Epileptic network of hypothalamic hamartoma: an EEG-fMRI study

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Abbreviations: HH, hypothalamic hamartoma; EEG-fMRI, EEG with functional MRI; GS, gelastic seizure; SISCOM, SPECT coregistered to MRI; DMN, default mode network

Highlights:

- We performed EEG-fMRI in eight HH patients with GS.
- EEG-fMRI revealed activation in or around the hypothalamus in 6/8 patients.
- Activation in subcortical tissues and deactivation including DMN were found.
- Subcortical network and DMN can be related each to GS and epileptic encephalopathy.
- EEG-fMRI enhances sensitivity in detecting the HH interface compared with SISCOM.

ABSTRACT

Objective: To investigate the brain networks involved in epileptogenesis/encephalopathy associated with hypothalamic hamartoma (HH) by EEG with functional MRI (EEG-fMRI), and evaluate its efficacy in locating the HH interface in comparison with subtraction ictal SPECT coregistered to MRI (SISCOM).

Methods: Eight HH patients underwent EEG-fMRI. All had gelastic seizures (GS) and 7 developed other seizure types. Using a general linear model, spike-related activation/deactivation was analyzed individually by applying a hemodynamic response function before, at, and after spike onset (time-shift model = $-8 \sim +4$ s). Group analysis was also performed. The sensitivity of EEG-fMRI in identifying the HH interface was compared with SISCOM in HH patients having unilateral hypothalamic attachment.

<u>Results:</u> EEG-fMRI revealed activation and/or deactivation in subcortical structures and neocortices in all patients. 6/8 patients showed activation in or around the hypothalamus with the HH interface with time-shift model before spike onset. Group analysis showed common activation in the ipsilateral hypothalamus, brainstem tegmentum, and contralateral cerebellum. Deactivation occurred in the default mode network (DMN) and bilateral hippocampi. Among 5 patients with unilateral hypothalamic attachment, activation in or around the ipsilateral hypothalamus was seen in 3 using EEG-fMRI, whereas hyperperfusion was seen in 1 by SISCOM.

Significance: Group analysis of this preliminary study may suggest that the commonly activated subcortical network is related to generation of GS and that frequent spikes lead to deactivation of the DMN and hippocampi, and eventually to a form of epileptic encephalopathy. Inter-individual variance in neocortex activation explains various seizure types among patients. EEG-fMRI enhances sensitivity in detecting the HH interface compared with SISCOM.

Keywords: Gelastic seizure; Epileptic encephalopathy; Subcortical epilepsy; EEG-fMRI; Ictal SPECT

1 1. INTRODUCTION

2 Hypothalamic hamartoma (HH) is a rare developmental malformation that has provided 3 important insight about epileptology (Berkovic et al., 1988; Mullatti et al., 2003; Striano et 4 al., 2012). HH is characterized by gelastic seizures (GS) which occur in almost all patients. 5 Such patients often develop other seizure types, i.e., partial and generalized seizures, and 6 cognitive/behavioral problems including memory deficits and mental retardation. Previous 7 studies showed that HH per se accounts for the generation of these seizures (Kahane et al., 8 2003; Kameyama et al., 2010; Kameyama et al., 2009; Kuzniecky, 2004; Munari et al., 1995; 9 Striano et al., 2012; Wethe et al., 2013). Therefore, the unique spectrum of symptoms in HH 10 has been regarded as the model of subcortical epilepsy and epileptic encephalopathy 11 (Kameyama et al., 2010; Striano et al., 2012). However, it remains unknown how the 12 epileptic activity propagates, and how cognitive/behavioral dysfunction develops. For 13 example, the networks associated with GS remain elusive, although the mammillo-thalamo-14 cingulate tract from HH or the pathway from the HH to the brainstem and cerebellum has 15 been postulated (Kahane et al., 2003; Kameyama et al., 2010). Therapeutically, stereotactic 16 radiofrequency thermocoagulation (SRT) has become one of the most useful surgical 17 interventions. SRT of the HH interface has yielded better outcomes for seizure freedom and 18 fewer surgical complications than a direct approach (Kameyama et al., 2010; Kameyama et 19 al., 2009). Subtraction ictal SPECT coregistered to MRI (SISCOM) has been used for 20 locating the HH interface (Kameyama et al., 2010), however, it is time-consuming and the 21 likelihood of detecting the HH interface individually is not high. 22 Here, we performed EEG with fMRI (EEG-fMRI) on eight patients with HH. EEG-fMRI 23 can detect blood-oxygen-level dependent (BOLD) changes that are related to interictal 24 discharges identified from scalp EEG (Lemieux et al., 2001; Warach et al., 1996). It was

