Brain correlates of spike and wave discharges in GLUT1 deficiency syndrome

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Keywords: GLUT1 deficiency syndrome; GLT1DS; GLT1; Generalized spike-and-wave discharges; Basal ganglia; Genetic generalized epilepsy; Absence seizure

1. Introduction

Glucose transporter type 1 deficiency syndrome (GLUT1DS) is a treatable encephalopathy, caused by a defect of glucose uptake, mediated by GLUT1, at the blood brain barrier and into brain cells due to mutations in SLC2A1 gene (solute carrier family 2 member 1) \cite{Wang2005}. The classical phenotype includes a range of complex movement disorders, drug-resistance epilepsy, mental retardation, and acquired microcephaly \cite{DeVivo1991,Seidner1998}. Less severe disorders mainly characterized by paroxysmal exercised-induced dyskinesia (PED) and ‘generalized epilepsies’, either alone or in combination, have been reported \cite{Arsov2012a,Striano2012,Suls2008,Suls2009}. When epilepsy is the main symptom, absence seizures (AS), mainly with early onset, seem to be the most frequent clinical expression of SLC2A1 mutations and in individual cases AS may be phenotypically indistinguishable from absences of the genetic generalized epilepsies (GGE; formerly known as idiopathic generalized epilepsies) \cite{Arsov2012a,Mullen2010,Striano2012}. Furthermore, during childhood, the most common EEG abnormalities are 2.5–4 Hz generalized spike and wave discharges (GSWD) \cite{Leary2003}. This pattern is usually more polymorphic and irregular than the classic 3–Hz spike-wave \cite{Leary2003}, but could be likewise similar and elicited by hyperventilation. Given this combination of EEG features and seizure type, GLUT1DS should be considered in the differential diagnosis of childhood (CAE) or juvenile absence epilepsy (JAE), particularly if atypical features are present \cite{Byrne2011}. Myoclonic-astatic and myoclonic absences epilepsy phenotypes have been also reported \cite{Larsen2015,Mullen2011}. A chronic metabolic disturbance caused by lowered glucose transport is considered to be

Purpose: To provide imaging biomarkers of generalized spike-and-wave discharges (GSWD) in patients with GLUT1 deficiency syndrome (GLUT1DS).

Methods: Eighteen GLUT1DS patients with pathogenetic mutation in SLC2A1 gene were studied by means of Video-EEG simultaneously recorded with functional MRI (VideoEEG-fMRI). A control group of sex and age-matched patients affected by Genetic Generalized Epilepsy (GGE) with GSWD were investigated with the same protocol. Within and between groups comparison was performed as appropriated. For GLUT1DS, correlations analyses between the contrast of interest and the main clinical measurements were provided.

Results: EEG during fMRI revealed interictal GSWD in 10 GLUT1DS patients. Group-level analysis showed BOLD signal increases at the premotor cortex and putamen. With respect to GGE, GLUT1DS patients demonstrated increased neuronal activity in the putamen, precuneus, cingulate cortex, SMA and paracentral lobule. Whole-brain correlation analyses disclosed a linear relationship between the GSWD-related BOLD changes and the levels of glycorrhachia at diagnosis over the sensory-motor cortex and superior parietal lobuli.

Conclusion: The BOLD dynamics related to GSWD in GLUT1DS are substantially different from typical GGE showing the former an increased activity in the premotor-striatal network and a decrease in the thalamus. The revealed hemodynamic maps might represent imaging biomarkers of GLUT1DS, being potentially useful for a precocious diagnosis of this genetic disorder.

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responsible for generalized epileptiform activities and epilepsy in patients with GLUT1 deficiency (Pascual et al., 2002; Seidner et al., 1998).

This study is the first investigation of the brain networks involved in the generation of GSWD in GLUT1DS by means of EEG co-registered with fMRI (EEG-fMRI). In particular, we aimed to isolate the BOLD correlates of GSWD in GLUT1DS with respect to the ones typically observed in GGE (Benuzzi et al., 2012; Gotman et al., 2005; Moeller et al., 2008; Li et al., 2009). This objective could be relevant for the following reasons: (a) to better understand the pathophysiology GLUT1DS; (b) to provide imaging biomarkers of the disorder that might orient the diagnosis; (c) to evaluate the mechanisms of ‘generalized’ SWD in a model not due to a channelopathy.

