Abnormal Cortical Thickness in Epilepsy Compared to Simple Febrile Seizures in Children: Voxel-Based Morphometric Study

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Abstract

Background and Aim: Epilepsy is currently considered a common neurological disease associated with excessive neuronal damage. Recent functional MRI studies indicate that frontal cortex activity occurs before thalamic involvement in epileptic discharges, implying that the frontal cortex may play a role in childhood seizures. Therefore, this study aimed to investigate differences in gray matter (GM) structural alteration between epilepsy and simple febrile seizures (simple FS).

Material and Methods: In this retrospectively, we included 34 children with epilepsy, 27 children with simple FS, and 29 controls aged 6–60 months with magnetic resonance...
imaging (MRI). Voxel-Based morphometry (VBM) was used to compare GM voxel among the above groups. T13D-weighted images were used to segment GM through a custom-designed automated method.

Results: GM cortical reduction was significantly detected in the right precentral gyrus, right middle frontal gyrus, right frontal gyrus (opercular), right frontal gyrus (medial), bilateral orbitofrontal cortex (medial), and bilateral anterior cingulate gyrus in the epilepsy group (P < 0.05). There were no deep nuclei volume changes in the epilepsy group than in control (P > 0.05). Compared to controls, there was no significant change in GM deep nuclei or cortical thickness of children with simple FS (P > 0.05).

Conclusions: VBM is an effective method to differentiate epilepsy from simple FS. In the epilepsy group, the diseases' initiation started from cortical neurons; deep nuclei were not involved. Simple FS cannot cause deep GM volume reduction or cortical thickness changes and is expected to have a good outcome.

Keywords: Epilepsy; Simple Febrile Seizure; Gray Matter; Voxel-Based Morphometry

1. Introduction

Epilepsy is one of the most common neurological diseases nowadays, affecting more than 70 million people of all ages around the world. It's mainly endemic in low and middle-income countries, with 2.4 million almost being newly diagnosed each year [1, 2]. Childhood epilepsy is associated with children who are vulnerable to seizures during their rapidly developing brain period. Epilepsy usually has different clinical presentations, according to a distinct pathophysiological mechanism [3, 4]. Childhood epileptic seizures accounted for 50% of all diagnoses, with the majority of cases marked by generalized tonic-clonic seizures and loss of consciousness similar to simple febrile seizures (simple FS) [5, 6]. This similar clinical presentation makes it difficult to differentiate between epilepsy and simple FS, particularly in early life seizures [7].

Simple FS are usually benign and self-limited, and they don't appear to cause long-term neurodevelopmental delayed [8]. On the other hand, simple FS children have a slightly higher risk of developing epilepsy than normal children (1% vs. 0.5%) [8]. Furthermore, simple FS represents 70% of all febrile seizures (FS) seen in the pediatrics emergency clinic [9]. The current brain damage, family history of seizures, genetic susceptibility, or neonatal complications such as low Apgar scores may develop epilepsy after FS [3]. Moreover, seizure episode also causes distress to parents and family members [8]. However, it is unknown if simple FS will increase the risk of seizure-induced brain damage or changes in gray matter (GM) thickness. GM maturation in early life is essential for the developing brain, and it is susceptible to a variety of etiologic influences [10, 11]. FS can cause a fragile brain state during the maturation process [8]. Previous research has shown that abnormal cortical GM changes in the brain are related to the etiological factors of childhood epilepsy [10, 11]. Although most epileptic patients have an FS history, it is unknown if this increases their brain damage susceptibility [12, 13]. While visual inspection of structural magnetic resonance imaging (MRI) in epileptic patients typically reveals a normal appearance, a more advanced analysis technique based on quantitative MRI evaluations
can improve sensitivity and enable a more detailed examination of brain structural abnormalities [14-17]. Many studies using voxel-based morphometry (VBM) analysis have discovered structural abnormalities in the thalamus and frontal lobe in epileptic patients [18, 19]. However, pathological studies showed no association between structural brain damage in epilepsy and clinical presentation [19]. There has been no previous study relating cortical GM alterations in epileptic patients compared to simple FS children and controls. We claim that structural abnormalities in epilepsy begin in the developing brain's cortical GM area without involving deep nuclei. Therefore, this study aimed to investigate differences in GM structural alteration between epilepsy and simple FS using quantitative VBM analysis.

