



Decreased Resting-State Functional Connectivity of Periaqueductal Gray in Temporal Lobe Epilepsy Comorbid With Migraine

Long Wang^{1,2†}, Xin-Ting Cai^{1†}, Mei-Dan Zu^{1†}, Juan Zhang¹, Zi-Ru Deng¹ and Yu Wang^{1*}

¹ Department of Neurology, The First Affiliated Hospital of Anhui Medical University, Hefei, China, ² Department of Neurology, The Second People Hospital of Hefei, Hefei, China

OPEN ACCESS

Edited by:

Giovanni Assenza, Campus Bio-Medico University, Italy

Reviewed by:

Lorenzo Ricci, Policlinico Universitario Campus Bio-Medico, Italy Pierpaolo Croce, University of Studies G. d'Annunzio Chieti and Pescara, Italy

> ***Correspondence:** Yu Wang yw4d@hotmail.com

[†]These authors have contributed equally to this work

Specialty section:

This article was submitted to Epilepsy, a section of the journal Frontiers in Neurology

Received: 02 December 2020 Accepted: 15 April 2021 Published: 26 May 2021

Citation:

Wang L, Cai X-T, Zu M-D, Zhang J, Deng Z-R and Wang Y (2021) Decreased Resting-State Functional Connectivity of Periaqueductal Gray in Temporal Lobe Epilepsy Comorbid With Migraine. Front. Neurol. 12:636202. doi: 10.3389/fneur.2021.636202 **Objective:** Patients with temporal lobe epilepsy (TLE) are at high risk for having a comorbid condition of migraine, and these two common diseases are proposed to have some shared pathophysiological mechanisms. Our recent study indicated the dysfunction of periaqueductal gray (PAG), a key pain-modulating structure, contributes to the development of pain hypersensitivity and epileptogenesis in epilepsy. This study is to investigate the functional connectivity of PAG network in epilepsy comorbid with migraine.

Methods: Thirty-two patients with TLE, including 16 epilepsy patients without migraine (EwoM) and 16 epilepsy patients with comorbid migraine (EwM), and 14 matched healthy controls (HCs) were recruited and underwent resting functional magnetic resonance imaging (fMRI) scans to measure the resting-state functional connectivity (RsFC) of PAG network. The frequency and severity of migraine attacks were assessed using the Migraine Disability Assessment Questionnaire (MIDAS) and Visual Analog Scale/Score (VAS). In animal experiments, FluoroGold (FG), a retrograde tracing agent, was injected into PPN and its fluorescence detected in vIPAG to trace the neuronal projection from vIPAG to PPN. FG traced neuron number was used to evaluate the neural transmission activity of vIPAG-PPN pathway. The data were processed and analyzed using DPARSF and SPSS17.0 software. Based on the RsFC finding, the excitatory transmission of PAG and the associated brain structure was studied via retrograde tracing in combination with immunohistochemical labeling of excitatory neurons.

Results: Compared to HCs group, the RsFC between PAG and the left pedunculopontine nucleus (PPN), between PAG and the corpus callosum (CC), was decreased both in EwoM and EwM group, while the RsFC between PAG and the right PPN was increased only in EwoM group but not in EwM group. Compared to EwoM group, the RsFC between PAG and the right PPN was decreased in EwM group. Furthermore, the RsFC between PAG and PPN was negatively correlated with the frequency and severity of migraine attacks. In animal study, a seizure stimulation

1

induced excitatory transmission from PAG to PPN was decreased in rats with chronic epilepsy as compared to that in normal control rats.

Conclusion: The comorbidity of epilepsy and migraine is associated with the decreased RsFC between PAG and PPN.

Keywords: epilepsy, migraine, comorbidity, periaqueductal gray, pedunculopontine nucleus, functional magnetic resonance imaging

INTRODUCTION

Temporal lobe epilepsy (TLE) is the most common form of epilepsy which is often characterized by drug resistance (1). Imaging evidences have shown that brain network abnormalities in TLE are not limited within the epileptogenic focus but rather throughout the whole brain (2, 3). TLE is at high risk for having comorbid condition of migraine, and these two common diseases are proposed to have some shared pathophysiological mechanisms (4, 5). Moreover, comorbid migraine may be a predictor of medically refractory epilepsy, indicating its impact on epilepsy (6). Therefore, a better understanding of the neurophysiological mechanism related to the comorbidity of epilepsy and migraine is helpful to develop more effective treatments.

The midbrain periaqueductal gray (PAG) is a group of structural and functional heterogeneous nuclei, which is divided into four independent but interconnected subregions: ventrolateral PAG (vlPAG), lateral PAG (lPAG), dorsomedial PAG (dmPAG), and dorsolateral PAG (dlPAG) based on cytoarchitectonics, histochemistry, and neuroimaging studies (7, 8). As a functional heterogeneous nuclei, PAG is a key brainstem structure for modulation of multiple neurophysiological functions including pain (9, 10), emotion (11), respiration, and cardiovascular activity (12). Especially, PAG is the primary center of the descending pain inhibition system involved in regulation of both endogenous pain and trigeminovascular pathway pain (9, 10). Structural and functional alterations of PAG have been identified in migraine (13-17), and it has been proposed that PAG dysfunction results in trigeminovascular neuron hyperexcitability leading to migraine headache (18). This is further supported by multiple clinical case descriptions that PAG lesion, especially ventrolateral PAG (vlPAG) lesion, can initiate migraine-like headache attacks (19-21). In fact, the dysfunction of PAG has now been considered a "generator" of migraine headache (22, 23). Experimental studies have demonstrated vlPAG is critically involved in the development of post-ictal antinociception in animal study as well as in that of pain sensitivity decrement during the interphase of chronic epilepsy in human study (24, 25). In addition, we have recently demonstrated that vlPAG is not only involved in pain regulation in epilepsy but also in the development of epileptogenesis (26).

Another upper brainstem structure, pedunculopontine nucleus (PPN), which has functional afferent as well as efferent connections with multiple brain structures including vlPAG and structures outside of brainstem (27), is also closely related to the post-ictal antinociception (28, 29). Some studies have also demonstrated that stimulation of PPN has anti-epileptic effects, though this method of PPN stimulation has not been accepted in clinical practice (30, 31).

Taken together, it is clear that vlPAG and PPN are both involved in modulation of pain and epilepsy. Thus, we speculate that there may be an abnormal functional connection between vlPAG and PPN in epilepsy patients with comorbid migraine. However, vlPAG is difficult to be completely separated from PAG in neuroimaging because of the complexity and particularity of PAG structure, so we conducted the current study with a whole brain functional connectivity (FC) selecting PAG as the region of interest (ROI). And we will preliminarily verify the findings of the clinical imaging study through animal experiments.

MATERIALS AND METHODS

Patient Study Participants

A total of 32 TLE patients with bilateral tonic-clonic seizures were recruited from the outpatient department at the First Affiliated Hospital of Anhui Medical University in Hefei from August 2019 to January 2020. Sixteen TLE patients were comorbid with migraine (EwM group) and 16 without migraine (EwoM group). The inclusion criteria were as follows: (1) patients with epilepsy of only bilateral tonic-clonic seizures; (2) seizure types of bilateral tonic-clonic seizures were established according to the classification of the International League Against Epilepsy (ILAE) (32-34); (3) all patients accepted 3.0 T magnetic resonance imaging (MRI) examination to exclude intracranial tumor, changes after cerebrovascular disease, post traumatic changes, and other conditions associated with brain structure alterations; (4) Mini-mental State Examination (MMSE) score \geq 24 points (35); (5) all patients were righthanded. The exclusion criteria were as follows: (1) patients with a history of other neurological, mental, or chronic diseases, such as schizophrenia, diabetes, etc.; (2) pregnant and nursing women; (3) patients with a history of metal dentures or implants; (4) patients who experienced generalized bilateral tonic-clonic seizure within 1 week prior to fMRI scanning; (5) excessive motion (>2.5 mm, 2.5°) during fMRI scanning.