2. Materials and methods

2.1. Patients

Eighteen Italian GLUT1DS patients (mean age 19.22 years; range 6–43 years; 12 females) were enrolled from September 2012 to March 2015. The recruited subjects fulfilled the clinical criteria for diagnosis of GLUT1DS (Pearson et al., 2013) and all but one presented pathogenic mutations in SLC2A1 gene (Table 1).

The clinical, molecular methods and genetic features of this GLUT1DS cohort have been previously described (De Giorgis et al., 2015; Gagliardi et al., 2012; Ragona et al., 2014; Veggio et al., 2010; Zorzi et al., 2008). Briefly, mutation analysis of all exons and intron-exon boundaries of SLC2A1 was performed on genomic DNA by direct sequencing. Each fragment was sequenced on both strands. The effect of the newly detected GLUT1 mutations on protein structure or function was analyzed with the prediction programs ExPASy (http://www.expasy.org).

The mean age at diagnosis was 16.2 ± 11.1 years. Structural MRI scan (3T) was normal in all cases. All patients but 2 (#6; #18) experienced epileptic seizures (mean age of seizure onset was 34 ± 21 months). The most representative seizure’s type was AS (10 out of 18 patients) followed by generalized tonic-clonic and myoclonic seizures. Notably, four patients had AS alone and one patient had myoclonic seizures without movement disorders. Eleven out of 18 GLUT1 subjects presented complex movement disorders, like ataxia, dystonia, chorea and paroxysmal exercise-induced dyskinesia. An intellectual impairment of variable degree was detected in 8 patients (according to the Diagnostic and Statistical Manual of Mental Disorders-DSM V). The rest of the sample displayed normal intellectual abilities, although for most of them the IQ was in the low range of normality. Seven out of the patients were on ketogenic diet either at different regimens alone (patients #4, #6, #11) or in combination with antiepileptic drugs (#2, #8, #10, #18).

A group of 18 patients with GGE served as control (mean age 24.88 years; range 5–48 years; 12 females). Patients were extracted from our database of GGE patients who underwent an EEG-fMRI study between April 2007–December 2013. The patients’ inclusion criteria were the followings: 1) documented GSWD with AS during routinely video-EEG monitoring; 2) routine EEG showing normal background; 3) normal developmental milestones; 4) normal structural imaging on anatomic MRI at 3T. The GGE syndromes classification was obtained by two expert epileptologists after a careful revision of clinical and electrophysiological data of each patient (S.M., A.E.V.) according to the ILAE classification (1981). Table 2 summarizes the clinical and EEG findings of GGE population.

The human ethic committee of the University of Modena and Reggio Emilia approved this study and written informed consent was obtained from all the patients and from their parents if underage.

2.2. EEG with fMRI recordings and analysis

Video-EEG signals have been recorded by means of a 32 channels MRI compatible EEG recording system (Micromed, Italy). Patients were asked to rest with eyes closed and keep still. All recordings were performed in the early afternoon. Sedation was never used. After the fMRI acquisitions, both GLUT1DS and GGE patients were interviewed for discomfort or any clinical, even subjective, ictal events while scanning.

fMRI data (30 axial slices, TR/TE = 3000/50 ms) have been acquired using a 3T scanner (Philips) over three 10-min sessions (200 volumes/}

