. Additional current psychiatric diagnoses as measured by the Structured Clinical Interview (SCI) for DSM-IV-TR Axis I Disorders included major depressive disorder (n=8), dysthymia (n=2), panic disorder (n=10), generalised anxiety disorder (n=9), PTSD (n=5), other somatoform disorders (n=10) and alcohol abuse (n=1). Sixteen of the 23 subjects were taking psychoactive medications. Subjects provided signed informed consent, and this study was approved by the Partners Human Research Committee. See online Supplemental table 1 for participants’ demographics and diagnoses.

Subjects participated in two closely timed research visits. In one visit, subjects completed the SCI and self-report measures, including: the Patient Health Questionnaire-15 (PHQ-15),37 Childhood Trauma Questionnaire (CTQ),38 Life Events Checklist-5 (LEC-5), PTSD Checklist for DSM-5 (PCL-5),40 Beck Depression Inventory-II (BDI)41 and the Spielberger Trait Anxiety Inventory (STAI-T).37 On the day of scanning, subjects completed the Conversion Disorder subscale of the Screening for Somatoform Symptoms-7 scale (SOMS:CD).42

The PHQ-15 is a 15-item measure of somatic symptoms within the past 4 weeks; each item is scored on a 3-point scale ranging from ‘not bothered at all’ to ‘bothered a lot’. The SOMS:CD is a 14-item measure of motor and sensory functional neurological symptoms within the past 7 days; each item is scored on a 5-point scale regarding the perceived degree of impairment ranging from ‘not at all’ to ‘very severe’. To emphasise a trans-diagnostic, dimensional approach to brain-symptom relationships in this study, we refer to symptoms captured by both the PHQ-15 and the SOMS:CD as functional (psychogenic) neurological symptoms.

The CTQ is a 25-item measure of childhood/adolescent abuse and neglect; cumulative indices of abuse (sexual, physical and emotional (CTQ-Abuse)) and neglect (emotional and physical (CTQ-Neglect)) were calculated separately. The LEC-5 is a 17-category measure of lifetime adverse events, and the PCL-5 is a 20-item measure of PTSD symptoms that can be subdivided into re-experiencing, avoidance, negative alterations in cognition and mood, and hyperarousal subdomains.

MRI data acquisition

In the MRI session, subjects were placed in a Siemens 3 Tesla Trio scanner to acquire a 3D T1-weighted magnetisation prepared rapid acquisition gradient echo sequence with the following parameters: orientation sagittal; matrix size 256×256; voxel size 1×1×1 mm; slice thickness 1 mm, slices 160; repetition time 2300 ms; echo time 2.98 ms; and field of view 256 mm. Bi-temporal foam pads were used to restrict head motion.

MRI data preprocessing and analyses

Statistical Parametric Mapping 8 (SPM8; http://www.fil.ion.ucl.ac.uk/spm/) and the VBMM toolbox were used to analyse data in Matlab. Each MRI sequence was visually inspected for quality and reoriented along the anterior-to-posterior commissure. Thereafter, images were segmented into grey matter, white matter and cerebrospinal fluid components. The diffeomorphic anatomical registration through exponentiated lie algebra (DARTEL) was applied to the grey matter images for spatial normalisation. To control for individual whole-brain volume differences, we implemented non-linear modulation using the Jacobian determinants derived from the normalisation process. The modulated grey matter images were smoothed with an 8 mm full-width-at-half-maximum Gaussian kernel.
Relationship between Functional Neurological Symptoms and Childhood Abuse (N=23)

Figure 1 Scatter plots of the positive associations between functional neurological symptoms and the magnitude of previously experienced childhood abuse. Also shown is the scatter plot of the positive correlation between the Conversion Disorder subscale of the Screening for Somatoform Symptoms-7 scale and the Patient Health Questionnaire-15. Spearman correlation coefficients are displayed.