25 reported to be clinically useful for localizing the epileptic focus and investigating epileptic

1 network even in the deep brain structures (Gotman and Pittau, 2011; Vulliemoz et al., 2010). 2 In addition, it possibly predicts postsurgical outcome non-invasively although its clinical 3 utility in comparison with other techniques, e.g. ictal SPECT, has been not determined 4 (Chaudhary et al., 2013). With these in mind, we expected EEG-fMRI to clarify the common brain networks associated with subcortical epileptogenesis/encephalopathy in HH patients. 5 6 Furthermore, we evaluated the clinical usefulness of EEG-fMRI in locating the HH interface 7 in comparison with SISCOM (Kameyama et al., 2010). We thought EEG-fMRI would be an 8 option for presurgical investigation, because it could provide us with the interictal epileptic 9 network that could complement the findings of SISCOM in a relatively less time-consuming 10 fashion.

11

12 **2. METHODS**

13 2.1 Patients

14 Subjects were eight patients (pts.) with HH (age 1-27 years) (Table 1) who were examined at 15 the Kyoto University Hospital from August 2011 to June 2013. Subjects included three 16 patients who had received surgical intervention for HH in the past (pt. 2 as partial resection; 17 pt. 6 as partial resection and SRT; pt. 7 as partial resection followed by infarction of the left 18 hemisphere and gamma-knife surgery) with persisting seizures. Hamartomas were classified 19 by the Delalande classification, which is based on the plane of insertion on the hypothalamus 20 to help choose the best surgical route (Type I: horizontal implantation plane, Type II: vertical 21 insertion plane and intraventricular location, Type III: combination of Types I and II. Type IV: 22 all giant hamartomas for which no specific surgical procedures can be recommended) 23 (Delalande and Fohlen, 2003). All the patients or parents of patients who were not capable of 24 consent gave written informed consent. The protocol was approved by the Ethics Committee 25 of our institute (IRB#E217).

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2 2.2 EEG-fMRI acquisition

3 EEG was obtained during fMRI, using a custom-made MR-compatible cap with 19 (for pt. 5) 4 or 23 (for other patients) Ag/AgCl scalp electrodes based on the 10-20 System, including 5 Fp1, Fp2, F7, F3, F4, F8, T7, C3, C4, T8, TP9, P7, P3, P4, P8, TP10, O1, and O2 referenced 6 to Cz (T1, T2, Fz, and Pz were added optionally) (EASYCAP, EASYCAP GmbH, 7 Herrsching, Germany). Electrocardiogram was also recorded. All the signals were transmitted 8 from an MR-compatible amplifier (BrainAmp MR plus, Brain Products GmbH, Munch, 9 Germany: sampled at 5 kHz) through an optic cable to a computer outside the MR scanner 10 room and stored on the computer. The EEG system was synchronized with the MRI scanner 11 clock. 12 A 3 Tesla MR scanner was used (Trio, Siemens, Erlangen, Germany). The MRI sequence 13 for the echo planar imaging (EPI), i.e., blood oxygenation-level dependent (BOLD) fMRI 14 (repetition time [TR] = 3,000 ms; echo time [TE] = 30 ms; flip angle $[FA] = 90^{\circ}$; field of 15 view [FOV] = 192×192 mm; voxel size = $3 \times 3 \times 3$ mm) was acquired with a one-channel 16 bird-cage head coil. The initial two scans were discarded to ascertain the steady-state of the 17 magnetization. Motion artifacts were minimized as much as possible by stabilizing patients' 18 heads with a pillow filled with foam microspheres. Special care was taken to ensure the 19 patients' comfort. Scans lasted 30–90 min in several runs per patient. T1-weighted, 20 magnetization-prepared, rapid acquisition with gradient echo (MPRAGE) images (TR =21 2,000 ms; TE = 4.38 ms; FA = 8° ; FOV = 176×192 mm; voxel size = $1 \times 1 \times 1$ mm) were 22 also acquired with an eight-channel phased-array head coil for coregistration of the fMRI 23 results. By using this coil we expected to acquire a clearer anatomical image than one-24 channel coil does.

