Neurostructural brain imaging study of trait dissociation in healthy children

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Background
Trait dissociation has not been examined from a structural human brain mapping perspective in healthy adults or children. Non-pathological dissociation shares some features with daydreaming and mind-wandering, but also involves subtle disruptions in affect and autobiographical memory.

Aims
To identify neurostructural biomarkers of trait dissociation in healthy children.

Method
Typically developing 9- to 15-year-olds (n = 180) without psychological or behavioural disorders were enrolled in the Developmental Chronnecto-Genomics (DevCoG) study of healthy brain development and completed psychological assessments of trauma exposure and dissociation, along with a structural T1-weighted magnetic resonance imaging. We conducted univariate ANCOVA generalised linear models for each region of the default mode network examining the effects of trait dissociation, including scanner site, age, gender and trauma as covariates and correcting for multiple comparison.

Results
We found that the precuneus was significantly larger in children with higher levels of trait dissociation but this was not related to trauma exposure. The inferior parietal volume was smaller in children with higher levels of trauma but was not related to dissociation. No other regions of interest, including frontal and limbic structures, were significantly related to trait dissociation even before multiple comparison correction.

Conclusions
Trait dissociation reflects subtle cognitive disruptions worthy of study in healthy people and warrants study as a potential risk factor for psychopathology. This neurostructural study of trait dissociation in healthy children identified the precuneus as an essential brain region to consider in future dissociation research.

Keywords
Default mode network; mind-wandering; precuneus; magnetic resonance imaging; volumetrics.

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Daydreaming, mind-wandering and the default mode network

To that aim, Giesbrecht and colleagues7 reviewed cognitive processes in dissociation and found that fantasy proneness (including daydreaming), suggestibility and subtle cognitive failures such as a lapsed attention account for a significant proportion of the variance in dissociation. Butler described normative dissociation as a ‘forum for mental processing’,2,3 that is a passive, spontaneous absorptive experience, typically termed daydreaming, that occurs in the absence of environmentally cued cognitive demands. In her foundational article on normative dissociation, Butler10 noted that much of our stream of consciousness is filled with dissociative experiences such as daydreaming and fantasy involving a temporary separation from other mental processes. Therefore, considering constructs that have been operationally defined in previous neuroimaging research, we determined that trait dissociation shares notable commonalities with daydreaming, mind-wandering and undirected thought, all of which have been associated with default mode network (DMN) activation.2,25 The DMN is a network of brain regions with highly correlated mental activity when a person is awake but not engaged in task-focused work.2,25 Past research reports enhanced dynamic DMN functional connectivity and activity during daydreaming and mind-wandering,28 and DMN connectivity was recently implicated as a potential predictor of trauma-related dissociation after controlling for psychological symptoms and trauma.29 which reinforced our selection of the DMN.

Study aims

To date, brain regions structurally related to the specific concept of trait dissociation in healthy people are unknown – and very little is known about brain regions involved in any type of dissociation in child and adolescent samples. Our study goal was to identify potential neurostructural correlates of non-pathological trait dissociation in healthy children. To this aim, we designed an exploratory volumetric study of all regions in the DMN, including the hippocampus. We controlled for age, gender, scanner site and, most importantly, trauma exposure. Trauma exposure was evaluated as a covariate of interest to examine its potential interaction with dissociation in this sample.

Method

Participants

Typically developing children aged 9 to 15 years were enrolled in our Developmental Chronnecto-Genomics (DevCoG)30 study of healthy brain development after obtaining parental permission and consent and participant assent. Children completed psychological assessments and underwent structural magnetic resonance imaging (MRI) (n = 183). Three children were excluded because of unusable data, giving a final sample of n = 180. The sample was evenly distributed by gender, with 94 males and 86 females, by study site, with 89 at the University of Nebraska Medical Center (UNMC) and 91 at the Mind Research Network (MRN), and by age, with an average age of 11.97 years (s.d. = 1.73). Children were excluded from the study if parents reported that their child ever had a diagnosis of any psychiatric or behavioural disorder, a history of traumatic brain injury or other neurological condition, or the presence of metallic implants (e.g. orthodontia). The study was approved by both study sites’ institutional review boards (IRBs) and all research was conducted according to ethical principles including obtaining fully informed written parental consent and child assent. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. All procedures involving human participants/patients were approved by the IRBs at the University of Nebraska Medical Center in Omaha, Nebraska, USA (UNMC IRB #503-15-EP) and at the Mind Research Network in Albuquerque, New Mexico, USA.