After preprocessing, SPM-based multiple regression was used to examine associations between covariates of interest and grey matter volumes with age and gender as nuisance variables. Separate regression analyses were performed for PHQ-15, SOMS:CD, CTQ-Total, CTQ-Abuse, CTQ-Neglect, LEC-5 ‘happened to me’ score, PCL-5 total, and the four PCL-5 subscores. To consider gender effects, we also performed all analyses for the 18 women, controlling for data. Data were first reviewed at an uncorrected p≤0.001 and a cluster extent threshold of 50 voxels. Whole-brain corrections for multiple comparisons at the peak voxel level used a family-wise error (FWE) rate of p<0.05. Given our specific a priori hypotheses for cingulo-insular cortices, we defined bilateral regions of interest (ROIs) in the ACC and insula using the WFU Pickatlas to perform small-volume corrections (SVC). Since there is an extensive literature linking stress-related neuroplasticity to the hippocampus, bilateral hippocampi were also an ROI. A FWE rate of p<0.05 was used for SVCs at the peak-voxel level. To control for potential confounding effects of depression and anxiety, we performed separate analyses for statistically significant findings with BDI or STAI-T scores entered as nuisance variables.

RESULTS

Associations between FND symptoms and childhood abuse

Statistically significant associations were observed between PHQ-15 and CTQ-Abuse scores (p=0.001; Spearman correlation coefficient=0.642), and between SOMS:CD and CTQ-Abuse scores across all patients (p=0.003; Spearman correlation coefficient=0.587) (figure 1, see online Supplemental figure 1). Positive associations were also appreciated between PHQ-15 scores and SOMS:CD scores (p=0.018; Spearman correlation coefficient=0.488). See online Supplemental table 2 for clinical scores.

Grey matter associations with FND symptoms and childhood abuse

For the entire FND cohort, only a trend-level association was present between PHQ-15 scores and decreased left anterior insular volumes (p_{uncorrected}<0.001). A negative trend association was also observed between CTQ-Abuse scores and left anterior insular volumes (p_{uncorrected}<0.001) (see online Supplemental figure 2). In women, these relationships showed separate statistically significant inverse relationships between PHQ-15 (p_{corr}=0.032), SOMS:CD (p_{corr}=0.038), CTQ-Abuse (p_{corr}=0.021) scores and left anterior insular grey matter volumes (figure 2, table 1). Regression analyses secondarily controlling for STAI-T scores remained statistically significant, although these associations did not hold controlling for BDI scores (see online Supplemental tables 3, 4).

Given the statistically significant parallel associations of functional neurological symptom severity and childhood abuse burden each associated with left anterior insular volume reductions in women, we conducted two separate post-hoc SPM-based regression analyses to investigate the potential relationship between these findings. When controlling for CTQ-Abuse scores and age in regression analyses, associations between PHQ-15 or SOMS:CD scores and left anterior insular grey matter volume did not remain statistically significant.

Grey matter associations with PTSD symptoms and lifetime adverse events

For the entire FND cohort, PCL-5 hyperarousal subscores were negatively associated with dorsal ACC (p_{uncorr}=0.014) grey matter volumes. This finding remained negatively associated with hyperarousal controlling for BDI (p_{corr}=0.001) and STAI-T (p_{corr}=0.006). For women, PCL-5 avoidance subscores were negatively associated with dorsal/perigenual ACC (p_{corr}=0.046) grey matter volumes. This inverse association also remained statistically significant when controlling for BDI (p_{SVC}=0.012) and STAI-T (p_{SVC}=0.016).

Analyses probing associations with cumulative lifetime adverse events across all FND subjects showed an inverse association between the LEC-5 ‘happened to me’ score and left hippocampal volume (p_{corr}=0.010). This association held controlling for BDI (p_{corr}=0.017) and STAI-T (p_{corr}=0.015). In women, a similar negative association between the LEC-5 ‘happened to me’ score and left hippocampal volume was appreciated (p_{corr}=0.005). This finding also held, controlling for BDI (p_{corr}=0.009) and STAI-T (p_{corr}=0.006). See online Supplemental tables 3–5, figure 3 and online Supplemental figure 3 for complete findings.

DISCUSSION

Consistent with our a priori hypotheses, women with FND showed significant associations between functional neurological symptoms and reduced left anterior insular grey matter volumes. In women, the magnitude of experienced childhood abuse was also associated with decreased left anterior insular volume. Post-hoc analyses showed that the relationship between functional neurological symptom severity and decreased insular volume was explained mainly by the magnitude of reported childhood abuse. These findings lend early support for potential childhood abuse-related aberrant neuroplasticity in the pathophysiology of FND. In addition, while similar trend-level associations were observed, there were no statistically significant
associations between functional neurological symptoms, childhood abuse and left anterior insular volume in the mixed-gender FND cohort controlling for age and gender. These results suggest the potential for differential brain-trauma relationships across men and women. Also, in our entire mixed-gender FND cohort, PTSD symptom severity was associated with reduced ACC grey matter volumes. Lastly, consistent with the literature detailing associations between lifetime trauma burden and hippocampal volume, we observed an inverse relationship between the number of cumulative adverse events and left hippocampal volume across all FND subjects.