In all patients, SpO₂ was monitored in preparation for hypoxemia by a pulse oximeter
under constant observation by a medical doctor in the scanner room. In order to perform the
scan safely with minimal motion artifacts, trichrofos, midazolam, or pentobarbital were used
to sedate patients, though pentobarbital was mainly used (Table 1).

5

6 2.3 EEG-fMRI pre-processing

7 First, MR and ballistocardiogram artifacts were removed in an offline manner according to a 8 previously established method (Allen et al., 2000; Allen et al., 1998). EEG was down-9 sampled to 250 Hz by BrainVision ANALYZER software (Brain Products GmbH, Munich, 10 Germany). In addition, the EEG signals were processed with 30-Hz low-pass and 0.53-Hz 11 high-pass filters using an in-house script in Matlab. The filtered EEG signals were inspected 12 by a certified electroencephalographer (R.M.), and the occurrence times of interictal 13 paroxysmal discharges were marked (See an example of identified interictal discharges in 14 Figure 1A). The interictal findings eligible for marking were chosen based on the criteria that 15 (1) their waveform and location corresponded with those in EEG that was taken clinically 16 before and also based on 10-20 system, and (2) when spikes were defined from multiple foci, 17 the most predominant and frequent population was chosen for the robust BOLD changes. 18 Functional data were pre-processed and analyzed using the FMRIB Software Library v5.0 19 (FSL, www.fmrib.ox.ac.jk/fsl). The echo planar imaging (EPI) data underwent motion 20 correction, and were unwarped according to field-map data by FSL, and smoothed with a 21 Gaussian kernel with a full-width at half-maximum of 5 mm, and high-pass filtered (cutoff: 22 100 s).

23

24 2.4 Individual EEG-fMRI analysis

1 fMRI data sets were analyzed based on an event-related design using a general linear model 2 (FEAT program, part of FSL [FMRIB's Software Library, www.fmrib.ox.ac.uk/fsl]). For the 3 timing of spikes as events, a canonical hemodynamic response function (HRF) consisting of a 4 double gamma function was convolved (phase = 0 s; standard deviations = 3 s, mean lag = 65 s). Then, considering the possibility that the patterns of BOLD signal change differ in 6 different areas of the brain, multiple time-shift models in which HRF was applied before and 7 after spikes were also produced (t = -8, -6, -4, -2, +2, +4 s) (Figure 1B). Previous studies 8 revealed that this technique successfully revealed BOLD signal changes preceding focal- and 9 generalized spike discharges (Hawco et al., 2007; Jacobs et al., 2009; Moeller et al., 2008). 10 Each time-shift model related to spikes was used as the explanatory variable of interest, and 11 eight nuisance covariates (six head motion parameters, and signals from the CSF and white 12 matter [WM]) were regressed out as in a previous fMRI study (Fox et al., 2005). Basically, 13 the areas of the CSF and WM were extracted from a standard brain image in MNI standard 14 space (ICBM-152: "MNI152_T1_2mm.nii.gz") and non-linearly coregistered to the anatomy 15 image in each patient using FNIRT (www.fmrib.ox.ac.jk/fsl/fnirt). These processes were done 16 successfully with "fsl_anat" script in almost all cases. After binarization (0 or 1) of the CSF and WM voxels, the images were coregistered to EPI space. In this coregistration process, 17 18 each voxel has one probabilistic value (0–1) for each patient's CSF and WM. The time course 19 of the BOLD signal of the eigenvalues was calculated for each region where the value was \geq 20 0.95 after the voxels with artifacts around the air-tissue interface were excluded manually. 21 Each of these values was then used to regress out the effect of the CSF and WM, which 22 reportedly involve much noise caused by the cardiac and respiratory cycles (Dagli et al., 23 1999; Windischberger et al., 2002); thus, we expected that this process enabled us to 24 sensitively detect activation/deactivation of the small region near the ventricle, i.e., the 25 hypothalamus. Time-series analysis was carried out using FILM with local autocorrelation