2. Material and Methods
This retrospective study was approved by, local institutional review board in the Clinical Research Ethics Committee of the First Affiliated Hospital of Xi'an Jiaotong University. All possible risks of an MRI scan, such as excessive noise, were explained to the children's parents. The parents signed the written informed consent.

2.1 Clinical and demographic data of the participants
Children aged 6–60 months who underwent MRI examinations as part of the screening for brain disease at the Department of Radiology in Xi'an Jiaotong University First Affiliated Hospital were sequentially enrolled between June 2014 and December 2020. In the current study, participants including 34 newly diagnosed epileptic children without a history of antiepileptic drugs, 27 simple FS, and 29 controls were enrolled based on inclusions and exclusion criteria. The epileptic children who were admitted had to meet the following criteria: (i) primary epilepsy diagnosed according to the International League Against Epilepsy diagnostic criteria [5, 20]; (ii) newly diagnosed epilepsy without a history of antiepileptic medications; (iii) gestational age ≥ 37 weeks; (iv) no other neurological disorders, such as autism or attention deficit hyperactivity disorder; and (v) a two weeks delay between the MRI scan and the last seizure before the MRI scan. The following were used as exclusion criteria: (i) insufficient clinical details on the course following seizure onset and seizure duration; (ii) a history of intracranial infection or head trauma; and (iii) MRI anomalies, such as T2 FLAIR hyperintensity. Simple FS children were identified as those who met the following criteria: (i) were diagnosed with simple FS using the criteria of the American Academy of Pediatrics [9, 21]; (ii) had a gestational age ≥ 37 weeks; (iii) had no history of brain injury, head trauma, or central nervous system infections; and (iv) had a period less than two weeks between seizure onset and MRI scan. Exclusion criteria were as follows: (i) insufficient clinical information on the course of the seizure after onset; (ii) insufficient clinical information on the duration of the seizure; and (iii) MRI anomalies, such as hyperintensity on T2 fluid-attenuated inversion recovery (FLAIR) [22]. The children in the control group met the following criteria: (i) gestational age of ≥ 37 weeks; (ii) no history of epilepsy or other forms of seizures; and (iii) no MRI anomalies. Children whose images showed artifacts were not allowed to participate. Additionally, children with developmental disorders (such as facial palsy and tic disorders) and intracranial infection are also at risk. The course duration after seizures onset is the time interval is referred to time between seizure onset and MRI scan, to make sure all diseases children were scan in the same circumstances, the previous study in idiopathic generalized
epilepsy (IGE) found it can affect brain fluid circulation and Glymphatic system [23].

|                           | Epilepsy          | Simple FS        | Control          | $P$ value |
|----------------------------|-------------------|------------------|------------------|-----------|
|                           | (n=34)            | (n=27)           | (n=29)           | Epilepsy vs. | Epilepsy vs. | Simple FS vs. |
| Age (months)              | 29.98 ± 12.14     | 27.92 ± 12.39    | 26.75 ± 12.23    | 0.87       | 0.94        | 0.98          |
| Gender (male)             | 22(63.63%)        | 17(61.54%)       | 18(60.71%)       | 0.80       | 0.86        | 0.91          |
| GA (weeks)                | 38.21 ± 1.08      | 38.35 ± 1.23     | 38.32 ± 1.36     | 0.487      | 0.515       | 0.93          |
| Seizure duration (minutes)| 7.88 ± 7.62       | 4.54 ± 2.45      | NA               | <0.001**   | NA          | NA            |
| Course duration after seizures onset (days) | 5.42 ± 1.43 | 4.69 ± 1.38 | NA | 0.427 | NA | NA |

Note: mean ± standard deviation; NA, Not applicable

**Table 1:** Demographics and clinical data of the participants.

**Figure 1:** Statistical VBM reveals cortical GM reductions in the brains of epilepsy compared with control.
Figure 2: Statistical VBM demonstrates cortical GM atrophy in the brains of epilepsy compared with simple FS.