Abbreviations: TLE, temporal lobe epilepsy; EwoM, epilepsy patients without migraine; EwM, epilepsy patients with comorbid migraine; PAG, periaqueductal gray; RsFC, resting-state functional connectivity; VAS, visual analog scale/score; PPN, pedunculopontine nucleus; CC, corpus callosum; rs-fMRI, resting-state functional magnetic resonance imaging; MMSE, mini-mental state examination; HDRS, Hamilton depression rating scale; ROI, region of interest.

All epilepsy patients with or without migraine were on antiepileptic treatment with monotherapy or combination therapy. All epilepsy patients with migraine had never accepted special treatment for migraine prophylaxis though some antiepileptic drugs may have anti-migraine effects. Fourteen right-handed healthy participants matched with the epilepsy group in age, sex, and educational level were recruited as healthy controls (HCs group).

Clinical Assessment and Analysis

Clinical information including demographic data, education level, past history of medical, and using of anti-depressive drugs were collected from all patients. Mini-mental State Examination (MMSE) and Hamilton Depression Rating Scale (HDRS) were, respectively used to evaluate the cognitive function and depressive symptoms of the participants (35, 36). Migraine head pain severity was measured by means of a Visual Analog



FIGURE 1 | Location of seeds and ROI. (A) The ROI of the PAG was obtained from the Keuken and Forstmann's 7T atlas that provides ROIs obtained from high-resolution MP2RAGE and FLASH scans warped in MNI space (Keuken and Forstmann, 2015), Available at https://www.nitrc.org/project cts/atag/; (B) The ROI of double PPN was generated by placing two 5 mm radius spheres with centers at coordinates [±6.4, -27, -15] in MNI standard space.

TABLE 1 | Demographic and clinical characteristics of the participants.

Characteristics	HCs (<i>n</i> = 14)	EwoM (<i>n</i> = 16)	EwM (<i>n</i> = 16)	Р
Gender (M/F)	7/7	7/9	6/10	0.774*
Age, years, mean (SD)	25.6 (4.6)	28.8 (5.9)	25.3 (6.3)	0.169
Education, years, mean (SD)	12.6 (3.8)	14.0 (4.0)	11.9 (3.4)	0.276
MMSE, scores, mean (SD)	29.4 (0.9)	29.0 (1.8)	28.3 (1.9)	0.178
HDRS, scores, mean (SD)	1.4 (0.9)	2.5 (1.3)	7.6 (3.7)	< 0.001
Epilepsy duration, month, mean (SD)	-	81.3 (53.7)	75.3 (57.8)	0.766
Mean seizure frequency, times/year, mean (SD)		2.3 (1.8)	4.8 (2.7)	0.005
Mean migraine frequency, days/month, mean (SD)		-	1.5 (0.5)	
Form of migraine				
IIM (patients number)			7	
PIM		-	12	
MwA		-	1	
MwoA		-	15	
VAS, scores, mean (SD)		-	4.5 (1.4)	
Medications				
Monotherapy (patients number)		12	6	0.033*
Drug combination		4	10	

SD, standard deviation; HCs, healthy controls; EwoM, epilepsy without migraine; EwM, epilepsy with comorbid migraine; MMSE, Mini-mental State Examination; HDRS, Hamilton depression rating scale; IIM, inter-ictal migraine; PIM, post ictal migraine; MwA, migraine with aura; MwoA, migraine without aura; VAS, visual analog scale/score; n, number. Medications presents antiepileptic drugs.

*chi-square fisher exact test.

Scale (VAS) with 0 indicating no head pain and 10 indicating unbearable head pain (37). The statistical analyses of the clinical data were conducted with SPSS version 16.0 (SPSS lnc., Chicago, USA), with a significant threshold of P < 0.05.

Neuroimaging Data Acquisition

The resting-state functional magnetic resonance imaging (rsfMRI) was obtained using a 3.0 T MRI scanner (Discovery GE750w; GE Healthcare, Buckinghamshire, UK) from the University of Science and Technology of China. All participants were required to keep their eyes closed without either thinking about anything or moving the body during scanning. MR imaging system was composed of 217 echo-planar imaging sequences with the following parameters: repetition time/echo time (TR/TE): 2,000/30 ms; flip angle: 90°; matrix size: 64 × 64; field of view (FOV): 192 × 192 mm²; slice thickness: 3 mm; 46 continuous slices (voxel size: $3 \times 3 \times 3 \text{ mm}^3$). T1-weighted anatomic images with 188 slices were also obtained in sagittal orientation with the following parameters: TR: 8.16 ms; TE: 3.18 ms; flip angle: 12°; FOV: 256 × 256 mm²; slice thickness: 1 mm; voxel size: $1 \times 1 \times 1 \text{ mm}^3$.

rs-fMRI Preprocessing

Functional image preprocessing was carried out using the Data Processing Assistant for rs-fMRI toolkit software package based on Statistical Parametric Mapping software (SPM12; www.fil.ion.ucl.ac.uk/spm) and rs-fMRI Imaging Toolkit (38, 39). Software was preprocessed in the following steps: conversion of the DICOM data to NIFTI images; removal of the first 10 time points to exclude the influence of unstable longitudinal magnetization; slice timing correction; motion correction; realignment to respective structural image; nuisance regressors, spatial normalization based on the segmentation of structural images, spatial smoothing (Gaussian kernel, FWHM = 4 mm), and filtering temporal band pass (0.01-0.1 Hz).

RsFC Processing and Analysis

The mask of the PAG was obtained from the Cacciola and Keuken's 7T atlas that provides ROIs obtained from highresolution MP2RAGE and FLASH scans warped in MNI space available at www.nitrc.org/projects/atag/ (40, 41) (Figure 1A). The location of the PPN was generated by placing two 5 mm radius spheres with centers at coordinates $[\pm 6.4, -27, -15]$ in MNI standard space (42, 43) (Figure 1B). To obtain the value of RsFC, the mean time series of PAG was extracted and then used to calculate the Pearson's correlations (r) with voxels in the remaining brain in each participant. The Fisher's z r-to-z transformation was applied to improve the normality, and then the results were displayed with an RsFC map for each participant. To explore RsFC differences among 3 groups, independent samples t-tests and one-way analysis of variance (ANOVA) were performed, respectively using DPABI software within a mask of whole brain gray matter. Moreover, clinical data including HDRS, medications, and seizure duration and frequency may have influence on RsFC results (44) as these confounding factors were statistically different between groups (Table 1). Thus, HDRS, medications, and seizure duration and frequency were used as covariables for calculation. *Post-hoc* analyses were conducted using Gaussian Random Field method (GRF) with the significance of voxel level set at P < 0.001, and that of cluster level set at P < 0.05, two-tailed.

Correlation Analysis

The RsFC of the PAG with corresponding significant areas were extracted and then Pearson's correlation analysis was conducted to detect the relationship between the values of RsFC and seizure events. The correlation analysis was performed by SPSS version 16.0 with a significant threshold of P < 0.05.

