Altered regional homogeneity of spontaneous brain activity in idiopathic trigeminal neuralgia

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Abstract: The pathophysiology of idiopathic trigeminal neuralgia (ITN) has conventionally been thought to be induced by neurovascular compression theory. Recent structural brain imaging evidence has suggested an additional central component for ITN pathophysiology. However, far less attention has been given to investigations of the basis of abnormal resting-state brain activity in these patients. The objective of this study was to investigate local brain activity in patients with ITN and its correlation with clinical variables of pain. Resting-state functional magnetic resonance imaging data from 17 patients with ITN and 19 age- and sex-matched healthy controls were analyzed using regional homogeneity (ReHo) analysis, which is a data-driven approach used to measure the regional synchronization of spontaneous brain activity. Patients with ITN had decreased ReHo in the left amygdala, right parahippocampal gyrus, and left cerebellum and increased ReHo in the right inferior temporal gyrus, right thalamus, right inferior parietal lobule, and left postcentral gyrus (corrected). Furthermore, the increase in ReHo in the left precentral gyrus was positively correlated with visual analog scale ($r=0.54; P=0.002$). Our study found abnormal functional homogeneity of intrinsic brain activity in several regions in ITN, suggesting the maladaptivity of the process of daily pain attacks and a central role for the pathophysiology of ITN.

Keywords: trigeminal neuralgia, resting fMRI, brain, chronic pain, local connectivity

Introduction

Idiopathic trigeminal neuralgia (ITN) is characterized by paroxysmal, electric, lancinating pains, and a variety of sensory experiences. The annual incidence is 4–5 in 100,000, and it is estimated to affect one in 15,000–20,000 people worldwide. With the expansion of the aging population, the proportion of elderly patients with ITN has gradually increased. Despite its prevalence and potential medical and social burdens, the pathogenesis underlying these pains remains poorly understood.

The prevailing pathophysiology of trigeminal neuralgia emphasizes its peripheral nociception and central component mechanisms. Peripheral nociception describes the hyperexcitability of the nerve-vessel conflict, whereas the central component relates to central facilitation and results in central hyperexcitability of the trigeminal system that is initiated or sustained by peripheral mechanisms. An ambiguous relationship between these two mechanisms may exist, given that ITN patients endure typical paroxysmal neuralgic perception and concomitant dull background pain between the two attacks that cause the ITN patient’s brain to frequently reconfigure its intrinsic activity and to adapt its functional architecture following daily attacks. It is therefore reasonable to expect these patients to have an altered brain intrinsic activity.

Over the past two decades, neuroimaging research has improved our understanding of how chronic pain (eg, migraines, arthritis, fibromyalgia, chronic back pain, and visceralgia) can affect brain structure and functions. Of particular interest,
magnetic resonance imaging (MRI)-based brain imaging offers a suitable avenue for measuring the functional and structural alterations that underlie pain perception and chronification.10–12 Recent studies have found that ITN is associated with structural changes in pain, attention, and motor processing areas. Evidence from structural MRI studies on ITN patients have found that ITN is associated with cortical and subcortical gray matter atrophy in the somatosensory, orbitofrontal, insular, anterior cingulate, and dorsolateral prefrontal cortices in addition to the thalamus and cerebellum.13–16 In brain diffusion tensor imaging studies, lower fractional anisotropy and increased radial diffusivity, mean diffusivity, and axial diffusivity have been recorded in brain white matter including the corpus callosum, cingulum, posterior corona radiata, and superior longitudinal fasciculus of ITN patients.17 Furthermore, abnormal thalamic metabolism of N-acetylaspartate/creatine has been reported in ITN patients.17,18 Despite an increasing amount of research regarding the effects of ITN on brain structure and metabolism, little is known about the effects of ITN on spontaneous brain activity.

Regional homogeneity (ReHo) analysis is a data-driven method of resting-state functional MRI (rsfMRI) that can measure the similarity across a time series of a given voxel with its nearest neighbors, without a priori knowledge of the experimental design.19 ReHo analysis is primarily focused on an activity pattern within a specific brain region and has been widely used in various clinical populations and phenotypic variables to reveal the regional synchronization of spontaneous brain activity.20,21

In this study, we hypothesized that ITN patients would show altered ReHo across central pain processing areas. Our aim was 1) to detect ReHo changes in ITN patients to localize specific brain regions that may be related to the development and persistence of this debilitating facial pain and 2) to correlate clinical variables of ITN with the ReHo of each patient within significant group differences.

