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Resting-state functional connectivity in women with PMDD

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Abstract

Background: Premenstrual dysphoric disorder (PMDD) is an understudied, debilitating disorder of women. Given evidence for prefrontal cortical and limbic dysfunction in PMDD, we compared intrinsic connectivity of the executive control network (ECN), default mode network (DMN), and amygdala in women with PMDD vs. controls.

Methods: Thirty-six women (18 PMDD, 18 control) participated in fMRI during the follicular and luteal phases of the menstrual cycle. At each time, resting-state functional connectivity was evaluated both before and after participants performed an emotion regulation task. The ECN was identified using independent components analysis, and connectivity of left and right amygdala seeds was also evaluated.

Results: Nonparametric permutation testing identified a cluster in the left middle temporal gyrus (MTG) with significantly stronger connectivity to the left ECN in women with PMDD vs. controls in all four fMRI sessions. Women with PMDD exhibited no difference in functional connectivity between menstrual cycle phases. Amygdala connectivity did not differ between the groups but differed significantly with menstrual phase, with left amygdala connectivity to cingulate cortex being significantly stronger during the follicular vs. luteal phase. Right amygdala connectivity to the middle frontal gyrus was also stronger during the follicular vs. luteal phase, with no group differences. These findings suggest that women with PMDD have different intrinsic network dynamics in the left executive control network compared to healthy controls.

Introduction

Premenstrual dysphoric disorder (PMDD) is a severe variant of premenstrual syndrome, characterized by debilitating behavioral symptoms, including dysphoric mood, which occur during the luteal phase of the menstrual cycle and abate following the onset of menstruation, causing considerable impairments in quality of life. Brain imaging studies of task-related activation during fMRI have identified several regions of abnormal function in women with PMDD. During the symptomatic phase, women with PMDD show greater reactivity of the amygdala to negative stimuli and weaker top-down control of this activation compared to healthy controls. Consistent with this observation, lower pre/postcentral gyrus activation and lower dorsolateral prefrontal cortical (dLPFC) activity have been observed when women with PMDD perform an emotion regulation task during the symptomatic phase of the menstrual cycle compared to the asymptomatic phase. The latter finding generally supports previous evidence linking dLPFC activation to the etiology of PMDD, although that evidence was obtained using an ovarian suppression plus add-back hormone protocol, in which women with PMDD had greater dLPFC activation compared to healthy controls while performing a working memory task. Inasmuch as the dLPFC is strongly linked to emotion regulation (for meta-analyses,
see refs. 10 and 11) dysfunction in this brain region could plausibly lead to problems regulating emotions such as those described by women with PMDD12.

Resting-state functional connectivity increasingly has been measured to improve understanding of neuropsychiatric disorders13. Patients with Major Depressive Disorder (MDD) exhibit stronger connectivity between the subgenual cingulate cortex and the default mode network (DMN)14, and between the insula and amygdala as compared with healthy controls15. A meta-analysis extended these reports, documenting stronger DMN intra-network connectivity and weaker frontoparietal intra-network connectivity in MDD patients than in healthy control subjects16. PMDD shares a number of characteristics with MDD, including overlapping diagnostic criteria and comparable impairments in quality of life17. The neural features that have been most strongly linked to MDD and other affective disorders have not yet been observed in women with PMDD, and vice versa – but lack of evidence that these neural features overlap cannot be taken as proof that they do not. Some preliminary forays have investigated the neural features of menstrual-related mood disorders, and differences in functional connectivity between women with Premenstrual Syndrome (PMS, a milder syndrome than PMDD) and healthy controls have been observed. Women with PMS have stronger amygdala-prefrontal cortical connectivity, and connectivity between the right amygdala and right precentral gyrus, left ACC, and left medial prefrontal cortex correlates positively with the strength of symptoms in these patients18. A network-level analysis using independent component analysis (ICA) to identify the DMN found stronger connectivity between the DMN and both the superior temporal gyrus and precentral gyrus in women with PMS as compared to healthy controls19. These reports suggest that network connectivity may differ in women with PMDD compared with healthy controls, and that these differences may point to potential therapeutic targets.