Table 1

| PI ID | Age(yrs)/sex | Age at diagnosis (yrs) | Glycorrhachia (mg/dl) | CSF/plasma glucose ratio | Protein mutation | Seizure type | Seizure onset (mo) | Sporadic/Familiar | Motor Signs | IQ | Treatment at fMRI |
|------|--------------|------------------------|-----------------------|--------------------------|------------------|--------------|-------------------|------------------|-------------|----|------------------|
| 1    | 12/M         | 6                      | 35                    | 0.54                     | c1457delEG1_1delcSPL | GTC, AS       | 11                | S                | PED, Dy     | 75  | VPA              |
| 2    | 11/F         | 10                     | 33                    | 0.34                     | R126C            | Myo, AS      | 30                | S                | PED, Dy     | 44  | LEV, LTG, 3:1 KD |
| 3    | 14/M         | 11                     | 44                    | 0.51                     | R223W            | FS           | 11                | S                | PED, Cho    | 75  | OXC              |
| 4    | 9/F          | 5                      | 31                    | 0.39                     | R153C            | GTC, AS      | 12                | F                | A            | 104 | 1:8:1 KD         |
| 5    | 12/M         | 10                     | 40                    | 0.47                     | R458W            | FS           | 40                | F                | PED, Cho    | 85  | VPA              |
| 6    | 10/F         | 10                     | 33                    | 0.38                     | V165I            | /            | /                 | F                | PED, Dy, Cho| 66  | 3:1 KD           |
| 7    | 7/F          | 7                      | 32                    | 0.35                     | N34S             | AS           | 24                | F                | /           | 76  | ETS              |
| 8    | 12/F         | 9                      | 34                    | 0.38                     | R400C            | AS, Myo, Myo | 36                | S                | PED, Dy     | 50  | LTG, 3:1 KD      |
| 9    | 6/F          | 5                      | 40                    | 0.41                     | not found        | AS, GTC      | 39                | S                | /           | 112 | VPA, VTC, CLB    |
| 10   | 27/F         | 20                     | 31                    | 0.38                     | R126C            | AS, FS       | 8                 | S                | PED, A      | 60  | LEV, 3:1 KD      |
| 11   | 18/F         | 14                     | 40                    | 0.44                     | P36R             | AS           | 18                | S                | /           | 53  | 2.5:1 KD         |
| 12   | 28/M         | 17                     | 46                    | 0.54                     | R458W            | FS           | 60                | S                | PED, Dy     | 57  | ETS              |
| 13   | 43/F         | 43                     | N/A                   | N/A                      | R458W            | AS           | 72                | F                | /           | 78  | –                |
| 14   | 40/F         | 29                     | 41                    | 0.44                     | V165I            | AS, GTC      | 12                | F                | PED         | 55  | VPA              |
| 15   | 24/F         | 24                     | N/A                   | N/A                      | V165I            | AS           | 72                | F                | /           | 76  | –                |
| 16   | 19/F         | 19                     | 38                    | 0.5                      | 1166delV         | AS           | 48                | S                | PED, Dy     | 80  | LTG              |
| 17   | 39/M         | 39                     | 42                    | 0.43                     | N34S             | AS           | 48                | F                | PED, Dy     | 75  | CBZ              |
| 18   | 15/M         | 15                     | 44                    | 0.44                     | R400L            | /            | /                 | S                | PED, Dy     | 56  | VPA, 3:1 KD      |

Legend Table 1: M: Male; F: Female; A: Ataxia; AS: absences seizures; Cho: chorea; Dy: Dystonia; GTC: generalized tonic clonic seizures; FS: focal seizures; Myo: Myoclonic Seizures; N/A: not available; PED: paroxysmal exercise-induced dyskinesia. LEV: Levetiracetam; LTG: Lamotrigine; VPA: Valproate; ETS: Ethosuximide; CBZ: Carbamazepine; CLB: Clobazam; KD: ketogenic diet. S: sporadic; F: familiar.

Missense mutations were observed in the majority; case #1 and n #16 presented deletions. Amino acid mutations are represented by (capital) native amino acid (in one-letter code), amino acid number, and mutant amino acid. Nucleotide deletions are indicated by nucleotide number, del for deletion, nucleotide, or by native nucleotide, number, and mutant nucleotide.
2.2.2. Group level analysis

After offline correction of the gradient artifacts and filtering of the EEG signal (Allen et al., 2000), the EEG data were reviewed and pre-processed according to our previous published method (Avanzini et al., 2014; Ruggieri et al., 2015). Two experienced electroencephalographers reviewed the pre-processed EEG recordings independently (A.R., P.V.) in order to identify GSWD. After identification, epileptiform activities were manually marked on EEG and exported as single event (stick functions or blocks) within the General Linear Model (GLM) pre-processing and a Volterra expansion of these (Friston et al., 1996). All re-alignment parameters [six scan realignments parameters from image alignment parameters] were included in the model as confounds.