Psychological measures

The Trauma Symptom Checklist for Children (TSCC)31 is a self-report measure for children aged 8–16, with scores for five clinical scales, of which we used the TSCC dissociation scale. This 10-item self-report scale includes statements such as ‘Trying not to have any feelings’, ‘Pretending I’m somewhere else’, ‘My mind going empty or blank’ rated on a Likert scale from 0 (never) to 3 (almost all of the time). The Cronbach’s alpha for the dissociation scale in this sample was very good, at 0.80. Providing evidence of convergent validity, the TSCC dissociation scale correlates highly in adolescent samples with the Adolescent Dissociative Experiences Scale (e.g. r = 0.79).32

We used a modified version of the UCLA Trauma History Profile33 to assess the number of traumatic life events encountered by children in the study. Children answered ‘yes’ or ‘no’ to whether they had experienced each of 12 potentially traumatic events. We shortened the original 15 event measures to exclude items about sexual abuse or physical abuse that occurred specifically in the home, so that participation would be considered low risk by the IRBs. Both personally experiencing violence and witnessing violence
to family members were still assessed. The items used in the current study were: having someone close to them die; being hit, punched or kicked very hard; seeing a family member hit, punched or kicked very hard; seeing or hearing about violence to a loved one; being a victim of community violence; being in a war; being in a disaster; being in a bad accident; having a painful or scary medical procedure; seeing a dead body not at a funeral; and having anything else very scary or upsetting happen.

**Structural T1-weighted MRI**

Structural T1-weighted MRI images were acquired using a Siemens 3-Tesla Skyra (at UNMC) or a Siemens 3-Tesla TRIO (at MRN), both with 32-channel head coils and closely calibrated sequences. A three-dimensional magnetisation-prepared rapid gradient-echo (3D MP-RAGE) sequence was used with the following parameters: repetition time TR = 2400 ms; echo time TE = 1.94 ms; flip angle 8°; field of view FOV = 256 mm; slice thickness 1 mm; base resolution 256; 192 slices; voxel size 1.0 × 1.0 × 1.0 mm. The T1-weighted structural brain images of all participants were processed using the Freesurfer software version 5.3 on a Linux Ubuntu platform (http://surfer.nmr.mgh.harvard.edu). Regional volumes were computed for the automatic cortical parcellation (aparc)34 and automatic subcortical segmentation (aseg)35 atlases in Freesurfer. We followed the ENIGMA protocol for quality assurance, which included performing visual checks of all cortical segmentations (http://enigma.usc.edu/protocols/imaging-protocols) and checking for motion, among other artifacts. Participants whose MRI images had large motion artifacts were excluded (n = 3). In addition, histograms of all regional values were computed for visual inspection. All volumes were normalised by dividing each regional volume by the total intracranial volume (TIV) per participant, to avoid the bias of head size in the volumetric measurements.36

**Statistical analysis**

The DMN regions of interest were: the caudal middle frontal, hippocampal, inferior parietal, isthmus cingulate, medial orbital frontal, parahippocampal, posterior cingulate, precuneus, and rostral anterior cingulate regions. For each of the nine DMN regions of interest, we conducted a univariate ANCOVA generalised linear model (GLM). Each of the models utilised the TIV-corrected volume averaged across the brain hemispheres for each structure of the DMN, which was entered as a dependent variable per model. TSCC dissociation score was the independent variable, and scanner site (UNMC or MRN), age, gender (male or female) and number of traumatic events experienced were entered as covariates. Full models were corrected using the Benjamini–Hochberg false discovery rate multiple comparison correction (denoted as BH in corrected p-values) as implemented in the p.adjust function in R version 1.2.5019 on a Windows 10 platform.

**Results**

TSCC dissociation scale scores in the full sample (n = 180) ranged from 0 to 19 and averaged 5.33 (s.d. = 4.02). Both the median and modal dissociation scores were 4.00, and the sample dissociation scores had acceptable skewness and kurtosis. The pathological cut-off scores for the TSCC dissociation scale varied slightly by age and gender but centred around a score of 15.33 Importantly, less than 4% of our sample scored above the pathological cut-off score, supporting the non-pathological nature of dissociation in our sample. The most strongly endorsed dissociation items were daydreaming, forgetfulness, my mind going empty or blank, and going away in my mind/trying not to think. Trauma exposure scores ranged from 0 to 7 (mean 2.17, s.d. = 1.81), with the most commonly endorsed traumas including the death of a loved one, the violent or serious injury of a loved one, and being a victim of physical violence. As expected, dissociation and trauma exposure scores were significantly correlated (r = 0.36, p < 0.001).