Yet resting-state functional connectivity studies in women with PMDD per se (rather than PMS) had not been performed, and studies in women with milder forms of premenstrual disorders may not be informative regarding PMDD. We therefore compared resting-state functional connectivity in women with PMDD and healthy controls during the premenstrual and follicular phases of the menstrual cycle. Resting-state fMRI scans were performed once before and again after participants completed an emotion regulation task (see Petersen et al.8 for details of task methodology and findings). Given previous evidence linking abnormal task-related activation in prefrontal cortical regions to PMDD85, we selected the executive control network (ECN) as a network of interest. Amygdala and DMN connectivity were assessed due to evidence that menstrual phase20, ovarian hormones21, and PMDD22 may all influence amygdala function, and evidence that DMN connectivity is abnormal in major depression, which shares many symptoms with PMDD23,24.

Methods and materials

Data from these participants have been previously presented, and additional information regarding materials and methods are available in those publications8,12. The study protocols were approved by the UCLA Institutional Review Board, and all participants gave written, informed consent before any study procedures were carried out. Participants were recruited from the greater Los Angeles community through flyers and Internet advertisements. Eighteen healthy controls and 18 women with PMDD completed the study. Complete data were not available for one participant in the PMDD group, who asked to leave the scanner before completing the session due to discomfort; therefore, data from 17 PMDD participants are included in the imaging results presented below. All participants completed two experimental sessions: one during the follicular phase, 5–12 days after the onset of menstruation, and one during the luteal phase, 10–14 days after ovulation. Ovulation was estimated using at-home urinary luteinizing hormone detection kits (Clearblue® Digital Ovulation kit; SPD Swiss Precision Diagnostics GmbH, Geneva). The order of the first testing session (follicular or luteal) was determined randomly, leading to 61% of PMDD participants and 50% of controls beginning the study in the follicular phase, and the rest in the luteal phase.

Inclusion and exclusion criteria

The participants were required to be between the ages of 18 and 44 inclusive, non-smokers, fluent in English, right-handed, willing to use non-hormonal contraception or abstinence for the duration of the study, and to have regular menstrual cycles every 24–32 days. Participants were excluded if they reported any history of psychiatric diagnoses other than unipolar depression, or if unipolar depression had occurred within the past two years. Mental health history was assessed by a trained clinician using the Structured Clinical Interview for DSM-IV Axis I Disorders, Patient Edition25. Participants were also excluded if they endorsed a history of central nervous system, cardiovascular, hepatic, renal, endocrine, or autoimmune disease during the health history taken by a nurse practitioner. MRI contraindications, including non-removable metal or greater than minimal claustrophobia, were also exclusionary.

A PMDD diagnosis (for inclusion in the PMDD group) was assigned on the basis of scores on the Daily Record of Severity of Problems (DRSP) over the course of two complete menstrual cycles. Diagnosis and subsequent
inclusion decisions were made before participants were scheduled for additional data collection. Complete DRSPs were filled out daily online throughout the duration of the study,26; graphical depictions of overall symptom profiles for each participant are available as Supplementary Figs. S1–S18. Participants were required to endorse low (<3) scores on all DRSP items during the follicular phase, defined here as 7–12 days after the onset of menstruation, and high scores on DRSP items during the premenstrual phase, defined here as 6 days before menstruation. High scores were operationalized as:

(1) scores ≥3 for at least 4 days, and ≥4 for at least 2 days on mood symptoms (DRSP items 1 through 4), and

(2) scores ≥3 for at least 2 days, and ≥4 for at least 2 days on at least 5 of symptoms (DRSP items 1 through 11)

(3) scores ≥3 for at least 2 days, and ≥4 for at least 2 days on items measuring severity of impairment (DRSP items 12 through 14)

Participants in the healthy control group were required to endorse symptom scores <3 during the premenstrual phase, and <3 during the follicular phase. Absolute values on each day were evaluated rather than averaging together scores across each phase or cycle.