1 correction (Woolrich et al., 2001). In the individual EPI space, Z-statistic images were 2 thresholded using clusters determined by Z > 2.3 at voxel level and a corrected cluster 3 significance level of p < 0.007 (0.05/7; Bonferroni correction was made because we used 4 seven time-series models individually) (Worsley, 2001). The images were then coregistered 5 to each anatomy image. The laterality concordance between EEG spikes and cortical 6 activation was blindly evaluated by an author (R.M.) who did not analyze fMRI data. The 7 laterality of spikes was defined by the side on which focal spikes occurred, or the side on 8 which spikes had larger amplitude (twice the height of the other side), in the case of a 9 bilateral or generalized spike. The laterality of the fMRI cortical activation was defined by 10 the side containing the most significant cluster of cortical positive BOLD responses with 11 early time-shift models (t = $-8 \sim 0$ s). The results of earlier time shift models ($-8 \sim 0$ s) were 12 used for comparison based on previous reports (Hawco et al., 2007; Jacobs et al., 2009; 13 Moeller et al., 2008) because later models could involve the zones epileptic activity finally 14 spread to other than the areas that generate spikes.

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16 2.5 Group analysis

17 Excluding one patient (pt. 5, 1-year-old female) in whom coregistration of the CSF and WM 18 was impossible, a random-effects group analysis was carried out using the FEAT program in 19 seven patients. Before analysis, MRI images were flipped in the left-right dimension in the 20 case of (1) patients with only the left side attachment or (2) those with left dominant 21 activation of the hypothalamus in the individual analysis (in case of bilateral attachment). Z-22 statistic images were thresholded using clusters determined by the criteria of Z > 2.3 and a 23 (corrected) cluster significance threshold of p = 0.007. Results were coregistered to the 24 common MR image in MNI standard space. Thus, all the pathologic hemispheres were shown 25 in the right hemisphere in the MNI space. The impact of HH on image normalization was

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5 2.6 SPECT and SISCOM

6 In seven patients excluding pt. 6, interictal and ictal SPECT were performed. These patients 7 were not included in our previous studies (Kameyama et al., 2010) at Nishi-Niigata Chuo 8 Hospital. A T1-weighted volumetric scan (1.5 Tesla, Shimadzu, Kyoto, Japan) was used for 9 coregistration of SPECT images. The imaging techniques and method of SISCOM have been reported elsewhere (Kameyama et al., 2010). ^{99m}Tc-ethyl cysteinate dimer (ECD) was 10 11 injected interictally and ictally. The ictal and interictal SPECT images, normalized according 12 to the global mean voxel counts, were subtracted to obtain ictal-to-interictal state difference 13 images. Areas with counts above two standard deviations in subtraction images were adopted 14 as significantly increased perfusion areas. Next, EEG-fMRI and SISCOM results were 15 compared in terms of their sensitivity to identify the HH interface.

16

17 3. RESULTS

18 **3.1 Individual analysis**

The areas that showed positive/negative BOLD activity (activation/deactivation) included cortical (e.g., neocortices including the insula and anterior cingulate cortices) and subcortical regions (e.g., the hypothalamus, thalamus, the caudate, brainstem, and cerebellum) to various degrees in all patients. In all six patients who showed cortical BOLD responses with early time-shift models (-8 ~ 0 s), the laterality of cortical activation was concordant with that of EEG spikes (Table 2). In 6/8 patients, activation was observed at the HH interface of the hypothalamus, or at its adjacent area, with early time-shift models (Table 2, and see Figure 2 1 for representative patients). Two patients (pt. 3 and 4), who did not show any activation in or 2 around the hypothalamus, had an intraventricular hamartoma (Delalande classification type 3 II) of smaller size compared with other patients (diameter < 10 mm). Thalamic BOLD 4 responses were observed in 6/8 patients with individual variance in activation/deactivation 5 patterns (activation; n = 2, deactivation: n = 2, mixed: n = 2). The summary of 6 activation/deactivation is shown in Table 3.

7

8 3.2 Group analysis

Group analysis revealed activation regions in the ipsilateral hypothalamus and brainstem
tegmentum with earlier time-shift models, and bilateral cerebellar activation (that was
dominant in the contralateral side) with later models (Figure 3). Deactivation areas included
the cunei, bilateral thalami, caudate nuclei, hippocampi, paracentral gyri, and parts of the socalled default mode network (DMN), i.e., the precunei, and inferior lateral parietal lobules.