Materials and methods
Informed consent was obtained from all participants. The study was conducted in accordance with the Declaration of Helsinki. The study protocol was approved by the Institutional Review Board of the Second Hospital of Jiaxing City.

Participants
Seventeen ITN patients (seven males/ten females, age =63.41±7.25 years old) were recruited prospectively from the Department of Neurology at the Second Hospital of Jiaxing City between February 2014 and March 2015. The inclusive criteria for the ITN patients were as follows: 1) unilateral pain in the distribution of one or more branches (the ophthalmic [V1], the maxillary [V2], and the mandibular [V3]) of the trigeminal nerve; 2) stereotyped attacks, and characteristic (intense, sharp, superficial, or stabbing paroxysmal) pains precipitated from trigger areas or by trigger factors for each patient; 3) no additional neurological or sensory deficits in all patients; and 4) no previous surgical or other invasive procedures for ITN.

For each patient, demographic and clinical details were obtained via retrospective chart reviews, and the diagnosis of ITN was confirmed by experienced neurologists (Yanping Wang and Xiaoling Zhang) according to the International Classification of Headache Disorders–2nd edition (ICHD-II).22 The exclusion criteria were 1) patients with neural-associated diseases or chronic pain other than ITN, hypertension, diabetes, or a history of brain surgery and 2) patients unable to undergo MRI scanning because of metal implantation or those with a history of claustrophobia or other psychological disorders. In addition, we also recruited 19 age- and sex-matched healthy subjects, with no family history of neuralgia, no history of substance abuse, chronic pain, systemic, psychiatric, or neurological disorders as controls. All of the participants were right handed according to their self-report. Table 1 lists the demographic information of the ITN patients and the controls.

Questionnaires
Before MRI scanning, the following questionnaires were given to the ITN patients who were asked to respond based on his or her current pain. The duration of pains (year), pain days per month (times), as well as visual analog scales (VAS) for pain was determined by asking the patient to mark the pain severity on a 10 cm line. Additionally, all subjects completed the depression scores Beck’s Depression Inventory-II (BDI-II),23 the Hamilton Depression (HAMD)/

Table 1 Subject characteristics

|                | ITN            | Controls       | P-value |
|----------------|----------------|----------------|---------|
| Age (years old) | 63.41±7.25     | 62.53±7.41     | 0.72    |
| Sex (M/F)      | 7/10           | 9/10           | >0.99   |
| Handedness     | 17R            | 19R            | >0.99   |
| MMSE           | 28.9±1.7       | 28.1±1.3       | 0.105   |
| BDI-II         | 2.4±1.8        | 2.2±1.4        | 0.67    |
| HAMD           | 2.9±2.1        | 1.9±1.1        | 0.147   |
| HAMA           | 3.5±2.9        | 1.7±0.9        | 0.035*  |
| Duration       | 6.98±5.64 y, mid 5.00 y | NA            | NA      |

Note: *P<0.05.

Abbreviations: ITN, idiopathic trigeminal neuralgia; M/F, male/female; R, right; y, year; MMSE, Mini-Mental State Examination; BDI-II, Beck’s Depression Inventory-II; HAMA, Hamilton Anxiety Rating Scale; HAMD, Hamilton Depression Rating Scale; NA, not applicable.
Anxiety (HAMA) Rating Scale and the Mini-Mental State Examination. Details are presented in Table 1.