At each testing session, blood samples (5 mL) were collected by venipuncture for assay of progesterone levels by electrochemiluminescence (Roche Elecsys Immunoassay system, F. Hoffman-La Roche, Basel, Switzerland).

Detailed demographic information is presented in Supplementary Table S1. The ethnic composition of each group was somewhat different, with the PMDD group predominantly (78%) non-Hispanic white, and no group predominating among controls (Supplementary Table 1). Statistical comparisons were not performed between groups because of the small cell sizes involved. Group differences in self-reported years of education, self-reported annual income, and IQ estimated by performance on the Shipley-2 Vocabulary Test were tested in a two-way t-test in JMP(R) Pro 11.0.0 (SAS Institute Inc., Cary, NC, USA).

Study design specifications

The sample size (N = 18/group) was determined prospectively, and was similar in size to existing published studies evaluating brain activity in women with PMDD,9,22. Participants completed two fMRI resting-state scanning sessions on each test day separated by an emotion-regulation task. Details of the task and results have been reported (Petersen et al., 2017). Briefly, the task involved using a cognitive distancing strategy to reappraise negative stimuli. Participants were instructed that if they saw the instruction “Far”, they were to imagine that they were far away from the scene they were about to view, as if they were distant observers or reporters reporting on the details and facts of the scene. If they saw the instruction “Close,” they were to imagine that they were actually immersed in the scene they were about to view. Following the instruction, they were presented with either a negatively or neutrally valenced image, selected mostly from the International Affective Picture System. After each such trial (instruction + image), participants rated how bad they felt on a scale from 1 to 4, with 1 indicating “not at all bad” and 4 indicating “very bad” using a hand-held button box in the scanner.

The resting-state scans (152 T2*-weighted echoplanar images; repetition time = 2 s; echo time = 30 ms; slice thickness = 4 mm; flip angle = 90°; matrix: 64 × 64; field of view = 192 mm) were acquired over 5 min with a 3-T Siemens AG Trio MRI scanner (Erlangen, Germany). Participants viewed a black screen and were instructed to, “relax, try to stay as still as possible, and keep your eyes open.” A T1-weighted magnetization-prepared rapid-acquisition gradient echo scan (MPRAGE) and a T2-weighted matched-bandwidth anatomical scan were acquired to improve registration to standard space.

Preprocessing and analysis

All MRI preprocessing and analyses were performed in the FMRIB Software Library (FSL) version 5.0.9 (www.fmrib.ox.ac.uk/fsl). Non-brain matter was removed with FSL’s brain extraction tool, low-frequency trends were removed with high-pass filtering (100 s threshold), and functional images were registered to standard space through three steps1: FMRIB’s Linear Image Registration Tool (FLIRT) was used to register the resting-state functional scan to the matched-bandwidth image through affine transformation2; the same procedure was used to register the resting-state functional image to the MPRAGE image; and finally3 this image was transformed to the MNI152 template through FMRIB’s Nonlinear Image Registration Tool (FNIRT) with 12 degrees of freedom and a warp resolution of 10 mm.

Motion cleaning was performed with regressors entered as confound variables into a linear regression model. First, twenty-four motion regressors were used as per27. These included the three translational motion parameters along the X, Y, and Z axes and three in the rotational dimension (“pitch”, “roll” and “yaw”) (6 total); then the temporal difference (difference of the current and previous time-point) (6 total), and the quadratics of these 12. Next, framewise displacement (FD) was entered as “spike” regressors in the model28,29. To account for additional variance due to noise30, the global signal was included in the regression model31. The residuals produced by the regression model were then scaled and normalized at each voxel with the equation: [(residuals−mean)/standard deviation] + 100.
To examine intrinsic ECN connectivity, preprocessed images were entered as multi-session temporally concatenated data into FSL’s Multivariate Exploratory Linear Optimized Decomposition into Independent Components (MELODIC) tool with dimensionality estimation set to 20 components. The components of interest for further analysis (left and right ECN) were identified through visual inspection (similarity to intrinsic connectivity networks 15 and 18 in Laird et al.32) Individual subject-level maps of these components were determined using FSL’s dual regression tool. Effects of group (PMDD vs. control) and phase (follicular vs. luteal) on ECN connectivity were tested using Randomise, FSL’s nonparametric permutation-based statistical modeling tool33, using Threshold-Free Cluster Enhancement (TFCE), a cluster-extent threshold of \(Z = 2.3\) and 10,000 permutations in a mixed-model design with group entered as a between-subjects factor and phase as a within-subjects factor. The group-by-phase interaction was evaluated by testing for an effect of group on difference maps of the follicular-luteal differences. Group and phase effects on resting-state functional connectivity were tested on the scan that preceded the task (rest 1), and then the same analysis was repeated on the scan that followed the task (rest 2). Using the same model, exploratory analyses were also performed on a DMN component (see https://neurovault.org/collections/4829/ for spatial map of this component).