The insula, a paralimbic multimodal region implicated in viscerosomatic, visceromotor, homeostatic, affective and cognitive processes, is well positioned to play a central role in the multiplicity of symptoms experienced by patients with FND. Information from the lamina I spinothalomocortical pathway conveying physiological information from bodily tissues projects onto the posterior insula and ACC. Given this convergence of information, the posterior insula mediates the interoceptive representation of the physiological status of the body. Affectively and motivationally valenced information from the amygdala, ACC and orbitofrontal cortex synapse onto the mid-insula, with the left mid-insula mediating parasympathetic activity and the right mid-insula mediating sympathetic tone. Craig has suggested that a higher-order synthesis of this information occurs in the anterior insula, and this area may be critical for emotional and self-awareness. Given its structural and functional connectivity, the anterior insula is also implicated in the integration of internal feeling states, the detection and processing of differences between observed and expected bodily states, and the pathophysiology of mood and anxiety disorders.

Figure 2  Left anterior insula grey matter volume negatively correlated with functional neurological symptoms and previously experienced childhood abuse in women with functional neurological disorder. (A) Patient Health Questionnaire-15 scores were inversely associated with left anterior insular grey matter volume (psvc=0.032; peak z-score=−3.93). (B) Conversion Disorder subscale of the Screening for Somatoform Symptoms-7 scale scores were also inversely correlated with left anterior insular grey matter volume (psvc=0.038; peak z-score=−3.87). (C) The magnitude of childhood trauma as measured by the Childhood Trauma Questionnaire showed a parallel negative association with left anterior insular grey matter volume (psvc=0.021; peak z-score=−4.04). Images are thresholded at a voxel-wise uncorrected p value of 0.005 for visualisation purposes. To display in the scatter plots a measure of grey matter volume for visualisation purposes only, we used the MarsBar toolbox (http://marsbar.sourceforge.net/) to extract mean grey matter volumes for each subject at a 5 mm sphere centred at the peak coordinate of statistically significant findings.

Table 1

<table>
<thead>
<tr>
<th>n</th>
<th>Measures of interest</th>
<th>Cerebral regions (Brodmann area)</th>
<th>Peak coordinates in MNI space (mm)</th>
<th>Peak voxel z-score</th>
<th>p Value (SVC)</th>
<th>Cluster extent (mm³)</th>
</tr>
</thead>
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<tr>
<td>23</td>
<td>PTSD-Hyperarousal</td>
<td>Dorsal ACC (24)</td>
<td>0 x 30 y 24 z</td>
<td>−3.97</td>
<td>&lt;0.001 (0.014)</td>
<td>229</td>
</tr>
<tr>
<td></td>
<td>LEC—happened to me</td>
<td>L hippocampus (20)</td>
<td>−33 x −27 y −12 z</td>
<td>−4.01</td>
<td>&lt;0.001 (0.010)</td>
<td>176</td>
</tr>
<tr>
<td>18</td>
<td>PHQ-15</td>
<td>L anterior insula (48)</td>
<td>−40 x 12 y −11 z</td>
<td>−3.93</td>
<td>&lt;0.001 (0.032)</td>
<td>169</td>
</tr>
<tr>
<td></td>
<td>SOMS:CD</td>
<td>L anterior insula (48)</td>
<td>−42 x 9 y −12 z</td>
<td>−3.87</td>
<td>&lt;0.001 (0.038)</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>CTQ-Abuse</td>
<td>L anterior insula (48)</td>
<td>−39 x 9 y −12 z</td>
<td>−4.04</td>
<td>&lt;0.001 (0.021)</td>
<td>156</td>
</tr>
<tr>
<td></td>
<td>PTSD-Avoidance</td>
<td>R dorsal/perigenual ACC (24)</td>
<td>2 x 32 y 24 z</td>
<td>−3.66</td>
<td>&lt;0.001 (0.046)</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>LEC—happened to me</td>
<td>L hippocampus (20)</td>
<td>−33 x −27 y −12 z</td>
<td>−4.27</td>
<td>&lt;0.001 (0.005)</td>
<td>230</td>
</tr>
</tbody>
</table>