We conducted univariate ANCOVAs for each cortical structure of the DMN using the aparc atlas34 (Fig. 1) and added the subcortical hippocampus defined by the aseg atlas35 for completeness. Levene’s test of equality showed no significant differences in error variance in any model (p = 0.200–0.798). Significant effects were found in the precuneus (F(5,174) = 4.83, pBH = 0.003, R2 = 0.12) and inferior parietal lobes (F(5,174) = 3.98, pBH = 0.008, R2 = 0.10), and these results survived multiple comparison correction.

**Fig. 1** Cortical default mode network regions included in this study and labelled by the (aparc) atlas. Note that we also included the hippocampus, not shown in this figure.
In the precuneus, dissociation significantly predicted TIV-corrected volumes ($F_{1,174} = 3.05, p_{BH} = 0.002$) above and beyond the covariates in the model, such that greater volumes were found in those with higher dissociation values (Table 1). Age was a significant predictor in this model, such that older age was associated with smaller TIV-corrected volumes, but no other covariate significantly predicted precuneus volumes. In the inferior parietal region, trauma but not dissociation significantly predicted TIV-corrected volumes, but no other covariate significantly predicted precuneus volumes in this region were not significantly related to trauma. We found no non-pathological dissociation effect and decreased volumes in this region were not significantly related to trauma. We noted in previous trauma research. Interestingly, a study found reduced hippocampal volume.48

In pathological dissociation, including absorption in one’s internal world and alterations in one’s experience of self or others,45 so finding larger precunei in the current study is intriguing.

Functionally, the precuneus is known to be involved in episodic memory retrieval, mental imagery, self-referential processing tasks, perspective taking and consciousness.39 Interestingly, a study comparing individuals with DID with actors simulating DID found higher resting-state metabolism in regions of the DMN, including the precuneus, in those with DID, suggesting that they were more involved in self-referential thought than the actors during rest.40

The inferior parietal region is an important association area typically involved in internal sensory processing41 and it is also active during rest, especially during self-referential thought, along with the precuneus.42 Nardo and colleagues43 studied dissociation in a traumatised sample and controlled for the effects of psychological symptoms and trauma exposure. Contrary to our results, which found no non-pathological dissociation effect and decreased volumes with trauma exposure in the inferior parietal region, they found that both pathological and trait dissociation were associated with volumetric increases in this region.

Although negative results should be discussed with great caution, we believe it is important to note that the current study did not find any significant volumetric differences in the examined frontal,43-45 hippocampal46,47 or parahippocampal44,47 regions noted in previous trauma research. Interestingly, a study found that reduced hippocampal volume was related to both severity of dissociative symptoms and trauma exposure in people with PTSD and DID (who also met criteria for PTSD),46 however, a rare study of individuals with dissociative disorders who did not meet criteria for PTSD did not find reduced hippocampal volume.48

Perhaps structural findings related to diminished medial temporal and increased frontal volumes become apparent over the course of disorder development, or perhaps these regions are not implicated in non-pathological trait dissociation. These questions are clearly a matter for future research. For now, our absence of frontal and medial temporal findings – an absence noted even before multiple comparison correction – clearly separates our findings on trait dissociation from most work on pathological dissociation related to trauma. Although we are careful in drawing conclusions from negative findings, our results clearly call for future imaging research on dissociation in psychologically healthy individuals to avoid the trauma confound and isolate structural and functional origins of the process of normative dissociation.