MRI data acquisition

MRI data were acquired on a 1.5 T GE Signa Excite scanner (GE Medical Systems, Milwaukee, WI, USA) using a standard GE whole-head coil. Each subject laid supine, with the head in a neutral position and fixed comfortably by a belt and foam pads during the test. High-resolution T1-weighted anatomical datasets were obtained parallel to the AC–PC plane and covering the whole brain using three-dimensional spoiled-gradient recall (SPGR) sequence (repetition time [TR] / echo time [TE] / inversion time [TI]) = 10.68 ms / 4.87 ms / 380 ms, flip angle = 15°, field of view = 256 mm, 140 axial slices, voxel size 1×1×1 mm³). Resting-state functional datasets were recorded using a T2*-weighted gradient echo spiral pulse sequence (30 axial slices, thickness/gap = 5.0/1 mm, in-plane resolution = 64×64, TR = 2,000 ms, TE = 40 ms, flip angle = 90°, field of view (FOV) = 240 mm × 240 mm). The scan time for the rsfMRI session for each participant was approximately 7 minutes.

Data processing

rsfMRI data were preprocessed and analyzed using Data Processing Assistant for Resting-State fmRI (DPARSF, http://www.restfmri.net), which is based on Statistical Parametric Mapping (SPM8, http://www.fil.ion.ucl.ac.uk/spm) and Matlab. For the rsfMRI data for each subject, the first ten volumes were discarded to avoid the possible effects of scanner instability and the adaptation of subjects to the environment, and the remaining 190 volumes were corrected for timing offsets. The resultant images were then realigned and coregistered to the T1 structural image. The T1 structural image was spatially normalized to the Montreal Neurological Institute space using diffeomorphic anatomical registration through exponentiated lie algebra, a more sophisticated registration algorithm, which can achieve a more accurate inter-subject registration, realignment of small deformations, and better spatial normalization.25 The resulting parameter was used to normalize the functional images with voxel size 3×3×3 mm³. Temporal filter (0.01–0.08 Hz) was conducted to lessen low-frequency drifts and high-frequency physiological noise. Nuisance regression was performed using white matter, cerebrospinal fluid, and the six head motion parameters as covariates. No participant had head motion with > 2.0 mm maximum displacement or 2.0° of any angular motion.

The ReHo was calculated as previously described.26 ReHo evaluates the degree of local synchronization among fMRI time courses. ReHo is computed as Kendall’s coefficient of concordance (KCC) value of the ranked time series of a given voxel to its nearest neighbors in a voxel-wise way:

\[
W = \frac{1}{12} \sum_{i=1}^{n} R_i^2 - n \left( \frac{\sum_{i=1}^{n} R_i^2}{n^2} \right)
\]

where W is the KCC among time series of given voxels, the range is 0–1; Rᵢ is the sum rank of the iᵗʰ time point, K is the number of time series within a measured cluster (here K=27, one center voxel plus the number of its neighbors), and n is the number of ranks (here, n=190 time points). The individual ReHo map was obtained for each subject by calculating the KCC value within its whole brain voxels. These intracranial voxels were extracted to make a brain mask. Each individual ReHo map was converted to a z-score map by subtracting the global mean ReHo and dividing the group standard deviation within the brain mask. Finally, the z-score maps were smoothed with a 6 mm full-width at half-maximum Gaussian kernel.

Statistical analysis

Demographic and clinical data were analyzed with SPSS 16.0 (SPSS Inc., Chicago, IL, USA) and ReHo of rsfMRI data with SPM8. Voxel-wise ReHo differences between the ITN patient group (n=17) and the control group (n=19) were performed with independent two-sample t-test. The threshold was set at voxel-level P<0.01 and cluster-level P<0.05, Gaussian random field correction.

Results

Demographic and clinical data

Table 1 shows the demographic information of the ITN and control groups. Table 2 demonstrates the clinical features of the ITN patients. Between-group t-tests identified significant group differences in HAMA, with patients of the ITN group scoring higher in anxiety (HAMA, P<0.05). The two groups displayed no differences in age, sex, handedness, Mini-Mental State Examination, BDI, or HDMA.

Voxel-wise comparison of ReHo

Table 3 and Figure 1 show the group differences in ReHo between the ITN group and the control group (voxel-level P<0.01, cluster-level P<0.05, Gaussian random field correction). Compared with the healthy controls, ITN patients showed decreased ReHo in the left amygdala, right parahippocampal gyrus, and left cerebellum. Increased ReHo was observed in the right inferior temporal gyrus, right...
thalamus, right inferior parietal lobule, the left pre- and postcentral gyrus (Table 3 and Figure 1). In order to investigate the impact of anxiety-related effects on the ReHo results, we further regressed out the HAMA score of each region showing a significant between-group ReHo difference; we found that the between-group differences in ReHo were not substantially affected by the HAMA regression.