The effect of GSWD at the group level was tested for GLUT1DS and GGE patients separately by means of the 2.2.2. Group level analysis et al., 2008). In case of blocks, the duration of GSWD was specified and smoothed with 8 × 8 × 8 mm full-width half maximum Gaussian Kernel. Movement artifacts indistinctly identified by analysis of Video-EEG recordings (blinking, lip smacking, swallowing, head movements) were considered as confounders in the model and treated as stick functions or blocks as appropriated (Ruggieri et al., 2015). In addition, 24 re-alignment parameters [six scan realignments parameters from image pre-processing and a Volterra expansion] were included in the model as confounds.

2.2.1. First level analysis

Maps of the t-statistic were created using the timing of the GSWD as events (stick functions or blocks) within the General Liner Model (GLM) framework. In case of blocks, the duration of GSWD was specified. All regressors were convolved with the canonical hemodynamic response function (HRF) plus its temporal and dispersion derivatives (Lermieux et al., 2008).

2.2.2. Group level analysis

The effect of GSWD at the group level was tested for GLUT1DS and GGE patients separately by means of the fixed-effect or the random effect approach as appropriate (Friston et al., 1999). When using the fixed-effect model, a conjunction analysis of the BOLD increases and decreases effects for all the subjects was done with global null hypothesis, to evaluate whether the selected congruent contrasts were consistently high and jointly significant (Friston et al., 1999). For the random effect analysis, a full factorial design was used to test for within effects (1-way analysis of variance [ANOVA]) with 1 group of patients (18 GGE) and three basis functions (canonical HRF and its derivatives) as factors. Group analysis results were considered significant at the p < 0.05 corrected for FWE or, if any results were revealed, a less conservative statistical inference was considered (p < 0.001, uncorrected). In these cases, an additional threshold of 5-voxels extent was applied in order to discard BOLD changes occurring in scattered voxels. The fMRI t-maps were superimposed on the canonical T1-weighted images as implemented in SPM8.

Subsequently, we compared the BOLD changes obtained in the two populations in order to isolate the GSWD-related hemodynamic patterns specific of the two epileptic conditions. To this end, 10 age and sex matched GGE patients were extracted from the whole GGE sample (patient #7, #9, #10, #11, #12, #13, #14, #15, #17, #18). Within this subgroup, a total number of 149 GSWD were recorded (mean duration 4.59 s). A single fixed effect analysis that included both groups in one matrix (10 GLUT1DS and 10 GGE) was performed. The GSWD effect across the 10 GGE patients was obtained using fixed effect approach with conjunction analysis (Friston et al., 1999) similar to what previously described for GLUT1DS. Further, we explored the differences by applying an exclusive masking procedure between the conjunction contrasts each related to the specific patients' populations. In details, to isolate the brain regions that were significantly involved in the main effect “GSWD > baseline” in GLUT1DS population but not in the main effect “GSWD > baseline” in GGE, the conjunction contrast “GSWD > baseline” in GLUT1DS was exclusively masked by the mask conjunction contrast “GSWD > baseline” in GGE. By counterpart, to isolate the brain regions exclusively activated by GSWD in GGE, the conjunction contrast “GSWD > baseline” in GGE was exclusively masked by the mask conjunction contrast “GSWD > baseline” in GLUT1DS. SPM exclusive masks were thresholded at p < 0.05 uncorrected, whereas the contrasts to be masked were thresholded at p < 0.001. In this way, those voxels that reached a level of significance at p < 0.05 in the mask contrast were excluded from the analysis.

Finally, for the GLUT1DS patient group only, we performed additional whole brain second-level correlations analyses between the contrast (GSWD) and the main clinical measurements including: age (years); age at seizure onset (months); total IQ; glycorrhachia values (mg/dl) and the CSF/plasma glucose ratio at the time of diagnostic workup. We also explored the effect of the ketogenic diet (KD) on the GSWD-related BOLD changes. Indeed, five GLUT1DS patients (#6, #8, #10, #11, #18) were on KD (alone or with antiepileptic treatment-AED) (KD+) at the time of the fMRI study, while the remaining were naïve (#13) or only under AED (#3, #5, #9, #17) (KD−). In a separate analysis, we have performed a conjunction analysis, based on the fixed effect model, of the GSWD-related T contrast in the KD+ subgroup and then