Animal Study

Based on the functional imaging findings that the FC of PAG-PPN and PAG-CC were decreased in epilepsy patients, the animal study is aimed to address the alterations of the neural transmission from vIPAG to PPN in epileptic rats, using retrograde tracing and fluorescence immunohistochemistry.

Experimental Animals

Male Sprague-Dawley (SD) rats aged 4–5 weeks and weighed between 80 and 110 g, from the Animal Center of Anhui Medical University, were used for the experiment. The animals were kept in a temperature-controlled ($22 \pm 1^{\circ}$ C) room with free access to food and water. The room was maintained on a light and dark cycle (12 h: 12 h, light from 6:00 to 18:00) and kept air humidity around 50%. The research complied with the national legislation on the Care and Use of Animals.

Experimental Design and Group Division

To investigate the activity of neural transmission from vlPAG to PPN, a retrograde tracing agent, FluoroGold (FG, Fluorochrome, USA), was injected into PPN and the traced neurons will be visualized in vlPAG. The FG-traced neuron number within vlPAG will be used to identify the neural transmission activity of vlPAG-PPN pathway. This FG retrograde tracing method has been widely used to study the activity of neuronal connections between different parts within the brain associated with distinct functions (45-47). For comparison of the neural transmission activity of vlPAG-PPN pathway between normal rats and chronic epileptic rats, seizure stimulation activating vlPAG-PPN pathway should be similar in severity and duration. Thus, status epilepticus (SE) was evoked with pilocarpine injection in naïve rats (Pilo group) as well as in rats with chronic epilepsy (1 month after SE induction) developed after SE induced by pre-treatment of pilocarpine injection (SE+Pilo group). In Pilo group, rats were stereotactically injected with FG to PPN 2 days before SE induction. In SE+Pilo group, rats were firstly induced with SE by plocarpine, and 28 days later, the rats were stereotactically injected with FG to PPN, and then, 2 days after FG injection, re-injected with pilocarpine to evoke a second SE as seizure stimulation. One hour after SE induction, i.e., seizure stimulation, rats were sacrificed for immunohistochemical staining.

Induction of SE

As previously described (48, 49), rats were firstly pre-treated with scopolamine methylbromide [1 mg/kg, intraperitoneal (i.p.),

Sigma, USA] to prevent peripheral cholinergic effects caused by pilocarpine. Thirty minutes later, rats were then injected with pilocarpine (hydrochloride) dissolved in physiological saline (350 mg/kg/2 ml, i.p., Cayman, USA) to induce SE. For some rats, reinjections of 0.2 ml pilocarpine solution might be needed with an interval of 15 min. Behavioral seizures were recorded using previous five-stage scale modified mildly from Racine's scale (50): briefly, stage 1, face and vibrissae twitching, ear rubbing with forepaws, chewing; stage 2, head nodding or with unilateral limb clonus; stage 3, bilateral limb clonus, mild whole body convulsions; stage 4, lockjaw, rearing, whole body convulsions

TABLES	Dogione with		difforonco	of PoEC	botwoon groups
IADLE 2	negions with	i signincant	unierence	UI NSEC	between-groups.

Brain region	MNI coordinate (x,y,z)	Voxel size	P-value
EwoM vs. HCs			
Left PPN	10, -23, -17	107	-6.624
Right PPN	-5, -30, -6	62	7.528
Corpus callosum	21, 17, 15	67	-5.749
EwM vs. HCs			
Left PPN	6, -25, -14	72	-6.912
Corpus callosum	25, 16, 10	59	-7.324
EwM VS EwoM			
Right PPN	3, 16, -9	26	-6.940

HCs, healthy control; EwoM, epilepsy without migraine; EwM, epilepsy with comorbid migraine.

with tail hypertension; and stage 5, rearing with whole body convulsions and falling down with whole body rigidity. SE onset was defined as stage 3 or greater seizures that developed to repeated or prolonged behavioral seizures. Diazepam (10 mg/kg, i.p.) was administrated 60 min after SE induction to terminate convulsion. The duration of SE (seizure stimulation) in Pilo group rats was the same as that in SE+Pilo group rats as the seizures were terminated with diazepam when the SE duration reached 1 h. The stimulating seizure severity in Pilo group rats was close to that in SE+Pilo group rats as it reached SE in both groups.

Stereotactic Injection of FG

Rats were anesthetized by intraperitoneal injection of a mixed solution of 2% sodium pentobarbital (80 mg·kg⁻¹) and xylazine (10 mg·kg⁻¹). Fully anesthetized animals with regular respiratory rate were mounted on a stereotactic apparatus (RWD Life Science, China). A rostro-caudal incision was made in the skull skin after shearing hairs and full disinfecting with 1% iodophor. To expose bregma and posterior fontanelle completely, swabs were used to dip medical H_2O_2 to wipe soft tissues slightly. After adjusting bregma and posterior fontanelle to the same horizontal level, a hole was drilled and microinjection syringe was vertically introduced using the following coordinates based on Paxinos and Watson's rat brain in the stereotactic coordinates atlas: anteroposterior (AP), 0.2 mm; mediolateral (ML), 2.0 mm; and dorsoventral (DV), 7.0 mm to reach the PPN (28, 51). FG (Fluorochrome, USA) dissolved in physiological



FIGURE 2 | Group differences in resting-state functional connectivity (RsFC) to PAG. (A) Differences in RsFC to PAG between the EwoM group and HCs group; (B) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and HCs group; (C) Differences in RsFC to PAG between the EwoM group and EwoM group.

saline (4%) was unilaterally injected into the PPN at a rate of 0.25 μl per minute for 2 min using microinjection syringe fixed to the stereotaxic frame. The microinjection syringe remained there for 1 min to ensure FG was fully absorbed and then slowly pulled out. An intramuscular injection of penicillin (30 U/kg) was conducted to prevent infection after skull skin was sutured.

Tissue Preparation

One hour after SE (second SE for SE+Pilo group rats) was terminated, rats were anesthetized with pentobarbital (80 mg·kg⁻¹) and xylazine (10 mg·kg⁻¹) and perfused with 250 ml 0.9% physiological saline followed by 200 ml 4% paraformaldehyde in PBS (pH 7.4). Next, the brain was removed immediately, fixed in 4% paraformaldehyde at 4°C overnight, dehydrated in 30% sucrose at 4°C, and then quick frozen on dry ice and stored at -80° C. Serial coronal sections (10 µm) containing the vlPAG were obtained by a cryostat (CM 3050S, Leica, Germany). Each section has an interval of about 70 µm, and six sections taken from each rat were attached to poly-lysine-coated slides for immunofluorescence study.

Fluorescence Immunohistochemistry

Sections were incubated with 0.3% Triton X-100 in PBS for 30 min, blocked with 2% normal goat serum in PBS for 1 h at room temperature, and then incubated overnight at 4°C with primary antibodies: rabbit anti-c-fos (Abcam, 1:200) and guinea pig anti-vGlut1 (Millipore, 1:100). Sections were then incubated for 2 h at room temperature with the corresponding secondary antibodies: donkey anti-rabbit IgG conjugated with Alexa Fluor 594 (Invitrogen, 1:500) and goat anti-guinea pig IgG conjugated with tetraethyl rhodamine isothiocyanate (TRITC) (ImmunoReagents, 1:300). For negative controls, adjacent sections were treated with the same steps except by replacing the primary antibodies with PBS. Two symmetric images (\times 200) in the vIPAG area were acquired at the same exposure in each sight of 5–7 sections per subgroup using Nikon Eclipse 80i fluorescence microscope (Nikon, Japan).