15 3.3 Comparison between EEG-fMRI and SISCOM

Both recording and analyses were successfully performed in 8/8 patients with EEG-fMRI and
5/7 patients with SISCOM (no seizure was captured in two patients). The laterality of EEGfMRI (see above) was concordant to that of EEG spikes in 6/8 patients, while that of
SISCOM (hyperperfusion) in 2/7 patients. Both EEG-fMRI and SISCOM were examined in
five patients with unilateral HH attachment. The sensitivity to detect the activation area of the
ipsilateral hypothalamus (i.e., HH interface) or its adjacent area was present in 3/5 patients
with EEG-fMRI and 1/5 patients with SISCOM (Table 2).

24 4. DISCUSSION

25 4.1 Strength of this study

1 We applied EEG-fMRI to patients with HH and revealed that its epileptic network comprised 2 both neocortices and subcortical structures. Spike-related BOLD responses were observed 3 interictally in all patients in various regions either with activation (positive BOLD) or 4 deactivation (negative BOLD). In 6/8 patients, the hypothalamus with the HH interface or its 5 adjacent area showed activation with a time-shift model before spike onset. Group analysis 6 showed activation in the ipsilateral hypothalamus, brainstem tegmentum, and contralateral 7 cerebellum. Deactivation was observed in the DMN and hippocampi. Among 5 patients with 8 unilateral hypothalamic attachment, activation was observed at and around the attachment 9 (HH interface) in 3/5 patients with EEG-fMRI whereas hyperperfusion was detected in 1/5 10 patients by SISCOM.

11 Recently, several lines of evidence suggested the utility of the time-shift models before 12 spike onset for locating areas of cortical activation concordant with focal EEG spikes in 13 partial epilepsy (Bagshaw et al., 2004; Jacobs et al., 2007; Jacobs et al., 2009). Prespike 14 models were also reported to be useful for EEG-fMRI analysis of generalized epilepsy 15 (Moeller et al., 2008). In line with these studies, we successfully detected lateralized cortical 16 activation and hypothalamic activation using the prespike models. What generates the 17 prespike BOLD responses is not exactly known, but some mechanisms are suggested: the 18 different temporal profile of the hemodynamic response function for epileptic activity (Kang 19 et al., 2003), or the metabolic process of neuronal or non-neuronal origin involving glial 20 structures that precedes the epileptic spikes (Pittau et al., 2011).

Some evidence shows the relationship between HH and various types of seizures including
GS. One stereo-EEG study of a patient with HH revealed that GS was well correlated to ictal
discharges in the HH (Munari et al., 1995). Another study showed that direct stimulation of
HH through depth electrodes evoked GS (Kahane et al., 2003). Other studies revealed that
ictal SPECT showed hyperperfusion in HH (Kuzniecky et al., 1997) or its interface to the

hypothalamus (Kameyama et al., 2010). Finally, it has also been shown that GS, focal and
 generalized seizures, and associated cognitive/behavioral symptoms can improve after
 appropriate surgery in HH (Kameyama et al., 2010; Kameyama et al., 2009; Striano et al.,
 2012; Wethe et al., 2013).

5 Variable activation/deactivation patterns among individual patients most likely reflect 6 various seizure types other than GS, while the other clinical profiles, such as the size of the 7 HH and use of anti-epileptic drugs might be also responsible. Despite this variability, group 8 analysis showed activation in subcortical structures such as the hypothalamus and brainstem, 9 especially with early time-shift models, and the cerebellum with later time-shift models. Our 10 EEG-fMRI analysis based on interictal findings of HH emphasize the existence of a common 11 epileptic network often involved in patients having HH with GS, and reinforce the 12 importance of the subcortical epileptic network that Kameyama and colleagues addressed in a 13 previous SISCOM study (Kameyama et al., 2010). Ictal SPECT performed during GS 14 showed, at a group level, involvement of the ipsilateral hypothalamus, mediodorsal (MD) 15 nucleus of the thalamus and putamen, bilateral pontine tegmentum, and contralateral inferior 16 semilunar lobule of the cerebellum (Kameyama et al., 2010). Based on lesion studies (Parvizi 17 et al., 2001), it is suggested that the execution of laughter is automatically regulated by the 18 cerebropontocerebellar pathway, which decussates before entering the middle cerebellar 19 peduncle. Additionally, stimulation of a small fiber bundle within the dorsal midbrain 20 tegmentum reportedly induced a laughing-like reaction in primates (Weinstein and bender, 21 1943). The findings of previous EEG-fMRI studies with a smaller number of patients having 22 HH (Kokkinos et al., 2012; Leal et al., 2009) are consistent with the existence of a network 23 including the mammillo-thalamo-cingulate tract (Kahane et al., 2003). Some patients in our 24 study showed involvement of the hippocampi and thalamus (see Figure 2). The mammillo-25 thalamo-cingulate tract could be another epileptic network involved in HH. The present EEG-