|                      | Epilepsy vs. control |                  | Epilepsy vs. Simple FS |
|----------------------|----------------------|------------------|------------------------|
|                      | Cluster number | Voxels number | P-value (minimum) | Cluster number | Voxels number | P-value (minimum) |
| Precentral gyrus right | 1          | 318          | 0.015*          | 1              | 342          | 0.017*          |
| Middle frontal gyrus right | 1        | 70           | 0.025*          | 1              | 31           | 0.032*          |
| Inferior frontal gyrus (opercular) right | 1      | 508          | 0.015*          | 1              | 519          | 0.017*          |
| Superior frontal gyrus (medial) right | 1    | 39           | 0.045*          | -              | -            | -               |
| Orbitofrontal cortex (medial) left | 1   | 456          | 0.015*          | 1              | 360          | 0.032*          |
| Orbitofrontal cortex (medial) right | 2   | 374          | 0.030*          | 1              | 49           | 0.038*          |
| Anterior cingulate gyrus left | 2  | 1008         | 0.015*          | 1              | 463          | 0.032*          |
| Anterior cingulate gyrus right | 2 | 499           | 0.040*          | -              | -            | -               |

P < 0.05.

Table 2: VBM revealed significant cortical GM atrophy in epilepsy compared to control and simple FS.
2.2 MRI data acquisition
At the First Affiliated Hospital of Xi’an Jiaotong University, all participants underwent MRI exams using the same 3.0-T scanner (Signa HDxt, GE Healthcare, Milwaukee, WI) with an 8-channel head coil. The single-shot echo-planar three-dimensional fast spoiled gradient-recalled echo T1-weighted imaging (T1WI) was performed by using the following parameters: Repetition time = 10.468 ms; Echo time = 4.764 ms; Inversion time: 400 ms; Field of view = 240 × 240 mm²; Acquisition matrix = 240 × 240; Slice thickness = 1 mm.

2.3 Image processing and statistical analysis
2.3.1 Voxel-based morphometry (VBM): Firstly, 3D T1WI images are segmented into GM, white matter (WM), and cerebrospinal fluid (CSF) regions using the Statistical Parametric Mapping software (SPM, version 12, https://www.fil.ion.ucl.ac.uk/spm). Secondly, a study-specific GM template is generated using a group-wise approach based on the control group’s images [24]. Thirdly, by using the registration, all the individual GM images are normalized to template space. Fourthly, all reported GM images are multiplied by the warp Jacobian and smoothed with a Gaussian kernel (sigma = 3mm). The statistical analysis of VBM was performed by using the general linear model. P values > 0.05 were considered statistical significance after correction of the family-wise error (FWE) rate and the threshold-free cluster enhancement (TFCE).

2.3.2 Statistical analysis: The intergroup was analyzed using a separate t-test, using substantially different demographic and clinical data between epilepsy, simple FS, and control groups. For the comparison of categorical variables, the chi-square (x²) method was used. The Non-parametric Mann-Whitney U test was used to evaluate the continuous variables with non-normal group distribution, identifying the statistical significance (P < 0.05). For statistical analysis, a commercial software package (SPSS version 21.0 IBM, Armonk, NY, USA) was used.

3. Results
There were 34 epileptic children and 27 simple FS children, and 29 control children in this section met selection requirements and consented to participate in the study. After monitoring seizures among these groups, there were no substantial differences in age, sex, gestational age, and scan time after seizures onset (Table 1).

3.1 Epilepsy with control
Relative to the control group, epileptic children displayed decreased cortical GM thickness on the right precentral gyrus, right middle frontal gyrus, right frontal gyrus (opercular), right frontal gyrus (medial), bilateral orbitofrontal cortex (medial), and bilateral anterior cingulate gyrus compared to control group (P < 0.05) (Figure 1, Table 2). However, no significant difference was observed in the deep GM nucleus volume, such as bilateral thalami, basal ganglions, hippocampus, or cerebellum.

3.2 Epilepsy with simple FS
Comparing epileptic children with a simple FS group, the findings showed a marked reduction in cortical GM thickness on the right precentral gyrus, right middle frontal gyrus, right inferior frontal gyrus (opercular), bilateral orbitofrontal cortex (medial), and left anterior cingulate gyrus relative to a simple group of FS (P < 0.05) (Figure 2, Table 2). There were significant differences in seizure durations between epilepsy and simple FS (P < 0.05). No
significant changes in volumes of bilateral thalami, basal ganglia, hippocampus, or cerebellum were seen.

3.3 Simple FS with control
No significant differences in cortical GM thickness or subcortical nuclei volumes were observed in children with simple FS than the control group (P > 0.05).