Quantification and Statistical Analysis

Counting of FG-traced neurons, c-Fos- and vGlut1-labeled cells, and FG-traced neurons labeled by c-Fos or by vGlut1 was conducted on one of every 20 sections across the vIPAG, i.e., 5-7 sections per animal. On each section, two counting frames were symmetrically placed ventrolaterally to the midbrain aqueduct on images of ×200 magnification basing on the Rat Brain in Stereotaxic Coordinates (6th edition, Paxinos and Watson) (52). Cell counting was performed by two independent researchers to minimize counting bias. The results were recorded as the mean number of cells per frame. Fluorescence intensity of FG with and without c-Fos labeling within the frames mentioned above was measured separately, and those of FG with and without vGlut1 labeling measured separately using an Image J software (Image J 1.51n, USA). Here, thresholding background of the captured images for background subtracting was conducted. The measurements were made on 5-7 separate sections per animal, in 6 animals per group.

Statistical Analysis

All data were presented as mean \pm standard error of the mean (M \pm SEM). One-way analysis of variance (ANOVA) was applied to analyze the differences in cell number among groups and the differences between two groups for comparison. All statistical analyses were figured out by SPSS 16.0, and statistical significance was set at *P* < 0.05. Pairwise comparison was conducted by *post hoc* test after ANOVA, Tukey HSD, or Dunnett T3 depending on Test of Homogeneity of Variances. All histograms and graphs were performed with GraphPad Prism 6.02. Images were managed with Adobe Photoshop CS6.

RESULTS

Analysis of Clinical Data

As shown in **Table 1**, there was no significant difference in age, gender, years of education, duration of epilepsy, and MMSE score among the three groups (P > 0.05). However, mean seizure







FIGURE 5 | epilepsy rats (solid bar) was significantly higher than that in non-stimulated control rats (open bar), but the mean number of c-Fos immunoreactive cells after a seizure stimulation in chronic epilepsy rats (solid bar) was significantly lower than that in naïve rats (meshy bar) (**B**) The mean number of c-Fos immunoreactive cells traced by FG after a seizure stimulation either in naïve rats (meshy bar) or in chronic epilepsy rats (solid bar) was significantly lower than that in non-stimulated control rats (open bar), but the mean number of c-Fos immunoreactive cells traced by FG after a seizure stimulation in chronic epilepsy rats (solid bar) was significantly lower than that in non-stimulated control rats (open bar), but the mean number of c-Fos immunoreactive cells traced by FG after a seizure stimulation in chronic epilepsy rats (solid bar) was significantly lower than that in naïve rats (meshy bar) (**C**) A One-way ANOVA was performed. Different than the control group, ^{*}*P* < 0.05, ^{***}*P* < 0.001. Different than the Pilo group, [^]*A* < 0.01, [^]*P* < 0.05, *n* = 6 per group, scale bar = 100 µm.

frequency and HDRS scores in EwM group were higher than that in EwoM group (P < 0.05, **Table 1**).

Group Differences in RsFC of the PAG

Compared to the HCs group, both EwoM and EwM groups showed decreased RsFC between PAG and left PPN (PAG-left PPN) and decreased RsFC between PAG and corpus callosum (CC) (PAG-CC), whereas PAG exhibited increased RsFC with the right PPN (PAG-right PPN) in EwoM group but not in EwM group. When comparing with EwoM group, EwM group exhibited decreased PAG-right PPN FC. However, we did not find significant difference in PAG-CC RsFC between EwM and EwoM groups. The significance of RsFC of PAG and corresponding significant clusters are showed in **Table 2** and **Figure 2**.

Correlation Analysis Between RsFC of PAG and Clinical Seizure Events

A significant negative correlation (r = -0.526, P = 0.037) was observed between VAS scores and the RsFC value of PAG-left PPN, where patients with lower RsFC value in this circuit had greater VAS scores (**Figure 3**). The RsFC value of PAG-left PPN was negatively correlated with the migraine attack frequency (r = -0.502, P = 0.048; **Figure 4**). In other words, patients with lower RsFC in this circuit had higher migraine attack frequency. In addition, our results did not find significant correlations between RsFC value of PAG-CC and migraine attack frequency (r = -0.048, P = 0.861) or with VAS scores (r = 0.339, P = 0.198).

Alterations of FG-Traced Neurons Labeled by c-Fos After Seizure Stimulation in Rats With Chronic Epilepsy

The expression of c-Fos positive cells as well as FG traced cells in vlPAG were less in normal control group (Figure 5A). Seizure induced increase of c-Fos labeled cells in vlPAG both in naïve rats (Pilo group) and chronic epilepsy rats (SE+Pilo group) compared to control rats, but a significant lower increase in SE+Pilo group rats compared to Pilo group rats [Figure 5B, One-way ANOVA for comparison among groups, $F_{(2,18)}$ = 12.135, p = 0.001; Pilo vs. control, p = 0.000; SE+Pilo vs. control, p = 0.042; SE+Pilo vs. Pilo, p = 0.007]. Seizure induced increase of c-Fos labeled cells traced by FG both in Pilo group and in SE+Pilo group rats, but a significant lower increase in SE+Pilo group rats compared to Pilo group rats [Figure 5C, One-way ANOVA for comparison among groups, $F_{(2,18)} = 10.866$, p =0.001. Pilo vs. control, p = 0.000; SE+Pilo vs. control, p =0.029; SE+Pilo vs. Pilo, p = 0.040]. This indicates that fewer vlPAG neurons projecting to PPN were activated after seizure stimulation in rats with chronic epilepsy as compared to that in normal control rats.

Alterations of FG-Traced Neurons Labeled by vGlut1 After Seizure Stimulation in Rats With Chronic Epilepsy

In normal control rats, few vGlut1 labeled cells presented in vlPAG, and few vGlut1 labeled neurons traced by FG presented (Figure 6A). Seizure induced increase of vGlut1 labeled cells in vlPAG both in naïve rats (Pilo group) and in chronic epilepsy rats (SE+Pilo group) compared to control rats, but a significant lower increase in SE+Pilo group rats compared to Pilo group rats [Figure 6B, One-way ANOVA for comparison among groups, $F_{(2,18)} = 9.907$, p = 0.002; Pilo vs. control, p = 0.000; SE+Pilo vs. control, p < 0.046; SE+Pilo vs Pilo, p < 0.038]. Seizure induced increase of vGlut1 labeled cells traced by FG both in Pilo group and in SE+Pilo group rats, but a significant lower increase in SE+Pilo group rats compared to Pilo group rats (Figure 6C, One-way ANOVA for comparison among groups, $F_{(2,18)} = 11.542$, p = 0.001. Pilo vs. control, p = 0.000; SE+Pilo vs. control, p = 0.028; SE+Pilo vs. Pilo, p = 0.032). This indicates that fewer vlPAG excitatory neurons projecting to PPN were activated after seizure stimulation in rats with chronic epilepsy as compared to that in normal control rats.

DISCUSSION

We are the first to demonstrate the functional alterations of PAG and its correlation with migraine in patients with epilepsy. We found that the FC of PAG-left PPN and PAG-CC was decreased in EwoM as well as in EwM group comparing with HCs group. An interesting phenomenon is that the PAG-right PPN FC was increased in EwoM group comparing with HCs group, which was not shown in EwM group. When comparing with EwoM group, EwM group exhibited decreased PAG-right PPN FC. Finally, the FC between PAG and PPN was negatively correlated with the attack frequency and VAS scores of migraine headache in epilepsy.