Small-volume corrected (SVC) findings are listed in parentheses. ACC, anterior cingulate cortex; CTQ, Childhood Trauma Questionnaire; LEC, Life Events Checklist; MNI, Montreal Neurological Institute; PHQ-15, Patient Health Questionnaire-15; PTSD, post-traumatic stress disorder; SOMS:CD, Conversion Disorder subscale of the Screening for Somatoform Symptoms-7 scale. PTSD subscores are based on the PTSD Checklist for DSM-5.
Grey Matter Volume Associations with PTSD Symptoms and Lifetime Trauma Burden

(A) Across all subjects, hyperarousal as measured by the PTSD Checklist for DSM-5 was negatively associated with dorsal anterior cingulate cortex (ACC) grey matter volume ($p_a=0.014$, peak z-score=$-3.97$). (B) In women with FND, avoidance as measured by the PTSD Checklist for DSM-5 was negatively associated with dorsal/perigenual ACC grey matter volume ($p_a=0.046$, peak z-score=$-3.66$). (C) Across all subjects, lifetime adverse events as measured by the ‘happened to me’ item of the Life Events Checklist-5 (LEC) was negatively associated with left hippocampal grey matter volume ($p_a=0.010$, peak z-score=$-4.01$). Images are thresholded at a voxel-wise uncorrected p value of 0.005 for visualisation purposes. To display in the scatter plots a measure of grey matter volume for visualisation purposes only, we used the MarsBar toolbox (http://marsbar.sourceforge.net/) to extract mean grey matter volumes for each subject at a 5mm sphere centred at the peak coordinate of statistically significant findings.

While the insula is well positioned to play a role in the pathophysiology of FND, the ACC is also a paralimbic region implicated in the integration of affective, viscerosomatic and cognitive processes. Meta-analyses have shown that the dorsal ACC is a convergent zone for negative emotional processing, pain processing and cognitive control. A dorsal-ventral gradient exists in the ACC whereby dorsal regions are interconnected to lateral prefrontal and premotor regions and implicated in appraisal and behavioural expression of mood states. Perigenual and subgenual ACC subregions are reciprocally connected to the amygdala and facilitate top-down regulation of affective states. As detailed in our neurobiological framework for somatosensory processing and cognitive control, the cognitive-affective neuroscience literature has demonstrated cingulo-insular involvement in mediating aberrant somatic experiences through negative expectation bias, alexithymia, pain catastrophising and heightened visceral-somatic processing during negative mood states. We have also postulated the construct of ‘neural functional unawareness’ as a pathophysiological model for FND, which implicates cingulo-insular abnormalities in mediating impaired emotional and interoceptive awareness. Our findings support selective left anterior insular volume reductions pertaining to functional neurological symptoms, while dorsal and perigenual ACC grey matter reductions are related to PTSD symptom severity in patients with FND. We theorise that inverse associations between functional neurological symptoms and left anterior insular grey matter represent a neurobiological substrate for the impaired integration of affective, viscerosomatic and cognitive processing in patients with FND leading to deficits in emotional and bodily awareness.

We also observed an inverse relationship between childhood abuse burden and left anterior insular volume in women with FND. Post-hoc analyses showed that the relationship between insular volume and functional neurological symptoms in women could be explained by the magnitude of endorsed childhood abuse. This suggests that our parallel observations of functional neurological symptoms and childhood abuse each linked to reduced insular volume are inter-related. Given that the insula is also broadly implicated in the pathophysiology of mood and anxiety disorders, it is notable that the parallel associations between symptom severity and childhood abuse burden each associated with left anterior insular volume reductions remained significant when controlling for trait anxiety but not for depression scores. This implies a potential role for the left anterior insula in the convergence of functional neurological symptoms and negative mood in patients with FND. As shown in figure 1 and online Supplemental figure 1, individuals with more childhood abuse reported increased functional symptoms as measured by both the PHQ-15 and the SOMS:CD scales. These observations are consistent with the literature detailing relationships between adverse early-life events and functional neurological symptom severity. While requiring replication, our results suggest that aberrant neuroplastic changes in the left anterior insula potentially driven by childhood abuse may underlie aspects of the pathophysiology for FND.