| Clinical Data | GSWD during fMRI |
|--------------|-----------------|
| Pts ID | Age (years)/sex | Epilepsy syndrome | AEDs | N° events | Mean duration (sec) | Minimum Duration (sec) | Maximum Duration (sec) |
| 1 | 24/F | JAE | LEV | 45 | 3.24 | 1.5 | 5.4 |
| 2 | 33/F | JAE | VPA | 15 | 3.61 | 0.8 | 6.8 |
| 3 | 47/M | JAE | VPA | 21 | 3.71 | 2 | 5 |
| 4 | 48/F | CAE | VPA | 14 | 4.03 | 1.2 | 6.1 |
| 5 | 26/M | Adult onset IGE | PB/TPM/ZNZ | 12 | 4.99 | 2 | 7 |
| 6 | 37/F | JAE | LEV | 19 | 3.91 | 1.6 | 8.1 |
| 7 | 14/F | JAE | VPA | 16 | 3.18 | 1.1 | 6.9 |
| 8 | 23/F | Adult onset IGE | LEV | 3 | 3.07 | 0.9 | 5 |
| 9 | 5/F | CAE | Naïve | 5 | 5.20 | 2.2 | 7 |
| 10 | 20/F | JAE | Naïve | 9 | 3.14 | 1.1 | 5.2 |
| 11 | 12/M | CAE | Naïve | 5 | 7.29 | 2.2 | 10 |
| 12 | 9/F | CAE | Naïve | 3 | 7.63 | 1.1 | 15 |
| 13 | 11/M | JAE | VPA | 9 | 5.19 | 3.1 | 7.6 |
| 14 | 10/F | CAE | Naïve | 6 | 5.07 | 2.2 | 6.9 |
| 15 | 41/F | Adult onset IGE | LEV | 47 | 2.13 | 1 | 5 |
| 16 | 35/F | JAE | VPA | 15 | 2.85 | 2.2 | 3.6 |
| 17 | 15/M | JME | LEV | 8 | 3.40 | 2.1 | 4.4 |
| 18 | 38/M | JAE | VPA | 41 | 3.74 | 1.7 | 6.6 |

Legend Table 2: M: Male; F: Female; JAE: juvenile absence epilepsy; CAE: childhood absence epilepsy; GGE: genetic generalized epilepsy; JME: juvenile myoclonic epilepsy; AEDs: antiepileptic drugs; VPA: valproate; LEV: levetiracetam; PB: phenobarbital; TPM: topiramate; ZNZ: zonisamide; GSWD: generalized spike-wave discharge.
we compared this effect with the one obtained in KD-. The differences were explored by applying an exclusive masking procedure at a threshold of $p < 0.05$, uncorrected for multiple comparisons.

3. Results

3.1. EEG findings during fMRI

In GGE, EEG during fMRI shown symmetric and synchronous generalized 3–4 Hz SWD in all the patients. A total of 293 events (mean of 17 events per subject) were acquired, from a minimum of three GSWD in subject #8 and #12, to a maximum of 47 events in subject #15. The duration of GSWD ranged from 1 s to 15 s, with a mean duration across subjects of 4.2 s (see Table 2). The GSWD occurrence was not accompanied by clinical manifestations, even subjective, in any patients.

In GLUT1DS, EEG during fMRI revealed SWD in 10 subjects. Six patients (#3, #5, #6, #9, #11, #17) presented 3–4 Hz diffuse spike-and-waves discharges; four patients showed more focal discharges over the bilateral frontal leads (#8, #10, #13, #18). A total number of 152 SWD were recorded (range: 6–28). The duration of GSWD ranged from 1 s to 15 s, with a mean duration across GLUT1DS of 4.3 s (see Table 3). Fig. 1, panel A shows the EEG pattern in a representative case (#10). The GLUT1DS patients did not shown any clinical seizures, even subtle or subjective, during the EEG-fMRI. In addition, no persistent or paroxysmal movement disorders (dystonia, dyskinesia, chorea) were recorded during the experimental sessions.

For both GLUT1DS and GGE populations, the absence of clinical events (seizure and movement disorder) was verified by either the inspection of the Video recorded simultaneously during fMRI acquisitions and the post-scanning individual patient’s interview.