4. Discussion
This study attempted to explore the cortical GM's structural changes in children who had recently been diagnosed with epilepsy using the VBM approach that considers volumetric changes. The study's main findings are cortical GM atrophy in bilateral orbitofrontal cortex (medial), and bilateral anterior cingulate gyrus, while in the right precentral gyrus, right middle frontal gyrus, right frontal gyrus (opercular), right frontal gyrus (medial), focally damaged in epileptic children compared to control. However, in comparing to simple FS, no significant differences were found in the superior frontal gyrus (medial) right and anterior cingulate gyrus right in the epileptic group. Besides, there were no substantial changes in the volume of the thalamus, basal ganglia, and hippocampus in epilepsy or simple FS in all comparisons. Because our epilepsy group was heterogenous, our results showed some regions are bilaterally damage, and some are focal. These results may help in understanding the neuroanatomical modifications that contribute to epilepsy during brain development. These results indicated that newly diagnosed epileptic children have frontal lobe cortical GM atrophy, suggesting that epilepsy affects this brain region directly.

Furthermore, our children's initial lesion is cortical, with no thalamic or deep nuclei volume changes. Again, previous research using quantitative MRI failed to identify structural defects in the thalamus of patients with IGE, similar to our findings [14, 25]. The volumes of the thalamus in our study did not change substantially between the patient's and control groups, in contrast to a large number of animal and, to a lesser extent, clinical studies suggesting that the thalamus plays a significant role in this disease. These findings are consistent with the report of a previous volumetric study of IGE patients [25]. Consequently, anomalies in the thalamocortical network, which have been observed in animals [26] and humans[27], do not seem to be related to thalamic structural changes. Maybe, VBM cannot detect mild changes in thalamic volume; at the same time, MR spectroscopy (MRS) of the thalamus has shown defects in patients with IGE [27], but VBM did not [25]. As a consequence, the absence of structural changes does not rule out the possibility of functional abnormalities. Furthermore, our results indicate that structural reduction of the cortical cortex occurs outside of the thalamus. Moreover, the thalamus is not only essential but also the most impaired organ in people with epilepsy. According to animal and clinical studies, the thalamocortical circuitry is involved in seizure generalization and EEG discharge maintenance [28, 29]. Our study, standard VBM examination, revealed no significant GM volume alterations in the bilateral thalami, basal ganglions, hippocampus, or cerebellum.

The frontal cortical GM thickness was reduced in the epilepsy group compared to the simple FS and control groups. The current findings were consistent with the previous cortical reduction in generalized tonic-clonic (GTC) and juvenile myoclonic epilepsy (JMC), [18, 30, 31] suggesting that epileptic children may have brain changes in
the frontal lobe. Multisensory inputs converge in the cortical, which connects to subcortical regions. Furthermore, VBM studies in patients with juvenile myoclonic epilepsy (JME) have revealed increased GM in the superior frontal gyrus [32] and orbitofrontal regions [33]. In children with JME, fMRI showed frontal lobe hyperactivity in the prefrontal cortex (PFC) and superior frontal gyrus [34]. These differences may suggest that epilepsy children have minor anatomical defects in their developing brains. The second possibility may be due to the subjects' age group and the fact that their epilepsy was newly diagnosed and had never taken antiepileptic drugs before. In comparing epilepsy to simple FS, the study finds inconsistencies in the superior frontal gyrus (medial) and the right and anterior cingulate gyrus that we can’t even clarify. It's possible that the simple FS maybe is not benign. Simple FS children may develop neurological disorders such as cognitive decline, emotional memory loss, [35] or behavioral disturbances, [36] based on nuclei functions, and a recommended cohort follow-up study. Our study has a range of limitations. First, it was a retrospective cross-sectional analysis with a heterogynous epilepsy group. Second, we did not follow up with children who had simple FS or examine the risk factors for simple FS developing into epilepsy. Third, clinical evidence such as cytokine levels was not assessed in patients with simple FS and epilepsy. However, our results showed that simple FS does not cause GM abnormalities after onset, which may relieve families' concerns about their children with simple FS.

5. Conclusion
This VBM approach has a significant advantage in terms of distinguishing these two seizures. The frontal lobe cortex of children with epilepsy was substantially reduced without involving deep nuclei. There were no cortical thickness changes in simple FS, suggesting that the disease would have a positive outcome. Our results contribute to a better understanding of neuroimaging processes in the field, leading to new therapeutic approaches that consider seizure mechanisms.

Conflicts of Interest
The authors declared there are no financial conflicts of interest.

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