Correlations between ReHo and clinical variables

Regional correlation analyses were conducted between the clinical variables of ITN patients and the cluster mean z score of each patient within the mask of significant group differences. The increase in ReHo in the left precentral gyrus was positively correlated with VAS score ($r=0.54; P=0.002$, Figure 1). No other significant correlations were identified between the altered ReHo and the other clinical scores assessed.

Discussion

In this study, we assessed the local synchronization of resting-state fMRI signals in patients with ITN, a unique neuropathic pain that is called the most painful condition in humans. Altered ReHo was detected in ITN patients, who showed decreased ReHo in the limbic system and cerebellum and increased ReHo in the right inferior temporal gyrus, right thalamus, right inferior parietal lobule, and the left postcentral gyrus. More importantly, increased ReHo of the left postcentral gyrus was associated with self-reported pain intensity (VAS). This study highlights the impairment to local synchronization of pain perception, pain modulation, and motor-related brain systems that is associated with central pain processing in patients with ITN.

Neural homogeneity is considered an important and robust feature of brain activity at microscale (ie, single neuron)\(^{27}\) and a reliable measure of connectomics at macroscale (ie, voxel/parcellation).\(^{27,28}\) Several studies have suggested that ReHo is linked to various phenotypic variables (eg, age, sex, intelligence, and personality), and differs between clinical populations. More recently, Jiang et al\(^{29}\) and Jiang and Zuo\(^{29}\) systematically reviewed the regional (short-distance) functional interactions of the organizational principles of human brain function, and proposed that ReHo may stand as a neurobiologically meaningful network centrality of the local features of the brain connectome.

Table 2 Subject characteristics

| Patient | Sex | Age, years | Duration, years | Pain location | Pain days per month | Pain intensity (VAS) | Vessel-nerve contact on MRI |
|---------|-----|------------|-----------------|---------------|--------------------|---------------------|--------------------------|
| Sub01   | M   | 46         | 20              | V2            | 15                 | 6                   | R                        |
| Sub02   | F   | 69         | 10              | V2            | 10                 | 7                   | L                        |
| Sub03   | F   | 56         | 3               | V1+V2         | 10                 | 5                   | R                        |
| Sub04   | M   | 58         | 6               | V2+V3         | 8                  | 8                   | L                        |
| Sub05   | M   | 60         | 1               | V2            | 6                  | 5                   | R                        |
| Sub06   | F   | 70         | 3               | V2            | 7                  | 7                   | R                        |
| Sub07   | F   | 68         | 10              | V3            | 5                  | 7                   | L                        |
| Sub08   | M   | 69         | 5               | V1+V2         | 8                  | 4                   | R                        |
| Sub09   | M   | 64         | 1               | V2            | 11                 | 8                   | L                        |
| Sub10   | F   | 68         | 2.5             | V3            | 15                 | 7                   | R                        |
| Sub11   | F   | 63         | 3               | V2+V3         | 9                  | 8                   | L                        |
| Sub12   | F   | 63         | 20              | V2+V3         | 8                  | 6                   | R                        |
| Sub13   | F   | 75         | 3               | V3            | 6                  | 6                   | R                        |
| Sub14   | M   | 58         | 10              | V2            | 4                  | 3                   | R                        |
| Sub15   | F   | 55         | 10              | V3            | 3                  | 7                   | R                        |
| Sub16   | F   | 65         | 3               | V2            | 6                  | 6                   | R                        |
| Sub17   | M   | 71         | 7               | V2            | 7                  | 4                   | R                        |

Note: V1, V2, V3 is the first, second and the third branch of the trigeminal nerve, respectively.

Abbreviations: M, male; F, female; VAS, visual analog scale; MRI, magnetic resonance imaging; L, left; R, right.