Seed-based connectivity of the left and right amygdala was derived using the General Linear Model as implemented in FSL’s FEAT. First, amygdala regions of interest defined by the Harvard-Oxford Probabilistic Atlas were transformed into each participant’s native space. Time series data within the left and right amygdala were extracted from the motion-cleaned data (described above) and averaged across voxels, resulting in a single-time series variable for each of the two (left and right) amygdala regions. These time-series were separately entered into two GLM models, one for the right and one for the left amygdala, resulting in a “connectivity map” (GLM parameter estimate maps) for each of the two amygdala regions for each participant. These maps were analyzed for group and phase effects using Randomise with TFCE, 10,000 permutations, and a cluster-extent threshold of \(Z = 2.3\). The main effects of group and phase and their interaction were evaluated in the same manner as the ECN data (described above).

To test whether differences in connectivity were related to PMDD symptoms (i.e., average of total DRSP score during the symptomatic phase), the averaged parameter estimates from the dual-regression model were extracted from clusters where group differences in resting-state functional connectivity were detected. These values were entered into a linear mixed model as a fixed effect, and participant as a random effect. Because PMDD symptoms were nearly zero in controls, the analysis was performed only on data from women with PMDD. PMDD symptoms as reported on the DRSP were entered as the dependent variable in the model.

A similar model was used to test whether functional connectivity was related to successful emotion regulation. Connectivity strength between the left ECN and the cluster that differed between groups was extracted and entered into a linear mixed model as a fixed effect, with participant as a random effect. Here, group (PMDD vs. control) was entered as a covariate. To evaluate the effect of connectivity strength on emotion regulation, mean participant ratings for all trials in which they viewed negatively valenced stimuli and were instructed to use a distancing strategy (“negative, far” trials) were entered into the model as the dependent variable.

**Results**

**Demographics**

The groups did not differ significantly in age, years of education, income, or IQ as estimated by the Shipley-2 vocabulary test34, all \(ps > 0.1\) (see Supplementary Table S1).

**Progestrone**

Progestrone levels differed significantly from the follicular to luteal testing days \([F(1,71) = 44.52, p < 0.0001]\) but not between the PMDD and control groups during either the follicular \([F(1,35) = 2.25, p = 0.14]\) or luteal \([F(1,35) = 0.38, p = 0.54]\) phase.

**Network identification**

The ECN was visually identified from the 20 components generated by MELODIC. A left-lateralized frontoparietal network was identified as the left ECN, and a right-lateralized frontoparietal network was identified as the right ECN (Supplementary Fig. S19; all images presented in radiological orientation).