The migraine prevalence is significantly higher in patients with TLE than that in the general population (53). In fact, the interactive comorbid relationship between epilepsy and migraine has been well-noticed, but the underlying pathophysiological mechanism is not clear, though some possible shared mechanisms including genetic mechanism has been identified (54). For pain processing within brainstem, PAG is known to interact with many other brainstem nuclei such as the rostral ventromedial medulla and the subnucleus reticularis dorsalis that directly or indirectly project to the spinal dorsal



9

FIGURE 6 | or in chronic epilepsy rats (solid bar) was significantly higher than that in non-stimulated control rats (open bar), but the mean number of vGlut1 immunoreactive neurons after a seizure stimulation in chronic epilepsy rats (solid bar) was significantly lower than that in naïve rats (meshy bar) (**B**) The mean number of vGlut1 immunoreactive neurons traced by FG after a seizure stimulation either in naïve rats (meshy bar) or in chronic epilepsy rats (solid bar) was significantly higher than that in non-stimulated control rats (open bar), but the mean number of vGlut1 immunoreactive neurons traced by FG after a seizure stimulation either in naïve rats (meshy bar) or in chronic epilepsy rats (solid bar) was significantly higher than that in non-stimulated control rats (open bar), but the mean number of vGlut1 immunoreactive neurons traced by FG after a seizure stimulation in chronic epilepsy rats (solid bar) was significantly lower than that in naïve rats (meshy bar) (**C**) A One-way ANOVA was performed. Different than the control group, "P < 0.05, "P < 0.01, "**P < 0.01. Different than the Pilo group, $^{A}P < 0.05$. n = 6 per group, scale bar = 100 µm.

horn and trigeminal nucleus (55). It is also demonstrated that decreased FC between PAG and other brain regions is involved in migraine headache basing on clinical neuroimaging study and experimental animal study (14, 56). In the current study, we found that the FC of PAG with other brain regions was decreased in TLE patients. Further, the FC of PAG was more remarkably decreased in TLE patients with comorbid migraine, and this FC was negatively correlated with the attack frequency and VAS scores of migraine headache. This indicates that the dysfunction of PAG network may be involved in the comorbidity of migraine in epilepsy.

The PPN, an elongated nucleus located in the rostral locomotor region of the brainstem, has functional afferent and efferent connections with multiple brain structures including PAG and structures outside of brainstem (27, 57). PPN has been shown to be crucial for the regulation of pain (58) as well as the regulation of seizure-induced hypoalgesia, a phenomenon of postictal antinociception due to the activation of PPN by seizure (28, 29). We demonstrated, for the first time, that the FC between PAG and left PPN is decreased during the interictal phase of seizures in patients with epilepsy. As the activation of PAG and PPN both have a pain suppressing effect, the decreased FC between PAG and PPN may give rise to a reduced pain threshold underlying migraine occurrence in epilepsy. Neuroimaging study has shown decreased activity in PAG in animals with epileptic seizures (59). This is consistent with our previous findings that there is a continuing loss of vlPAG neurons in rats with chronic epilepsy and neurochemical lesion to PAG decreases the pain threshold in epileptic rats (26). Furthermore, our current animal study showed that fewer vIPAG cells doublelabeled with FG and c-Fos were activated in the chronic stage of epilepsy, indicating the decrease of excitatory transmission between PAG and PPN. The decreased FC between PAG and PPN in patients with TLE may be a representation of attenuated neural transmission from PAG to PPN in epilepsy rats. Previous studies have shown that both PAG and PPN activation have anti-epileptic effects (26, 60-62). Taken together, PAG and PPN both have antiepileptic and antinociceptive function, thus the decreased PAG-PPN functional connection might be proposed to be an underlying pathomechanism of the interactive comorbid relationship between epilepsy and migraine.

An interesting finding is the increased connectivity between PAG and the right PPN in patients without comorbid migraine but not in patients with comorbid migraine. This increased connectivity between PAG and right PPN seemed in conflict with the decreased connectivity between PAG and left PPN in patients without comorbidity of migraine. This discrepancy between the left and right PPN in connectivity with PAG might be due to the lateralization of seizure onset origin and a compensatory mechanism in epilepsy, though the sample size was not sufficient to analyze the influence of the lateralization of seizure onset origin. The finding that PAG exhibited increased RsFC with the right PPN in EwoM group but not in EwM group may also reflect the more generally favorable epilepsy outcome in the EwoM group of patients. This would be in line with the results from a recent work showing that TLE patients with increased level of connectivity had favorable outcome after anti-epileptic therapy (44).

CC connects the left and right cerebral hemispheres and mediates the interhemispheric transferring of signals, influencing on movements, sensory, emotion, behavior, cognition, memory, and complex integrative functions (63). In epilepsy, bilateral seizures with focal onset spread rapidly through CC and cause secondary structural damage related to demyelination in CC (64, 65). This is further supported by previous finding that decrement of CC connectivity is associated with disease history of TLE (66, 67). White matter including CC has been demonstrated with abnormalities in chronic pain including peripheral neuralgia and headache but less in episodic pain, indicating the CC abnormalities are the consequence of chronic pain attacking as the abnormalities are associated with pain intensity (68-70). This is further supported by the reversible CC lesion after headache attack in migraine patients (71-73). From literatures, the CC abnormalities are also associated with epilepsy as well as migraine attacks. Thus, the decreased PAG-CC connectivity is a logical accompaniment of epilepsy with or without comorbid migraine, though the decrement of the FC of PAG-CC was not correlated with the frequency of seizure or migraine attacks. In fact, this correlation cannot be excluded because of the limited size of the recruited patients in current study. The role of this decreased PAG-CC connectivity either in the development of migraine or in that of epilepsy is unknown, but this decreased connectivity might be associated with other clinical accompaniments of epilepsy and migraine, such as emotion, memory, sleep, etc., of which both PAG and CC are involved in the regulation (74–77).

In conclusion, an abnormal PAG neural network is present in TLE patients. It might be proposed that the decreased RsFC of PAG-PPN pathway is also associated with comorbidity of epilepsy and migraine. Thus, the disrupted connection between PAG and PPN may be an underlying pathophysiological mechanism for the interactive relationship between epilepsy and migraine.