1 fMRI study showed thalamic deactivation at a group level, with large inter-individual 2 variability in terms of BOLD response patterns. This observation is consistent with some 3 EEG-fMRI studies of generalized epilepsy, which have shown either activation or 4 deactivation of the thalamus, either ictally or interictally (Aghakhani et al., 2004; Carney et 5 al., 2010; Gotman et al., 2005; Hamandi et al., 2006; Moeller et al., 2008; Siniatchkin et al., 6 2011). Multiple variables is likely to determine the strength that the HH affect the thalamus 7 via its epileptic activity, including the patients' age, the size of the HH and the location of its 8 attachment. They may explain the inter-individual variability, but we need more patients for 9 analysis.

10 The regions of the DMN, which show increased brain activity at rest (Raichle et al., 2001), 11 showed activation/deactivation in our study. It has been reported these areas are involved in 12 epileptic discharges in frontal- and temporal lobe epilepsy, posterior quadrant epilepsy and 13 idiopathic generalized epilepsy (Fahoum et al., 2012; Gotman et al., 2005; Laufs et al., 2007). 14 The network was evaluated in terms of the impairment of attention (Gotman et al., 2005) or cognitive dysfunction when damaged in neurodegenerative disorders such as Alzheimer's 15 16 disease (Johnson et al., 1998). Deactivation was also found in the bilateral hippocampi in the present study, which is thought to be related to memory encoding, consolidation, and retrieval 17 18 (Carr et al., 2011). If frequent epileptic discharges from HH involve both the DMN and 19 hippocampi, the core regions of the memory network from early childhood, there might be an 20 interruption of the normal cognitive development process, and eventually a part of epileptic 21 encephalopathy.

Ictal SPECT generally has a diagnostic value in locating the seizure onset zone among
 neuroimaging techniques (Spencer, 1994); however, in our pilot study, interictal EEG-fMRI
 of HH had no less sensitivity than SISCOM in detecting regional activation in and around the
 hypothalamus or lateralized cortical activation concordant with EEG spikes. This is partly

due to the higher spatiotemporal resolution of fMRI, which is sensitive enough to detect the
small and transient HH-related epileptic brain activity. Furthermore, EEG-fMRI requires a
short recording time of ~2 hours including preparation, as opposed to SISCOM, which
requires seizure recording that usually takes a long time for patients and medical staffs.
Therefore, we suggest that EEG-fMRI is a more practical and useful tool to identify the
laterality of HH attachment for surgical treatment by SRT.

7

8 4.2 Limitations

9 We should also be cautious in interpretation of the present data as follows. First, the variety 10 of the age, drugs that were used for sedation, and the status of pre-/post-operation of the 11 patients are all confounding factors that may influence the results in this preliminary study as 12 mentioned earlier. The sample size of the patients is small in the present study, and future 13 studies with a larger cohort warrant the validity of the findings in this small pilot study. Some 14 patients might have slept during recording and it would affect results in terms of BOLD 15 response through metabolic change. However, event-related design that our study used is 16 immune to sleep-related change compared to block design because individual spikes are 17 extremely transient than sleep. Second, the difference in activation patterns in terms of the 18 location of HH attachment to the hypothalamus could not be evaluated due to the small 19 number of patients. Third, we arbitrarily analyzed the most frequent interictal discharges in 20 each patient. However, these discharges could contain epileptic activities that have no direct 21 relationship with HH, or activities from secondary epileptic foci. Last, EEG-fMRI has an 22 intrinsic methodological limitation. Spikes must be large enough in amplitude to be detected 23 in the filtered EEG, and frequent enough to reveal the areas of statistical significance in fMRI 24 analysis. This contrasts with ictal SPECT that does not require EEG for analysis. A large

3

4 5. Conclusion

In this study, we showed that EEG-fMRI in patients with HH detected brain areas possibly
involved in epileptogenesis/encephalopathy, and that it had comparable sensitivity with
SISCOM in detecting the HH interface. Future studies using EEG-fMRI would further
expand our understanding about HH and its epileptogenesis, and corroborate the clinical
usefulness of this technique.