**Group differences in resting-state functional connectivity**

Effects of group and phase were tested in a mixed-model design for the left ECN, right ECN, and default mode network. In both the phase comparison (follicular vs. luteal) and the group-by-phase interaction test, no voxels survived permutation testing at the 0.01 level, suggesting that there was no effect of phase and no group-by-phase interaction on resting-state functional connectivity within the left ECN. We employed an additional layer of correction for the 5 comparisons performed (left ECN, right ECN, DMN, right amygdala, left amygdala), further bringing the significance threshold for these familywise-error corrected clusters down to \(p = 0.01 (\alpha = 0.05/5)\). No effects of group, phase, or their interaction were found in the right ECN or DMN.
In the left ECN analysis only, a significant main effect of group emerged in a very small functional group of four non-contiguous voxels in the left middle temporal gyrus (peaks of two-voxel clusters at MNI x, y, z = −60, −54, −4 and −54, −56, 0), indicating significantly stronger connectivity between these middle temporal gyrus voxels and the rest of the left ECN in the PMDD group compared to healthy controls, irrespective of menstrual phase. The GLM parameter estimates in these voxels were extracted and the means for each group and phase were plotted (Fig. 1). No effects of group or phase were found in right ECN. Notably, this cluster was not significant after correcting for the number of tests performed ($p = 0.036$).

When the same analysis was repeated for the second rest scan (following the emotion regulation task), the same relationship was observed: significantly stronger connectivity between a middle temporal gyrus cluster and the rest of the left ECN in the PMDD group compared to controls, irrespective of menstrual phase. The cluster showing a significant group difference included 112 voxels (peak voxel at $−58, −60, −10$). Again, there was no effect of phase or group-by-phase interaction on resting-state functional connectivity in the left or right ECN (Fig. 2). This cluster survived correction for the total number of tests performed ($p = 0.002$).

A seed-based connectivity analysis comparing connectivity of the amygdala in the follicular vs. luteal phases showed significantly stronger connectivity between the left amygdala and a number of parietal and midline clusters focused around the posterior cingulate cortex, mid-cingulate cortex, and right angular gyrus in the follicular phase. A mixed-model analysis revealed a significant group-by-phase interaction, with a smaller follicular-luteal change in connectivity in women with PMDD compared to healthy controls (Fig. 3; Supplementary Table S2). This follicular-luteal connectivity difference was only observed in the first fMRI session of the day, before the emotion regulation task was performed. No effects of group or phase were found during the second, post-task resting-state session.

Similarly, a seed-based connectivity analysis evaluating right amygdala connectivity also indicated stronger connectivity between the right amygdala and clusters in the middle temporal gyrus in the follicular phase than the luteal phase that survived correction for the total number of comparisons (Fig. 4; Supplementary Table S3). No effect of group or group-by-phase interaction was detected, and this menstrual phase effect was observed only in the first resting-state session, which took place before the emotion regulation task.

**Linking resting-state connectivity to behavior**

To test whether connectivity of the MTG cluster to IECN was related to PMDD symptom severity, the mean parameter estimates of the cluster in both rest 1 and rest 2 were extracted and correlated with PMDD symptoms as reported on the DRSP. No significant relationship was observed ($p = 0.68$).

The relationship between this connectivity measurement and successful emotion regulation through cognitive reappraisal was also tested using this model. The mean parameter estimates of this cluster during rest 1 were unrelated to ratings on “negative, far” trials ($p = 0.18$) but were significantly related during rest 2 ($p = 0.0054$). This relationship (Fig. 5) was significant in the PMDD group ($p = 0.04$) but not the control group ($p = 0.28$).
Left and right amygdala connectivity was not related to behavior on the emotion regulation task.

**Counterbalancing**

To verify that order effects did not influence the results, a linear mixed model testing the main effect of session (timepoint 1 versus timepoint 2) was run; no effects of session on any resting-state functional connectivity parameter were observed.

**Effect size maps**

Effect size maps illustrating the outcome of all analyses described above are available at https://neurovault.org/collections/4829/.

**Discussion**

Researchers have speculated that PMDD symptoms may be explained by “impaired ovarian hormones-mediated sensitivity of emotional and cognitive brain networks” \(^{35}\), yet the integrity and dynamics of such networks in women with PMDD have been only minimally explored. Neural models of affective disorders have started to converge on findings that network-level dynamics may underlie affective symptoms, with top-down control from prefrontal cortical regions failing to effectively regulate bottom-up affective processes from limbic regions (for review, see Disner et al. \(^{36}\)) to produce these symptoms. Here, we show that intrinsic network connectivity in women with PMDD is broadly consistent with this model of prefrontal cortical dysregulation insofar as women who meet DSM-5 criteria for PMDD have significantly stronger connectivity between a region of the left middle temporal gyrus and the left ECN relative to healthy controls when challenged by an emotional task. This effect appears to be stable across menstrual cycle phases.