LIMITATIONS

Some limitations should be noted in this study. Firstly, the relatively few participants may limit the statistical power. For example, the lateralization of seizure onset origin cannot be used

to analyze its association with lateral alteration of PAG-PPN connectivity and the correlation of PAG-CC connectivity with the frequency of seizure or migraine attacks. Thus, it is necessary to repeat our experiment by enlarging the sample size to better interpret the current results. Secondly, all the participants were on antiepileptic medication which may affect the FC as described in previous studies (78, 79). Thirdly, this is a cross-sectional study which cannot provide longitudinal alteration data of the participants, thus it cannot be determined whether the PAG functional alteration could be used to predict treatment outcomes in TLE patients comorbid with migraine. Fourthly, a group of patients with only migraine was not included in this study, and this might weaken the conclusion that the dysfunction of PAG network may play a causative role in the comorbidity of migraine in epilepsy.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Ethics Committee of the First Affiliated Hospital of Anhui Medical University. The patients/participants provided their written informed consent to participate in this

REFERENCES

- Tellez-Zenteno JF, Hernandez-Ronquillo L. A review of the epidemiology of temporal lobe epilepsy. *Epilepsy Res Treat.* (2012) 2012:630853. doi: 10.1155/2012/630853
- Alvim MK, Coan AC, Campos BM, Yasuda CL, Oliveira MC, Morita ME, et al. Progression of gray matter atrophy in seizure-free patients with temporal lobe epilepsy. *Epilepsia*. (2016) 57:621–29. doi: 10.1111/epi.13334
- Englot DJ, Konrad PE, Morgan VL. Regional and global connectivity disturbances in focal epilepsy, related neurocognitive sequelae, and potential mechanistic underpinnings. *Epilepsia*. (2016) 57:1546–57. doi: 10.1111/epi.13510
- Keezer MR, Bauer PR, Ferrari MD, Sander JW. The comorbid relationship between migraine and epilepsy: a systematic review and meta-analysis. *Eur J Neurol.* (2015) 22:1038–47. doi: 10.1111/ene.12612
- Kingston WS, Schwedt TJ. The relationship between headaches with epileptic and non-epileptic seizures: a narrative review. *Curr Pain Headache Rep.* (2017) 21:17. doi: 10.1007/s11916-017-0617-9
- Velioglu SK, Boz C, Ozmenoglu M. The impact of migraine on epilepsy: a prospective prognosis study. *Cephalalgia*. (2005) 25:528– 35. doi: 10.1111/j.1468-2982.2005.00912.x
- Menant O, Andersson F, Zelena D, Chaillou E. The benefits of magnetic resonance imaging methods to extend the knowledge of the anatomical organisation of the periaqueductal gray in mammals. *J Chem Neuroanat.* (2016) 77:110–20. doi: 10.1016/j.jchemneu.2016.06.003
- Beitz AJ. The midbrain periaqueductal gray in the rat. I. Nuclear volume, cell number, density, orientation, and regional subdivisions. J Comp Neurol. (1985) 237:445–59. doi: 10.1002/cne.9023 70403
- Yamamotova A. Endogenous antinociceptive system and potential ways to influence It. *Physiol Res.* (2019) 68(Suppl. 3):S195– S205. doi: 10.33549/physiolres.934351

study. The animal study was reviewed and approved by the Ethics Committee of the First Affiliated Hospital of Anhui Medical University.

AUTHOR CONTRIBUTIONS

LW, X-TC, and YW made substantial contributions to conception and design of the study. LW, M-DZ, JZ, and Z-RD were responsible for subject recruitment, case diagnosis, and fMRI data acquisition and analysis. LW and X-TC were primarily responsible for animal experiments. LW, X-TC, and M-DZ contributed to statistical analysis and interpretation of data. LW, X-TC, and YW were responsible for drafting the manuscript and revising it. All authors approved the final version of the manuscript.

FUNDING

This work was supported by Natural Science grants to YW (Grant number: 81671290) from the National Natural Science Foundation of China.

ACKNOWLEDGMENTS

We thank the Research and Experimental Center of Anhui Medical University for providing the experimental platform.

- Henssen D, Dijk J, Knepfle R, Sieffers M, Winter A, Vissers K. Alterations in grey matter density and functional connectivity in trigeminal neuropathic pain and trigeminal neuralgia: a systematic review and meta-analysis. *NeuroImage*. (2019) 24:102039. doi: 10.1016/j.nicl.2019.102039
- Venkatraman A, Edlow BL, Immordino-Yang MH. The brainstem in emotion: a review. Front. Neuroanat. (2017) 11:15. doi: 10.3389/fnana.2017.00015
- Dampney R. Emotion and the cardiovascular system: postulated role of inputs from the medial prefrontal cortex to the dorsolateral periaqueductal gray. *Front Neurosci.* (2018) 12:343. doi: 10.3389/fnins.2018.00343
- Li Z, Liu M, Lan L, Zeng F, Makris N, Liang Y, et al. Altered periaqueductal gray resting state functional connectivity in migraine and the modulation effect of treatment. *Sci Rep.* (2016) 6:20298. doi: 10.1038/srep20298
- Chen Z, Chen X, Liu M, Liu S, Ma L, Yu S. Disrupted functional connectivity of periaqueductal gray subregions in episodic migraine. *J Headache Pain*. (2017) 18:36. doi: 10.1186/s10194-017-0747-9
- Guaschino E, Ghiotto N, Tassorelli C, Bitetto V, Nappi G, Moglia A, et al. P043. Hyperechogenicity of the periaqueductal gray in chronic migraine and episodic migraine as a potential marker of progressive dysfunction: preliminary results with transcranial sonography. *J Headache Pain.* (2015) 16(Suppl. 1):A61. doi: 10.1186/1129-2377-16-S1-A61
- Chen Z, Chen X, Liu M, Liu S, Ma L, Yu S. Nonspecific periaqueductal gray lesions on T2WI in episodic migraine. J Headache Pain. (2016) 17:101. doi: 10.1186/s10194-016-0695-9
- Ito K, Kudo M, Sasaki M, Saito A, Yamashita F, Harada T, et al. Detection of changes in the periaqueductal gray matter of patients with episodic migraine using quantitative diffusion kurtosis imaging: preliminary findings. *Neuroradiology*. (2016) 58:115–20. doi: 10.1007/s00234-015-1603-8
- Vecchia D, Pietrobon D. Migraine: a disorder of brain excitatory-inhibitory balance? *Trends Neurosci.* (2012) 35:507–20. doi: 10.1016/j.tins.2012.04.007
- Wang Y, Wang XS. Migraine-like headache from an infarction in the periaqueductal gray area of the midbrain. *Pain Med.* (2013) 14:948– 9. doi: 10.1111/pme.12096