10

11 **Conflicts of Interest:** none.

12

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- 26

1 FIGURE LEGENDS

2 Figure 1.

(A) An example of identified interictal paroxysmal discharges from pt. 2 is displayed (arrow).
(B) Time-shift model of hemodynamic response function (HRF). For the timing of spikes as
events (such as the spike shown in A), the canonical HRF consisting of a double gamma
function was convolved (yellow trace). The model was shifted in steps of 2 s, creating 6
additional models to account for the differences in the time-course of blood oxygenationlevel dependent (BOLD) signal changes associated with different brain regions.

9

10 **Figure 2.**

11 Representative findings of EEG-fMRI analysis and SISCOM. In two patients, representative 12 slices with time-shift models are shown. Arrowheads show the hamartoma and attachment 13 side in each patient. The voxels with significant activation [positive blood oxygenation-level 14 dependent (BOLD)] /deactivation (negative BOLD) by cluster-level statistics (p < 0.007 after 15 Bonferroni correction) are overlaid on each anatomical image. (A) In pt. 2 with left side 16 attachment to the hypothalamic hamartoma (HH), activation was observed in the HH 17 interface of the left hypothalamus and left lateral frontal cortex (t = -2). The result was 18 comparable with that of the subtraction ictal SPECT coregistered to MRI (SISCOM). Other 19 activations included the bilateral cerebellum (contralateral side dominant), and lateral 20 temporal lobes. Deactivations included the bilateral cunei, bilateral precunei, bilateral 21 hippocampi, and bilateral thalami. (B) In pt. 8 with bilateral attachments to the HH, 22 activations included the HH interface of the left hypothalamus, and left hippocampus and 23 amygdala. SISCOM did not yield significant hyperperfusion at and around HH. Deactivation 24 included the precunei, medial frontal cortices, and bilateral thalami. The regions satisfying p 25 < 0.007 by cluster-level statistics are shown for the EEG-fMRI analysis.

1

2 Figure 3.

3 Group analysis of EEG-fMRI. Group analysis of seven patients revealed the common 4 epileptic network associated with epilepsy with HH. Before group analysis, MRI images were 5 flipped in the left-right dimension so that the right hemisphere shows the side of attachment 6 (unilateral attachment) or that of predominant hypothalamic activation (bilateral attachment). 7 The voxels with significant activation/deactivation by cluster-level statistics (p < 0.007 after 8 Bonferroni correction) are overlaid on the group mean anatomical image in the MNI standard 9 space. Activations included the ipsilateral hypothalamus, brainstem tegmentum, bilateral 10 precunei, and lateral parietal lobes, mainly with early time-shift models, and the contralateral 11 dominant bilateral cerebellum (contralateral dominant) with later time-shift models. 12 Deactivations included the ipsilateral cuneus, bilateral caudate nuclei, thalami, hippocampi, 13 precunei, and lateral parietal lobes.

Detiont		Hamartoma						
No., Age/gender	Attachment side	tachment Delalande side classification		Gelastic Sz	Other types of Sz	Complication	AED	
1.15/M	L	III	20	+	GTC	PP	VPA, CBZ	
2. 8/M	L	III	16	+	GTC, TC, CPS	PP, BD, MR	CZP, CLB, LTG	
3. 3/M	R	II	7	+	-	BD	VPA, LTG	
4. 27/M	R	II	8	+	GTC, SPS	-	CBZ	
5. 1/F	R	II	10	+	GTC	-	ZNS, LEV	
6. 19/F	Bilateral	Ι	13	+	AT	PP, MR	VPA, CBZ, GBP, TPM	
7. 13/M	Bilateral	III	30	+	TC, CPS	PP, BD, MR	CBZ, TPM, SL	
8. 23/F	Bilateral	Ι	16	+	GTC, CPS	PP, BD	VPA	

1 Table 1 Clinical profiles of patients

2 Abbreviations: Sz = seizure, GTC = generalized tonic clonic, CPS = complex partial Sz, TC = tonic Sz, SPS = simple partial Sz, AT = atonic Sz,