The cluster that shows a group difference in connectivity appears to be related to regulation of negative emotion in women with PMDD only, in whom the mean parameter estimates of this cluster correlated with ratings of negative emotion obtained during an emotion regulation task, implying that stronger connectivity reduced negative emotion produced by the task.
controls), the cluster that differed between women with PMDD and controls increased in size after participants performed an emotionally demanding task, and the connectivity of this cluster differed considerably between women with PMDD and controls. These findings are also broadly consistent with previous evidence showing differences in intrinsic network connectivity between healthy controls and women with PMS, a milder syndrome similar to PMDD. Similar to our finding of stronger connectivity between the left middle temporal gyrus and left ECN compared to controls, Liu et al. reported stronger connectivity between the left middle temporal gyrus and default mode network in women with PMS compared to controls. Intriguingly, in both studies, the effect was independent of menstrual cycle phase, despite the dramatic difference in symptom presentation between the follicular and late luteal phases.

The seed-based connectivity analysis did not indicate a significant effect of PMDD on amygdala connectivity. Instead, it revealed a menstrual phase-related shift in amygdala connectivity as the cycle progressed from the follicular to late luteal phases in both healthy controls and women with PMDD. Left amygdala connectivity to the posterior cingulate cortex, mid-cingulate area, right angular gyrus, and right superior parietal cortex was significantly weaker in both groups (controls and PMDD) during the late luteal phase compared to the follicular phase. Similarly, right amygdala connectivity to the cerebellum and left middle temporal gyrus were significantly weaker during the late luteal phase compared to the follicular phase. Because these connectivity reductions were observed in healthy controls as well as women with PMDD, they appear to reflect normal cyclicity in brain connectivity. The functional significance of this change is not evident from these data, and warrants further investigation.

Some limitations affected this study. Resting-state connectivity datasets are information-rich and provide the opportunity to test for effects within many networks, as well as between any number of seed regions. The small sample size available here provided limited statistical power, which constrained the number of hypotheses that could be tested; therefore, we limited hypothesis testing to the highest priority networks and seeds. We have made these data available for other investigators to search for trends or test their own hypotheses (see https://neurovault.org/collections/4829/). It also bears considering that the small sample may obscure effects produced by subtypes of PMDD, as the group was too small to further subdivide. Since these data were collected, temporal subtypes reflecting unique timescourses of symptom presentation (i.e., symptoms constrained to a relatively brief premenstrual window; symptoms persisting through the entire luteal window; or symptoms that persist into the onset of the next cycle) have been described. These and other subtypes of PMDD patients may experience different patterns of intrinsic network connectivity that could not be addressed in this study and represent a target for future investigations.

The data presented here point to a fundamental neurobiological difference between women with PMDD and healthy controls. Identifying the network dynamics that manifest differently in women with PMDD moves the field toward identifying a biomarker for this under-researched condition. As experts in the field have called for novel therapeutic approaches, this report identifies targets for brain-based therapies, such as neurostimulation or neurofeedback, which have shown promise for treatment of other affective disorders.

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Conflict of interest
The authors declare that they have no conflict of interest.