Table 3 Brain areas with ReHo differences between the ITN and control groups

| Brain regions             | MNI coordinates | BA | Number of Voxels | t-value |
|---------------------------|------------------|----|------------------|---------|
| Inferior temporal gyrus   | -51 -57 -15      | R  | 37               | 71      | 3.848 |
| Amygdala                  | -12 3 -30 28     | L  | 82               | -4.2591 |
| Cerebellum                | -9 -87 -21 18    | L  | 48               | -3.883 |
| Parahippocampal gyrus     | -12 -12 -21 34   | R  | 42               | -3.9466 |
| Thalamus                  | 6 -15 6         | R  | 43               | 4.0861  |
| Inferior parietal lobule  | 39 -45 42 40     | R  | 71               | 4.1893  |
| Postcentral gyrus         | -21 -51 69 7    | L  | 121              | 4.8598  |

Notes: Coordinates x, y, z (mm) are given in standard stereotactic MNI space. All regions listed are statistically significant at $P<0.05$, AlphaSim corrected.

Abbreviations: ReHo, regional homogeneity; ITN, idiopathic trigeminal neuralgia; MNI, Montreal Neurological Institute; BA, Brodmann area; L, left; R, right.
In our study, altered patterns of ReHo were detected in patients with ITN, including decreased ReHo in the left amygdala, right parahippocampal gyrus, and left cerebellum and increased ReHo in the right inferior temporal gyrus, right thalamus, right inferior parietal lobule, and left postcentral gyrus. This is consistent with several structural brain imaging studies that have reported gray matter changes in pain perception, pain modulation, and motor areas.\(^\text{14,16}\) For example, reduced gray matter size was documented in the somatosensory cortex, thalamus, parahippocampus, and temporal lobe in ITN patients.\(^\text{30}\) Another study\(^\text{14}\) reported greater gray matter volume in the thalamus, amygdala, periaqueductal gray matter, and basal ganglia. By contrast, ITN patients had thinner cortex in the anterior cingulate, the insula, and the orbitofrontal areas. Although inconsistency does exist in morphometric studies, the involved regions may also suggest a possible neuropathology in ITN.

The amygdala is a complex and deep brain structure involved in emotional processing. Fear conditioning and adjustment are the most noted emotional processes related to the amygdala,\(^\text{9,31}\) although its functional role has been extended to multiple dimensions including reward learning and motivation.\(^\text{9}\) Additionally, amygdala has been associated with a variety of psychiatric disorders, including anxiety\(^\text{32}\) and depression.\(^\text{33}\) Decreased ReHo in the left amygdala and parahippocampus is also reported in patients with experimentally induced lower back pain.\(^\text{34}\) Based on an investigation of the neurobiological significance of ReHo,\(^\text{21}\) decreased ReHo in the amygdala may indicate reduced functional integration of integrative information processing.

We also found that ITN patients had increased ReHo in the thalamus compared to controls. This result is consistent with a recent metabolic MRI study on ITN patients that showed a lower N-acetylaspartate/creatine ratio in the affected-side thalamus and a negative relationship with VAS and disease duration.\(^\text{17}\) In addition, previous studies of ITN have reported greater patient thalamic volume relative to controls.\(^\text{14}\) It is reasonable to assume that the increased local synchronization of blood oxygenation level dependent (BOLD) time series

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**Figure 1** Group comparison of ReHo and its correlation with clinical variables.

**Notes:** (A) Voxel-wise comparison of ReHo. Group differences in ReHo between the ITN group and the control group (voxel-level \(P<0.01\), cluster-level \(P<0.05\), GRF correction). Blue-green colors indicate the ITN patients who had reduced ReHo while red-yellow indicates the ITN patients who had increased ReHo compared to the controls. (B) Plots of the significant clusters. The mean and standard deviation of ReHo values in significant clusters. \(P<0.05\). (C) Correlations between ReHo and clinical variables. A positive correlation between the ReHo increase in the left sensorimotor areas and pain intensity in ITN patients, ie, the higher the pain intensity the ITN patient, the greater the ReHo (\(r=0.54, P=0.002\)).

**Abbreviations:** L, left; R, right; GRF, Gaussian random field; ITN, idiopathic trigeminal neuralgia; ReHo, regional homogeneity; VAS, visual analog scale.
in the thalamus reflects sustained nociceptive input, fitting with trigeminal neuropathy symptomology.