- Haas DC, Kent PF, Friedman DI. Headache caused by a single lesion of multiple sclerosis in the periaqueductal gray area. *Headache*. (1993) 33:452– 5. doi: 10.1111/j.1526-4610.1993.hed3308452.x
- Fragoso YD, Brooks JB. Two cases of lesions in brainstem in multiple sclerosis and refractory migraine. *Headache*. (2007) 47:852–4. doi: 10.1111/j.1526-4610.2007.00823.x
- Zhu J, Li JL, Liu YM, Han YY, Wang Y. Unilateral headache with visual aura from a Wallenberg syndrome. CNS Neurosci Ther. (2012) 18:598– 600. doi: 10.1111/j.1755-5949.2012.00330.x
- Schulte LH, May A. Of generators, networks and migraine attacks. Curr Opin Neurol. (2017) 30:241–5. doi: 10.1097/WCO.00000000000441
- 24. de Freitas RL, de Oliveira RC, de Oliveira R, Paschoalin-Maurin T, de Aguiar Correa FM, Coimbra NC. The role of dorsomedial and ventrolateral columns of the periaqueductal gray matter and in situ 5-HT(2)A and 5-HT(2)C serotonergic receptors in post-ictal antinociception. *Synapse*. (2014) 68:16–30. doi: 10.1002/syn.21697
- de Freitas RL, Medeiros P, Khan AU, Coimbra NC: micro1 -Opioid receptors in the dorsomedial and ventrolateral columns of the periaqueductal grey matter are critical for the enhancement of post-ictal antinociception. *Synapse*. (2016) 70:519–30. doi: 10.1002/syn.21926
- Wang L, Shen J, Cai XT, Tao WW, Wan YD, Li DL, et al. Ventrolateral periaqueductal gray matter neurochemical lesion facilitates epileptogenesis and enhances pain sensitivity in epileptic rats. *Neuroscience*. (2019) 411:105– 18. doi: 10.1016/j.neuroscience.2019.05.027
- Mena-Segovia J, Bolam JP. Rethinking the pedunculopontine nucleus: from cellular organization to function. *Neuron.* (2017) 94:7–18. doi: 10.1016/j.neuron.2017.02.027
- de Oliveira RC, de Oliveira R, Falconi-Sobrinho LL, Biagioni AF, Almada RC, Dos Anjos-Garcia T, et al. Neurotoxic lesions of the pedunculopontine tegmental nucleus impair the elaboration of postictal antinociception. *Physiol Behav.* (2018) 194:162–9. doi: 10.1016/j.physbeh.2018.05.011
- Mazzei-Silva EC, de Oliveira RC, dos Anjos Garcia T, Falconi-Sobrinho LL, Almada RC, Coimbra NC. Intrinsic connections within the pedunculopontine tegmental nucleus are critical to the elaboration of post-ictal antinociception. *Synapse.* (2014) 68:369–77. doi: 10.1002/syn.21749
- Jaseja H. Pedunculopontine nucleus stimulation: a novel adjunctive therapeutic approach in intractable epilepsy. *Epilepsy Behav.* (2013) 27:279. doi: 10.1016/j.yebeh.2012.12.005
- Liu TT, Feng J, Bu HL, Liu C, Guan XH, Xiang HB. Stimulation for the compact parts of pedunculopontine nucleus: an available therapeutic approach in intractable epilepsy. *Epilepsy Behav.* (2013) 29:252–53. doi: 10.1016/j.yebeh.2013.06.033
- 32. Fisher RS, Cross JH, French JA, Higurashi N, Hirsch E, Jansen FE, et al. Operational classification of seizure types by the international league against epilepsy: position paper of the ilae commission for classification and terminology. *Epilepsia*. (2017) 58:522–30. doi: 10.1111/epi.13670
- Fisher RS, Cross JH, D'Souza C, French JA, Haut SR, Higurashi N, et al. Instruction manual for the ILAE 2017 operational classification of seizure types. *Epilepsia*. (2017) 58:531–42. doi: 10.1111/epi.13671
- Scheffer IE, Berkovic S, Capovilla G, Connolly MB, French J, Guilhoto L, et al. ILAE classification of the epilepsies: position paper of the ILAE commission for classification and terminology. *Epilepsia*. (2017) 58:512– 21. doi: 10.1111/epi.13709
- Hamilton M. A rating scale for depression. J Neurol Neurosurg Psychiatry. (1960) 23:56–62. doi: 10.1136/jnnp.23.1.56
- Guerrero-Berroa E, Luo XD, Schmeidler J, Rapp MA, Dahlman K, Grossman HT, et al. The MMSE orientation for time domain is a strong predictor of subsequent cognitive decline in the elderly. *Int J Geriatr Psych.* (2009) 24:1429–37. doi: 10.1002/gps.2282
- Gallagher EJ, Liebman M, Bijur PE. Prospective validation of clinically important changes in pain severity measured on a visual analog scale. *Ann Emerg Med.* (2001) 38:633–8. doi: 10.1067/mem.2001.118863
- Chao-Gan Y, Yu-Feng Z. DPARSF: a MATLAB toolbox for "pipeline" data analysis of resting-state fMRI. Front Syst Neurosci. (2010) 4:13. doi: 10.3389/fnsys.2010.00013
- Song XW, Dong ZY, Long XY, Li SF, Zuo XN, Zhu CZ, et al. REST: a toolkit for resting-state functional magnetic resonance imaging data processing. *PLoS* ONE. (2011) 6:e25031. doi: 10.1371/journal.pone.0025031

- Cacciola A, Bertino S, Basile GA, Di Mauro D, Calamuneri A, Chillemi G, et al. Mapping the structural connectivity between the periaqueductal gray and the cerebellum in humans. *Brain Struct Funct.* (2019) 224:2153–65. doi: 10.1007/s00429-019-01893-x
- Keuken MC, Forstmann BU. A probabilistic atlas of the basal ganglia using 7 T MRI. Data Brief. (2015) 4:577–82. doi: 10.1016/j.dib.201 5.07.028
- 42. Janzen J, van 't Ent D, Lemstra AW, Berendse HW, Barkhof F, Foncke EMJ. The pedunculopontine nucleus is related to visual hallucinations in Parkinson's disease: preliminary results of a voxel-based morphometry study. *J Neurol.* (2012) 259:147–54. doi: 10.1007/s00415-011-6149-z
- Cho KH, Cho YJ, Lee BI, Heo K. Atrophy of the pedunculopontine nucleus region in patients with sleep-predominant seizures: a voxel-based morphometry study. *Epilepsia*. (2016) 57:e151–4. doi: 10.1111/epi.13431
- 44. Ricci L, Assenza G, Pulitano P, Simonelli V, Vollero L, Lanzone J, et al. Measuring the effects of first antiepileptic medication in temporal lobe epilepsy: predictive value of quantitative-EEG analysis. *Clin Neurophysiol.* (2021) 132:25–35. doi: 10.1016/j.clinph.2020.10.020
- Yamada S, Kawata M. Identification of neural cells activated by mating stimulus in the periaqueductal gray in female rats. *Front Neurosci.* (2014) 8:421. doi: 10.3389/fnins.2014.00421
- Reyes BAS, Zhang XY, Dufourt EC, Bhatnagar S, Valentino RJ, Van Bockstaele EJ. Neurochemically distinct circuitry regulates locus coeruleus activity during female social stress depending on coping style. *Brain Struct Funct.* (2019) 224:1429–46. doi: 10.1007/s00429-019-01837-5
- 47. Menant O, Prima MC, Morisse M, Cornilleau F, Moussu C, Gautier A, et al. First evidence of neuronal connections between specific parts of the periaqueductal gray (PAG) and the rest of the brain in sheep: placing the sheep PAG in the circuit of emotion. *Brain Struct. Funct.* (2018) 223:3297–316. doi: 10.1007/s00429-018-1689-y
- Wang QQ, Zhu LJ, Wang XH, Zuo J, He HY, Tian MM, et al. Chronic trigeminal nerve stimulation protects against seizures, cognitive impairments, hippocampal apoptosis, and inflammatory responses in epileptic rats. J Mol Neurosci. (2016) 59:78–89. doi: 10.1007/s12031-016-0736-5
- Li LY, Li JL, Zhang HM, Yang WM, Wang K, Fang Y, et al. TGFbeta1 treatment reduces hippocampal damage, spontaneous recurrent seizures, and learning memory deficits in pilocarpine-treated rats. J Mol Neurosci. (2013) 50:109–23. doi: 10.1007/s12031-012-9879-1
- Racine RJ. Modification of seizure activity by electrical stimulation. II. Motor seizure. *Electroencephalogr Clin Neurophysiol.* (1972) 32:281– 94. doi: 10.1016/0013-4694(72)90177-0
- Mai J, Majtanik M, Paxinos G, Paxinos G, Watson C, Calabrese E, et al. *The Rat Brain in Stereotaxic Coordinates. Rat Brain Stereotaxic Coordinates.* (2008) 3:6. doi: 10.1016/0165-0270(80)90021-7
- 52. Paxinos G, Watson C. *The Rat Brain in Stereotaxic Coordinates. 6th ed.* San Diego, CA: Academic Press (2007).
- Wang XQ, Lang SY, He MW, Zhang X, Zhu F, Dai W, et al. High prevalence of headaches in patients with epilepsy. J Headache Pain. (2014) 15:70. doi: 10.1186/1129-2377-15-70
- 54. Winawer MR, Connors R. Evidence for a shared genetic susceptibility to migraine and epilepsy. *Epilepsia*. (2013) 54:288–95. doi: 10.1111/epi.12072
- Napadow V, Sclocco R, Henderson LA. Brainstem neuroimaging of nociception and pain circuitries. *Pain Rep.* (2019) 4:e745. doi: 10.1097/PR9.00000000000745
- 56. Jia Z, Tang W, Zhao D, Yu S. Disrupted functional connectivity between the periaqueductal gray and other brain regions in a rat model of recurrent headache. *Sci Rep.* (2017) 7:3960. doi: 10.1038/s41598-017-04060-6
- 57. Cong F, Wang JW, Wang B, Yang ZY, An J, Zuo ZT, et al. Direct localisation of the human pedunculopontine nucleus using MRI: a coordinate and fibre-tracking study. *Eur Radiol.* (2018) 28:3882–92. doi: 10.1007/s00330-017-5299-5
- Genaro K, Prado WA. Neural correlates of the antinociceptive effects of stimulating the anterior pretectal nucleus in rats. J Pain. (2016) 17:1156– 63. doi: 10.1016/j.jpain.2016.07.002
- 59. Kommajosyula SP, Faingold CL. Neural activity in the periaqueductal gray and other specific subcortical structures is enhanced when a selective serotonin reuptake inhibitor selectively prevents seizure-induced sudden death in the