Our findings can be contextualised based on basic science and human literature detailing maladaptive experience-dependent neuroplastic changes following chronic stress. Animal models of chronic stress have shown that the medial prefrontal cortex and the hippocampal CA3 region undergo dentritic...
spine density reductions following prolonged stress exposures. In a VBM study of 130 healthy individuals, cumulative lifetime adverse events were associated with reduced medial prefrontal, insular and subgenual ACC volumes. Adverse early-life events have been specifically associated with smaller ACC, insular, orbitofrontal cortex, caudate and hippocampal volumes. A meta-analysis in psychiatric populations with childhood maltreatment also demonstrated reduced insular, orbitofrontal, parahippocampal, amygdalar, middle temporal, inferior frontal and post-central gyri grey matter in patients compared with controls. Our finding of an association between cumulative adverse life events and reduced left hippocampal volume adds to the literature linking trauma burden to hippocampal volumes across several psychopathologies. Given that our findings also suggest specific grey matter-symptom severity relationships, the effects of trauma type, age of onset, trauma burden, gender effects and gene by experience relationships should be further studied in future investigations. These subsequent research efforts would help clarify the degree to which experience-dependent neuroplasticity increases a general vulnerability to the development of FND and suggests potential biological convergence between the magnitudes of experienced childhood abuse, functional neurological symptoms and reduced left anterior insular volume.

There are a few potential limitations to this study. Our FND cohort had mixed symptoms including medically unexplained abnormal movements, weakness, non-epileptic seizures, pain, fatigue and non-dermatomal sensory deficits. While we note that there have been increasing calls for greater research integration across FNDs, we acknowledge that this is debated in the field and that this study is underpowered to include FND subtypes as nuisance variables. As is common in FND, our cohort also had significant mood and anxiety comorbidities, and the majority were on psychoactive medications, which are potential confounds. It is possible that our FND cohort recruited across an integrated behavioural neurology-neuropsychiatry clinic could tend towards a population with greater psychopathology compared with FND patients recruited from general neurology settings. We note that patients with current major depressive disorder have been removed by some groups from study inclusion; however, it is our viewpoint that the frequent co-occurrence of depression and anxiety in FND likely reflects aspects of a shared pathophysiology. Comorbid psychiatric disorders are the rule, not the exception, in patients with FND. Inclusion of subjects with prominent psychiatric comorbidities is more representative of the FND spectrum and increases the external validity of our sample to clinical practice. Furthermore, a strength of this study is our ability to secondarily control for depression or anxiety. It is also important to highlight that there are multiple risk factors for FND apart from adverse life events (and not all patients endorse past traumatic experiences), suggesting that more research is needed to investigate brain-disease risk relationships. Also, additional research is needed to clarify if the overlapping brain-symptom severity and brain-childhood abuse relationships appreciated in this study reflect gender-specific neural mechanisms of disease or rather, in part, reflect the increased frequency of childhood abuse in women compared with men (see online Supplemental table 2).

Our reliance on self-report measures of symptom severity (PHQ-15, SOMS:CD) may also not fully capture the degree of physical disability in a given FND patient, and PHQ-15 scores may not reliably differentiate FND from other neurological populations. The lack of a healthy control group could be seen as a limitation; however, this addition would not aid the study of functional neurological symptoms as continuous variables, and the over-reliance on healthy subject group comparisons has recently been criticised in neuropsychiatric research.

The neurobiology of FND spans multiple networks, particularly given that right temporoparietal junction functional alterations in functional movement disorders have been linked to deficits in motor intention awareness and self-agency. In addition, it is likely that ACC alterations play important roles in predisposing factors such as alexithymia and dissociation. More research is needed to investigate biological similarities and differences in the pathophysiology of FND across gender, as well as to clarify if the observed brain-symptom relationships represent disease-related or compensatory structural alterations.

In conclusion, we used a trans-diagnostic, symptom-based and disease-risk approach to characterise inverse associations between left anterior insular volume and indices of functional neurological symptoms and childhood abuse in patients with FND. We also observed differential cingulo-insular structural alterations, delineating inverse associations between PTSD symptoms and ACC volume. This study, while requiring replication, may advance our understanding of the neuropsychopathophysiology of FND and suggests potential biological convergence between the magnitudes of experienced childhood abuse, functional neurological symptoms and reduced left anterior insular volume.

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