Both the parahippocampal gyrus and the lateral parietal cortex are regions of the default mode network (DMN), which is engaged in the internal processing including autobiographical memory, self-reference, and stimulus-independent thought. The DMN is generally active when it is not engaged with the external environment. The parahippocampus is involved in multiple behaviors, including stress and anxiety and the integration of complex information. Decreased parahippocampal ReHo in ITN patients supports its integrative role in internal processing and suggests that ITN involves decreased neural synchronization during internally directed or stimulus-independent mental states, which may be associated with an increased awareness of external nociceptive pain stimuli and anticipation. However, we also observed an increase in ReHo in the right lateral parietal cortex, and because they both constitute parts of DMN, the disparity between the parahippocampal decrease and the lateral parietal increase in ReHo could reflect maladaptive brain activity that is induced by pain. The lateral parietal cortex has been reported to participate in pain anticipation via meta-analysis, and it may represent a second level of the pain information processing circuit that supports the active, conscious, cognitive evaluation of pain sensation.

We also found increased ReHo in brain regions that generally encodes the discriminative properties of pain, including the pre- and postcentral gyri. The postcentral gyrus has been involved in the anticipation, intensity, discriminative, spatial and temporal summation aspects of pain processing. In patients with ITN, even simply non-painful movements can elicit attacks of pain, so restriction of facial movements would therefore ease the pain. The precentral gyrus (primary motor cortex) could therefore mirror sensory pain responses to repeated trigeminal neuralgia, motor inhibition of the maxilla and facial muscle tension. In our study, a positive relationship between pain intensity and ReHo in the left pre- and postcentral gyrus was found, suggesting a link between local synchronization of intrinsic brain activity and pain modulation.

It is also imperative to speculate whether altered ReHo occurs as an epiphenomenon that is secondary to daily stimulation or represents a functional predisposition to ITN in the light of peripheral vascular compression. However, our current study did not identify cingulate and insular involvement, which has been frequently reported in studies of pain anticipation and investigation of the interplay between intrinsic brain networks. Our data directly link ratings of self-reported pain to local homogeneity of spontaneous brain activity in the sensorimotor areas, which may reflect the maladaptive process of daily pain attacks, while other ReHo changes across distributed brain regions may at least indicate a central role for the pathophysiology of ITN. Broadly, our findings have implications for understanding how ITN affects local intrinsic brain activity.

Limitations
The current analysis was restricted to local brain activity (local synchronization of resting BOLD signals). Consequently, only changes in regional connectivity could be detected, and it was not possible to determine how ITN affected the interactions among brain networks systems, which are more widely used. Studies of functional connectivity and topology could be used in the future to explore these questions. Furthermore, the patients in our study were not divided into subgroups, as described in Obermann et al. As such, we could not clarify a causal relationship for the pathophysiology of ITN. Last but not least, the functional MRI data were acquired on a 1.5 T scanner, negative influences for low-magnetic field MR in detecting between-group findings may exist.

Conclusion
In summary, our results provide novel evidence for the altered local synchronization of low-frequency spontaneous activity across central pain processing brain regions in patients with ITN. Furthermore, a positive relationship between pain intensity and ReHo in the sensorimotor areas was identified. This may reflect the maladaptive process of daily pain attacks, while other ReHo changes across distributed brain regions may indicate a central role for the pathophysiology of ITN. Broadly, our findings have implications for understanding how ITN affects local intrinsic brain activity.

These findings are in agreement with recent structural brain imaging results that have indicated that individuals with ITN have altered brain function involved in pain-processing and increased motor output, to control facial movements and to limit pain attacks.

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Disclosure
The authors report no conflicts of interest in this work.

References
1. Jannetta PJ. Observations on the etiology of trigeminal neuralgia, hemifacial spasm, acoustic nerve dysfunction and glossopharyngeal neuralgia. Definitive microsurgical treatment and results in 117 patients. Neurochirurgia (Stuttg). 1977;20(5):145–154.
2. Kondo A. Follow-up results of microvascular decompression in trigeminal neuralgia and hemifacial spasm. *Neurosurgery*. 1997;40(1):46–51.