DBA/1 mouse model of sudden unexpected death in epilepsy. *Epilepsia.* (2019) 60:1221–33. doi: 10.1111/epi.14759

- Andrews JP, Yue Z, Ryu JH, Neske G, McCormick DA, Blumenfeld H. Mechanisms of decreased cholinergic arousal in focal seizures: *in vivo* wholecell recordings from the pedunculopontine tegmental nucleus. *Exp. Neurol.* (2019) 314:74–81. doi: 10.1016/j.expneurol.2018.11.008
- Nolte MW, Loscher W, Gernert M. Pedunculopontine neurons are involved in network changes in the kindling model of temporal lobe epilepsy. *Neurobiol. Dis.* (2006) 23:206–18. doi: 10.1016/j.nbd.2006.03.003
- Jaseja H. Deep-brain stimulation in intractable epilepsy: pedunculopontine nucleus versus thalamic nuclei: a perspective. World Neurosurg. (2014) 82:e568-9. doi: 10.1016/j.wneu.2014.05.021
- van der Knaap LJ, van der Ham IJ. How does the corpus callosum mediate interhemispheric transfer? A review. *Behav Brain Res.* (2011) 223:211– 21. doi: 10.1016/j.bbr.2011.04.018
- Chen PC, Messina SA, Castillo E, Baumgartner J, Seo JH, Skinner H, et al. Altered integrity of corpus callosum in generalized epilepsy in relation to seizure lateralization after corpus callosotomy. *Neurosurg Focus*. (2020) 48:E15. doi: 10.3171/2020.1.FOCUS19791
- Kim H, Piao Z, Liu P, Bingaman W, Diehl B. Secondary white matter degeneration of the corpus callosum in patients with intractable temporal lobe epilepsy: a diffusion tensor imaging study. *Epilepsy Res.* (2008) 81:136– 42. doi: 10.1016/j.eplepsyres.2008.05.005
- 66. Uribe-San-Martin R, Ciampi E, Di Giacomo R, Vasquez M, Carcamo C, Godoy J, et al. Corpus callosum atrophy and post-surgical seizures in temporal lobe epilepsy associated with hippocampal sclerosis. *Epilepsy Res.* (2018) 142:29–35. doi: 10.1016/j.eplepsyres.2018.03.001
- Ashraf-Ganjouei A, Rahmani F, Aarabi MH, Nazem-Zadeh HSMMR, Davoodi-Bojd E, Soltanian-Zadeh H. White matter correlates of disease duration in patients with temporal lobe epilepsy: updated review of literature. *Neurol Sci.* (2019) 40:1209–16. doi: 10.1007/s10072-019-03818-2
- Liu J, Zhu J, Yuan F, Zhang X, Zhang Q. Abnormal brain white matter in patients with right trigeminal neuralgia: a diffusion tensor imaging study. J Headache Pain. (2018) 19:46. doi: 10.1186/s10194-018-0871-1
- Coppola G, Di Renzo A, Tinelli E, Petolicchio B, Di Lorenzo C, Parisi V, et al. Patients with chronic migraine without history of medication overuse are characterized by a peculiar white matter fiber bundle profile. *J Headache Pain*. (2020) 21:92. doi: 10.1186/s10194-020-01159-6
- Dun W, Yang J, Yang L, Ma S, Guo C, Zhang X, et al. Abnormal white matter integrity during pain-free periovulation is associated with pain intensity in primary dysmenorrhea. *Brain Imaging Behav.* (2017) 11:1061– 70. doi: 10.1007/s11682-016-9582-x

- Lewis R, Ruiz A, Monteith T. Reversible lesion of the corpus callosum in a patient with migraine with aura: a case study. *Headache*. (2020) 60:791– 2. doi: 10.1111/head.13768
- 72. Lin FY, Yang CY. Reversible splenial lesion of the corpus callosum in migraine with aura. *Neurologist.* (2011) 17:157–9. doi: 10.1097/NRL.0b013e31821733c2
- Samanta D. Transient lesion in the splenium of the corpus callosum in status migrainosus. Acta Neurol Belgica. (2014) 115:397–8. doi: 10.1007/s13760-014-0355-2
- 74. Li XL, Fang YN, Gao QC, Lin EJ, Hu SH, Ren L, et al. A diffusion tensor magnetic resonance imaging study of corpus callosum from adult patients with migraine complicated with depressive/anxious disorder. *Headache*. (2011) 51:237–45. doi: 10.1111/j.1526-4610.2010.01774.x
- Tovote P, Esposito MS, Botta P, Chaudun F, Fadok JP, Markovic M, et al. Midbrain circuits for defensive behaviour. *Nature*. (2016) 534:206– 12. doi: 10.1038/nature17996
- David AS, Wacharasindhu A, Lishman WA. Severe psychiatric disturbance and abnormalities of the corpus callosum: review and case series. J Neurol Neurosurg Psychiatry. (1993) 56:85–93. doi: 10.1136/jnnp.56.1.85
- Della Valle R, Mohammadmirzaei N, Knox D. Single prolonged stress alters neural activation in the periacqueductal gray and midline thalamic nuclei during emotional learning and memory. *Learn Memory.* (2019) 26:1– 9. doi: 10.1101/lm.050310.119
- Pang XM, Liang XL, Zhou X, Liu JP, Zhang Z, Zheng JO. Alterations in intra- and internetwork functional connectivity associated with levetiracetam treatment in temporal lobe epilepsy. *Neurol Sci.* (2020) 41:2165–74. doi: 10.1007/s10072-020-04322-8
- Hermans K, Ossenblok P, van Houdt P, Geerts L, Verdaasdonk R, Boon P, et al. Network analysis of EEG related functional MRI changes due to medication withdrawal in focal epilepsy. *NeuroImage*. (2015) 8:560– 71. doi: 10.1016/j.nicl.2015.06.002

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Wang, Cai, Zu, Zhang, Deng and Wang. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.