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References
1. Freeman, E. W. Premenstrual syndrome and premenstrual dysphoric disorder: definitions and diagnosis. Psychoneuroendocrinology 28, 25–37 (2003).
2. Rapkin, A. A review of treatment of premenstrual syndrome and premenstrual dysphoric disorder. Psychoneuroendocrinology 28, 39–53 (2003).
3. Hartlage, S. A., Freels, S., Gotman, N. & Yonkers, K. Criteria for premenstrual dysphoric disorder: secondary analyses of relevant data sets. Arch. Gen. Psychiatry 69, 305–5 (2012).
4. Association AP. Diagnostic and Statistical Manual of Mental Disorders: Diagnostic and Statistical Manual of Mental Disorders 5th edn (American Psychiatric Association, Arlington, VA, 2013).
5. Yamada, K. & Kamagata, E. J. Reduction of quality-adjusted life years (QALYs) in patients with premenstrual dysphoric disorder (PMDD). Qual. Life Res. 26, 3069–73 (2017).
6. Wittchen, H. U., Becker, E., Lieb, R. & Krause, P. Prevalence, incidence and stability of premenstrual dysphoric disorder in the community. Psychol. Med. 32, 119–32 (2002).

7. Protopopescu, L. et al. Toward a functional neuroanatomy of premenstrual dysphoric disorder. J. Affect. Disord. 108, 87–94 (2008).

8. Petersen, N. et al. Brain activation during emotion regulation in women with premenstrual dysphoric disorder. Psychol. Med. 48, 1795–802 (2018).

9. Baller, E. B. et al. Abnormalities of dorsolateral prefrontal function in women with premenstrual dysphoric disorder: a multimodal neuroimaging study. Am. J. Psychiatry 170, 305–14 (2013).

10. Buhle, J. T. et al. Cognitive reappraisal of emotion: a meta-analysis of human neuroimaging studies. Cereb. Cortex 24, 2981–2990 (2014).

11. Kohn, N. et al. Neural network of cognitive emotion regulation — an ALE meta-analysis and MACM analysis. Neuron Image 87, 345–355 (2014).

12. Petersen, N. et al. Emotion regulation in women with premenstrual dysphoric disorder. Arch. Women’s Ment. Health 19, 891–8 (2016).

13. Greicius, M. Resting-state functional connectivity in neuropsychiatric disorders. Curr. Opin. Neurol. 21, 424–30 (2008).

14. Greicius, M. D. et al. Resting-state functional connectivity in major depression: abnormally increased contributions from subgenual cingulate cortex and thalamus. Biol. Psychiatry 62, 429–37 (2007).

15. Connolly, C. G. et al. Resting-state functional connectivity of subgenual anterior cingulate cortex in depressed adolescents. Biol. Psychiatry 74, 896–907 (2013).

16. Kaiser, R. H. et al. Dynamic resting - state functional connectivity in major depression. Neuropsychopharmacology 41, 1822–30 (2016).

17. Rapkin, A. J. & Winer, S. A. Premenstrual syndrome and premenstrual dysphoric disorder: quality of life and burden of illness. Expert Rev. Pharmacoecon. Outcomes Res. 9, 157–70 (2009).

18. Ding, D. et al. Larger volume and different functional connectivity of the amygdala in women with premenstrual syndrome. Eur. Radiol. 28, 1900–8 (2018).

19. Liu, Q., Li, R., Zhou, R., Li, J. & Gu, Q. Abnormal resting-state connectivity at functional MRI in women with premenstrual syndrome. PLoS ONE 10, e0136029 (2015).

20. Petersen, N. & Cahill, L. Amygdala reactivity to negative stimuli is influenced by oral contraceptive use. Soc. Cogn. Affect. Neurosci. 10, 1266–72 (2015).

21. van Wingen, G. A. et al. Progesterone selectively increases amygdala reactivity in women. Mol. Psychiatry 13, 325–33 (2008).

22. Gingnell, M., Morell, A., Bannbers, E., Wikstrom, J. & Sundstrom-Poromaa, I. Menstrual cycle effects on amygdala reactivity to emotional stimulation in premenstrual dysphoric disorder. Hormones Behav. 62, 400–6 (2012).

23. Sheline, Y. I. et al. The default mode network and self-referential processes in depression. Proc. Natl Acad. Sci. USA 106, 1942 (2009).

24. Zhu, X. et al. Evidence of a dissociation pattern in resting-state default mode network connectivity in first-episode, treatment-naive major depression patients. Biol. Psychiatry 71, 617–7 (2012).