3.2. GSWD related BOLD changes in GLUT1DS

Table 3 summarizes the fMRI findings at single subject level. GSWD-related hemodynamic changes were distributed homogenously across patients, revealing the involvement of the same cortico(frontal)-subcortical(basal ganglia) network. In particular, all GLUT1DS demonstrated BOLD changes at a frontal network, encompassing premotor (prefrontal cortex, supplementary motor area-SMA, cingulate cortex) (9 patients) and motor brain regions (5 subjects). The frontal BOLD response was positive in all the cases except four patients (#6, #8, #13, #18). Subcortical positive hemodynamic changes were mainly confined at the basal ganglia (putamen) (5 patients) while only one patient (#13) demonstrated thalamic BOLD signal increases. Finally, 7 GLUT1DS demonstrated significant BOLD responses at the posterior part of the Default Mode Network, essentially as signal increases (6 subjects). Fig. 1, panel B shows the GSWD-related fMRI map in a representative case (#10).

At the group-level, fixed effect analysis ($p < 0.05$ corrected for FWE) showed BOLD positive changes over the bilateral premotor cortex (global maxima-GM) (Brodmann Area-BA9), precuneus (BA7), inferior frontal gyri (BA46) and SMA (BA6), while decreases involved the bilateral motor cortex (BA2) and the pericentral lobule (BA5) (Fig. 2A). At a less conservative threshold ($p < 0.001$ uncorrected) a bilateral thalamic deactivation was detected. The conjunction analysis ($p < 0.001$ uncorrected) revealed commonalities in BOLD signal increases between the ten subjects in the bilateral premotor cortex (GM) and right putamen; negative BOLD changes were detected in the thalamus, primary motor cortex, inferior frontal gyri. The BOLD increase in the premotor cortex survived at a more conservative threshold ($p < 0.05$ corrected for FWE) (Fig. 2B).

3.3. GSWD-related BOLD changes in GGE

At the group level (n. of patients = 18), random effect analysis ($p < 0.05$ corrected for FWE) revealed BOLD signal increases in the bilateral thalami (GM) and cerebellum, while negative changes were observed in the posterior part of the Default Mode Network (DMN) encompassing the bilateral precuneus (BA7), and inferior parietal lobule (BA40) (Supplementary Fig. 1). When the analysis was limited to 10 GGE, the conjunction approach based on the fixed-effect model

| PT ID | Type and location | Number/mean duration (sec) | Minimum duration (sec) | Maximum duration (sec) | Basal Ganglia | Frontal operculum | Premotor cortex | SMA | Cingulate cortex | Pericentral cortex | DMN** | Other |
|-------|------------------|-----------------------------|------------------------|------------------------|----------------|-------------------|-----------------|-----|----------------|-------------------|-------|-------|
| PT3   | B F SWD          | 12/11.89                    | 2.68                   | 18.1                   | B (R)          | B (L)             | B (R)           | L (4.50) | –              | –                 | L (4.90) | –     |
| PT5   | Diffuse SWD      | 6/2.1                       | 3.24                   | –                      | –              | –                 | –               | –              | –              | –                 | –      | –     |
| PT6   | Diffuse SWD      | 12/2.17                     | 0.36                   | 4.53                   | B (L) (2.58)   | –                 | L (3.46)        | (d)*         | B (L) (2.89)   | (d)*              | –      | B cerebellum |
| PT8   | B F SWD          | 11/2.14                     | 0                      | 4.01                   | –              | –                 | R (d)*          | –              | –              | –                 | –      | –     |
| PT9    | Diffuse SWD      | 15/3.2                      | 0                      | 15.24                  | L (2.72)       | –                 | R (4.35)        | (i)          | L (3.02)       | (i)               | B (R) (3.75) | (i)*     |
| PT10  | B F SWD          | 18/4.28                     | 1                      | 17.84                  | B (L) (4.86)   | B (L) (3.99)     | B (L) (7.59)    | (i)          | L (7.59)       | (i)               | B (L) (5.07) | (i) |
| PT11  | Diffuse SWD      | 15/1.34                     | 0                      | 2.05                   | –              | B (4.92)         | –               | B (4.94)     | –              | –                 | B (L) (6.15) | –     |
| PT13  | B F SWD          | 28/3.6                      | 0                      | 15.8                   | L (3.82)       | (i)*             | B (R) (5.67)    | (d)          | B (R) (6.41)   | (d)               | B (R) (7.81) | (d) |
| PT17  | Diffuse SWD      | 10/3                        | 1                      | 4.5                    | –              | –                 | B (L) (3.32)    | (i)          | B (L) (3.17)   | (i)*               | R (3.77) | –     |
| PT18  | B F SWD          | 25/3.5                      | 8.4                    | –                      | –              | –                 | B (L) (5.39)    | (d)          | –              | –                 | –      | –     |