3. Gronseth G, Cruccu G, Alksne J, et al. Practice parameter: the diagnostic evaluation and treatment of trigeminal neuralgia (an evidence-based review): report of the Quality Standards Subcommittee of the American Academy of Neurology and the European Federation of Neurological Societies. *Neurology*. 2008;71(15):1183–1190.

4. Katusic S, Williams DB, Beard CM, Bergstralh EJ, Kurland LT. The human amygdala and pain: evidence from neuroimaging. *Hum Brain Mapp*. 2001;9(2):229–237.

5. Nakata H, Sakamoto K, Kakigi R. Meditation reduces pain-related neural activity in the anterior cingulate cortex, insula, secondary somatosensory cortex, and thalamus. *Front Psychol*. 2014;5:1489.

6. Wang Y, Li D, Bao F, et al. Impaired trigeminal nociceptive processing in patients with trigeminal neuralgia. *Neurology*. 2007;69(9):835–841.

7. Beck AT, Ward C, Mendelson M, Beck depression inventory (BDI). *Arch Gen Psychiatry*. 1961;4(6):561–571.

8. Hamilton M. The assessment of anxiety states by rating. *Br J Med Psychol*. 1959;32(1):50–55.

9. Sauro KM, Williams DB, Beard CM, Bergstralh EJ, Kurland LT. The human amygdala and pain: evidence from neuroimaging. *Hum Brain Mapp*. 2001;9(2):229–237.

10. Raichle ME, MacLeod AM, Snyder AZ, Powers WJ, Gusnard DA, Shulman GL. A default mode of brain function. *Proc Natl Acad Sci U S A*. 2001;98(2):1157–1160.

11. Li D, Yang C, Yao L, Xiong Y, Li Y, Zang Y. Using coherence to measure regional homogeneity of resting-state fMRI signal. *Front Syst Neurosci*. 2010;4:24.

12. Bantick SJ, Wise RG, Ploghaus A, Clare S, Smith SM, Tracey I. Functional congruence: an index of neural homogeneity and a new measure of brain activity. *Science*. 1968;159(3820):1257–1259.

13. LeDoux JE. Emotion circuits in the brain. *Annu Rev Neurosci*. 2000;23:155–184.

14. Bai XY, Zhang GY, Yan SM. Cuneus and fusiform cortices thickness is reduced in trigeminal neuralgia (tic doloureux). *PLoS One*. 2013;8(6):e66340.

15. Price M, Kubo TT, Doring TM, Tukamoto G, Vincent M, Gasparetto EL. Cuneus and fusiform cortices thickness is reduced in trigeminal neuralgia. *J Headache Pain*. 2014;15:17.

16. Nakata H, Sakamoto K, Kakigi R. Meditation reduces pain-related neural activity in the anterior cingulate cortex, insula, secondary somatosensory cortex, and thalamus. *Front Psychol*. 2014;5:1489.

17. Wang Y, Li D, Bao F, et al. Thalamic metabolic alterations with cognitive dysfunction in idiopathic trigeminal neuralgia: a multivoxel spectroscopy study. *Neuroradiology*. 2014;56(8):685–693.

18. Zuo XN, Xu T, Jiang L, et al. Toward reliable characterization of functional homogeneity in the human brain: pre-processing, scan duration, imaging resolution and computational space. *Neuroimage*. 2013;65:374–386.

19. Zuo XN, Xu T, Jiang L, et al. Toward reliable characterization of functional homogeneity in the human brain: preprocessing, scan duration, imaging resolution and computational space. *Neuroimage*. 2013;65:374–386.

20. Zuo XN, Xu T, Jiang L, et al. Toward reliable characterization of functional homogeneity in the human brain: preprocessing, scan duration, imaging resolution and computational space. *Neuroimage*. 2013;65:374–386.

21. Jiang L, Xu T, He Y, et al. Toward neurobiological characterization of functional homogeneity in the human cortex: regional variation, morphological association and functional covariance network organization. *Brain Struct Funct*. 2015;220(5):2485–2507.