25. First M. B., Spitzer R. L., Gibbon M., Williams J. B. W. Structured Clinical Interview for DSM-IV-TR Axis I Disorders, Research Version, Patient Edition (New York State Psychiatric Institute, 2002).

26. Endicott, J., Nee, J. & Harrison, W. Daily Record of Severity of Problems (DRSP): reliability and validity. Arch. Women’s Ment. Health 9, 41–9 (2006).

27. Friston, K. J., Williams, S., Howard, R., Frackowiak, R. S. & Turner, R. Movement-related effects in fMRI time-series. Magn. Reson. Med. 35, 346–355 (1996).

28. Power, J. D., Barnes, K. A., Snyder, A. Z., Schlaggar, B. L. & Petersen, S. E. Spurious but systematic correlations in functional connectivity MRI networks arise from subject motion. Neuron Image 59, 2142–54 (2012).

29. Power, J. D., Barnes, K. A., Snyder, A. Z., Schlaggar, B. L. & Petersen, S. E. Steps toward optimizing motion artifact removal in functional connectivity MRI: a reply to Carp. Neuron Image 76, 439–41 (2013).

30. Satterthwaite, T. D. et al. An improved framework for confound regression and filtering for control of motion artifact in the preprocessing of resting-state functional connectivity data. Neuron Image 64, 240–56 (2013).

31. Murphy, K. & Fox, M. D. Towards a consensus regarding global signal regression for resting state functional connectivity MRI. Neuro Image 154, 169–73 (2017).

32. Laird, A. R. et al. Group-based trajectory modeling to identify individual differences in symptom change. Psychol. Med. 23, 1–9 (2019).

33. Wittchen, H. U., Becker, E., Lieb, R. & Krause, P. Prevalence, incidence and stability of premenstrual dysphoric disorder in the community. Psychol. Med. 32, 119–32 (2002).

34. Kohn, N. et al. Neural network of cognitive emotion regulation — an ALE meta-analysis and MACM analysis. Neuron Image 87, 345–355 (2014).

35. Petersen, N. et al. Emotion regulation in women with premenstrual dysphoric disorder. Arch. Women’s Ment. Health 19, 891–8 (2016).

36. Greicius, M. Resting-state functional connectivity in neuropsychiatric disorders. Curr. Opin. Neurol. 21, 424–30 (2008).

37. Greicius, M. D. et al. Resting-state functional connectivity in major depression: abnormally increased contributions from subgenual cingulate cortex and thalamus. Biol. Psychiatry 62, 429–37 (2007).

38. Connolly, C. G. et al. Resting-state functional connectivity of subgenual anterior cingulate cortex in depressed adolescents. Biol. Psychiatry 74, 896–907 (2013).

39. Kaiser, R. H. et al. Dynamic resting - state functional connectivity in major depression. Neuropsychopharmacology 41, 1822–30 (2016).

40. Rapkin, A. J. & Winer, S. A. Premenstrual syndrome and premenstrual dysphoric disorder: quality of life and burden of illness. Expert Rev. Pharmacoecon. Outcomes Res. 9, 157–70 (2009).

41. Ding, D. et al. Larger volume and different functional connectivity of the amygdala in women with premenstrual syndrome. Eur. Radiol. 28, 1900–8 (2018).

42. Liu, Q., Li, R., Zhou, R., Li, J. & Gu, Q. Abnormal resting-state connectivity at functional MRI in women with premenstrual syndrome. PLoS ONE 10, e0136029 (2015).

43. Petersen, N. & Cahill, L. Amygdala reactivity to negative stimuli is influenced by oral contraceptive use. Soc. Cogn. Affect. Neurosci. 10, 1266–72 (2015).

44. van Wingen, G. A. et al. Progesterone selectively increases amygdala reactivity in women. Mol. Psychiatry 13, 325–33 (2008).

45. Gingnell, M., Morell, A., Bannbers, E., Wikstrom, J. & Sundstrom-Poromaa, I. Menstrual cycle effects on amygdala reactivity to emotional stimulation in premenstrual dysphoric disorder. Hormones Behav. 62, 400–6 (2012).