| Table 1 | ANCOVA results for precuneus and inferior parietal regions of the DMN |
|---|---|---|---|---|
| Predictors | Estimates | 95% CI | $t$ | $p$ | Partial $\eta^2$ |
| Precuneus | | | | | |
| Intercept | 0.00934 | 0.00845 to 0.01023 | 20.78626 | <0.001 | 0.725 |
| Site | 0.00007 | −0.00017 to 0.00032 | 0.60292 | 0.547 | 0.002 |
| Age | −0.00012 | −0.00019 to −0.00005 | −3.49982 | 0.001 | 0.066 |
| Gender | −0.00006 | −0.00027 to 0.00018 | −0.46998 | 0.639 | 0.001 |
| Trauma history | −0.00005 | −0.00012 to 0.00002 | −1.27660 | 0.203 | 0.009 |
| Dissociation | 0.00005 | 0.00002 to 0.00008 | 3.05445 | 0.003 | 0.061 |
| Inferior parietal lobes | | | | | |
| Intercept | 0.01411 | 0.01273 to 0.01549 | 20.18396 | <0.001 | 0.713 |
| Site | 0.00016 | −0.00022 to 0.00054 | 0.82065 | 0.413 | 0.004 |
| Age | −0.00019 | −0.00029 to −0.00008 | −3.44657 | 0.001 | 0.064 |
| Gender | −0.00011 | −0.00048 to 0.00026 | −0.57094 | 0.569 | 0.002 |
| Trauma history | −0.00015 | −0.00026 to −0.00004 | −2.66851 | 0.008 | 0.039 |
| Dissociation | 0.00003 | −0.00002 to 0.00008 | 1.08488 | 0.279 | 0.007 |

**Bold denotes significant results at $p < 0.01$.**

**Discussion**

To the best of our knowledge, this is the first structural MRI study of brain regions associated with non-pathological dissociation in healthy children. Our most important finding is that the precuneus was larger in children with higher levels of trait dissociation and that volumes in this region were not significantly related to trauma. We also found that the inferior parietal region was smaller in healthy children who had experienced more traumatic events, but this region was not associated with trait dissociation. Research on trait dissociation in healthy people is lacking, however, a meta-analysis of 24 functional neuroimaging studies of the relatively similar concept of mind-wandering38 identified the importance of the precuneus/posterior cingulate cortex during this mental state. Similarly, our results are bolstered by two reviews of neuroimaging studies of pathological dissociation. Roydeva & Reinders11 concluded that posterior association areas were functionally relevant in pathological dissociation, and specifically called for more research on the precuneus. In a review of pathological dissociation in borderline personality disorder, Krause-Utz et al45 explicitly suggested the importance of the precuneus owing to its role in self-referential processing. Known functions of the precuneus38 are consistent with the phenomenological experience of trait dissociation, including absorption in one’s internal world and alterations in one’s experience of self or others,45 so finding larger precunei in the current study is intriguing.
In identifying some – but not unmitigated – overlap between brain areas implicated in pathological and non-pathological dissociation, and given the subtle cognitive errors noted in trait dissociation, our findings align with Loewenstein’s suggestion that studying dissociation may address puzzling gaps in psychology and neuroscience. These gaps are relevant for understanding not only cognitive function in general, but also pathological disorders because research suggests that mild cognitive impairments may pose a potential risk for PTSD or dissociative disorder in healthy people with high dissociation. Such risks are consistent with previous research associating alterations in the DMN with psychopathology in general, and specifically structural variations in the precuneus with subclinical symptoms of clinical disorders.

Limitations and future research

Despite the novel contribution of the current study, it has limitations. The exploratory research utilised structural MRIs in healthy children. By design, our study did not include a sample of children diagnosed with dissociative disorders. Although such a sample would certainly be smaller, replicating this study in children with dissociative disorders would improve understanding of potential structural alterations related to pathological versus non-pathological dissociation. Future neurofunctional research should include blood oxygen level-dependent (BOLD) or arterial spin labelling (ASL) perfusion or magnetencephalography resting-state studies and functional assessments specifically targeting the precuneus in healthy child and adult samples and pathological samples. Finally, our analyses are cross-sectional, so future work should examine how longitudinal changes in dissociation correspond with changes in brain structure and function. Longitudinal follow-up should also assess whether trait dissociation relates to risk for psychopathology in the current sample. Such future work would allow us to evaluate the extent to which normative dissociation is a risk or protective factor during development.

Implications

We hope our findings on non-pathological trait dissociation in children may inform brain mapping research and also inform research on transdiagnostic symptoms of pathological dissociation, independent of psychiatric diagnoses and trauma exposure. We believe the key implications of this study are the identification of a previously unknown psychological function of the precuneus and the suggestion of this structure as a promising target for future neuropsychological and psychopathological research in healthy and psychopathological groups.

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References