3 PP = precocious puberty, BD = behavioral disorder, MR = mental retardation.

4 AED = antiepileptic drug

5

			SISCOM						
Patient No., Age/Gender	Spike location	Sedation: pentobarbital (mg/kg)	Total recording time (min)	No. of analyzed spikes	Laterality concordance between EEG spike and cortical activation from t = -8 to 0 model	Activation in or around the hypothalamus which has the HH interface (The earliest model that showed activation)	Laterality concordance between EEG spike and cortical hyperperfusion (the cortex showing most significant hyperperfusion)	Hyperperfusion in or around the hypothalamus which has the HH interface	
1.15/M	L PQ	2.8	70	73	n.d.	✓ (-2)	No Sz recorded		
2. 8/M	L FC	2.5	56	41	✓ (L LF/ P)	✓ (-2)	✓ (LT, LP)	\checkmark	
3. 3/M	LFC (ST)	16.2	56	9	n.d.	n.d.	- (Not clear, R O)	n.d.	
4. 27/M	R AQ	-	90	17	✓ (R O)	n.d.	No Sz recorded		
5. 1/F	G (R > L)	2.6 (+trichlofos 0.5 mg/kg)	77	9	✓ (R MF)	✓ (-2)	✓ (Not clear, R insula)	n.d.	
6. 19/F	Bilateral F (L > R)	1.31 (+midazolam 0.65 mg/kg)	56	451	✓ (L LF)	✓ (-8)	Not done		
7.13/M	G(L > R)	1.9	30	434	✓ (L LF)	✓ (-8)	- (L G, R MF)	\checkmark	
8. 23/F	LAQ	-	56	39	✓ (L P)	✓ (-2)	- (Not clear, R FP)	n.d.	

 Table 2
 Activation (positive BOLD) revealed by EEG-fMRI and SISCOM results

2 Abbreviations: Sz = seizure. PQ = posterior quadrant, FC = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, ST = sharp transients, AQ = anterior quadrant, G = generalized, F = frontocentral, F = generalized, F = frontocentral, F = generalized, F = generalized,

3 frontal. MF = medial frontal, LF = lateral frontal, P = parietal, T = temporal, O = occipital, FP = frontal pole, "n.d." = no cortical BOLD activity

4 (EEG-fMRI) / no hyperperfusion (SISCOM) detected.

1

1 Table 3 Summary of Activation/Deactivation by EEG-fMRI

	Deep brain					Frontal-Parietal						
Patient No., Age/Gender	Hypo- thalamus	Thalamus	Caudate	Midbrain	Cerebellum	Central sulcus	Oper- culum	Lateral Frontal	Medial Frontal	Insula	Lateral parietal	Precuneus /posterior cingulate
1.15/M	A ^a	A ^a		A ^a	A ^c	D	D	D^{b}	D	D	D	D
2. 8/M	A ^b	A ^b /D	D	D^{c}	A ^c	A ^c	A ^c /D	A ^a , ^b /D	A ^b /D	A/D	A ^{a, b} ∕D	A^b/D
3. 3/M					D						D^{b}	D^b
4. 27/M									D		A ^b	A ^b
5. 1/F	A ^{a, b}	A ^{a, b}	A ^a	A ^a			D	D	A ^{a, b}		D^b	А
6. 19/F	А	A/D	D	А	A ^{a, c} /D	A/D	A^a/D	A ^{a, b} /D	A^a/D	A/D	A^a/D	A^a/D
7. 13/M	A^a/D	D	D	А	A ^{a, c} /D	A^a/D	A^a/D	A ^{a, b} /D	D	D	A^a/D	A^a/D
8. 23/F	A ^b	D				D	D		A ^b /D	A ^b /D	A ^a /D	D
Patient No.,	Temporal		Ossinital									
Age/Gender	Lateral	medial	Occipital									
1.15/M	D		D		Abbreviation; A	A: activation,	D: deactiv	ation.				
2. 8/M	A ^c /D	$\mathbf{D}^{\mathbf{b}}$	D									
3. 3/M			D^b		a: the earliest time-shift model activation,							
4. 27/M			A ^{a, b}		1 • • / •1 .							
5. 1/F	D	A ^b			b: 1ps1/un1latera	l dominant,						
6. 19/F	A ^a /D	D	A/D		c: contralateral	dominant						
7. 13/M	A ^a /D	A^a/D	A^a/D		c. contratatoral	dominunt.						
8. 23/F	A^b	A^b	A ^b /D									











Fig. 3			Activat	tion (positive	BOLD)	2.3	4.5
Time-shift			Deactivati	on (negative	BOLD)	2.3	4.5
model t =	-8	-6	-4	-2	0	+2	+4
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