Legend: Table 3: fMRI results are displayed using a threshold set at $p < 0.05$ corrected for FWE and $p < 0.001$ uncorrected if no BOLD changes were detected at the more conservative statistical inference (the results obtained using a $p < 0.001$ uncorrected are indicated by the asterisk [*].) The numbers in brackets indicate the maximal t-values. ED: epileptic discharges; SW: Slow Waves; SWD: spike-wave discharges; L: Left; R: Right; B: Bilateral; F: Frontal; TP: Temporo-Parietal. (i) increase BOLD signal; (d) decrease BOLD signal; L: Left; R: Right; B: Bilateral; SMA: Supplementary Motor Area; **DMNp: Default Mode Network which includes: posterior part of DMN (precuneus/posterior cingulate, bilateral inferior parietal lobuli).
revealed commonalities in BOLD signal increases in the bilateral thalami (more right) and cerebellum.

3.4. GLUT1 versus GGE

GLUT1DS patients demonstrated during GSWD increased neuronal activity in the bilateral putamen, precuneus, middle cingulate cortex, right SMA and right paracentral lobule, while GGE do not (Fig. 3A). On the contrary, no GGE-exclusive GSWS-BOLD correlates were observed at the considered threshold.

3.5. Correlations between BOLD signal and clinical measures

Whole-brain correlation analyses using individual clinical characteristics of the GLUT1DS patients disclosed a linear positive relationship between the interictal discharges-related BOLD changes and the levels of glycorrhachia (and also CSF/plasma glucose ratio) over a bilateral cortical network including the opercular-insular cortex, the post-central cortex, and superior parietal lobule (Fig. 3B). No significant correlations were discovered with the age, age at seizure onset, and the total IQ.

The conjunction analysis limited to KD+ patients ($p < 0.001$, uncorrected for FWE) demonstrated common BOLD positive responses over the bilateral precuneus, bilateral inferior parietal lobule, right precentral gyrus and a small spot in the left putamen (data not shown). No negative responses were detected. The comparison between the hemodynamic changes in GLUT1DS KD+ versus KD-, did not isolated any BOLD cluster specific of GLUT1DS KD+ and viceversa, at the considered threshold.

4. Discussion

This work represents the first attempt to investigate the hemodynamic correlates of interictal epileptiform activity (IED) in a cohort of patients with GLUT1DS. To note, previous functional neuroimaging studies in these patients (Akman et al., 2015; Pascual et al., 2002, 2007; Suls et al., 2008) were unable to obtain the EEG recordings at the time of imaging and the interictal/ictal state was assessed only by clinical inspection. In our study, the video-EEG recorded simultaneously during fMRI acquisitions allowed both to record the cerebral brain activity and to exclude the presence of any clinical seizure as well as paroxysmal movement disorder during the experimental sessions. Consequently, the resulted maps represent a real-time picture of brain function during IED in GLUT1DS.

The studied population is representative of the complex phenotypical spectrum of the disorder and particularly with early onset absence seizures as the main seizure type (Arsov et al., 2012b; Byrne et al., 2011; Leary et al., 2003; Striano et al., 2012; Wang et al., 2005) and paroxysmal exertion-induced dystonia (PED) as the most frequent movement disorder (Suls et al., 2008). However, epilepsy and motor signs were not homogenously distributed across patients as well as treatment and age at the time of the fMRI study. This heterogeneous presentation reflects the intrinsic and natural variability of the syndrome, as largely documented (Brockmann, 2009; De
The clinical variability of GLUT1DS represents a difficulty to diagnose the disorder, especially for patients with mild or non-classical phenotype, who are likely to be under-diagnosed (Brockmann, 2009; De Giorgis et al., 2015). In this contest, the present work might be of importance to underlie the pathophysiological fingerprints of this metabolic encephalopathy.

As far as the fMRI analyses, the results of our work are innovative for two main aspects. Firstly, the observed BOLD findings allowed delineating a well-defined network of cortical-subcortical regions that are involved in the generation of GSWD in GLUT1DS. The results at the second level analysis showed an increase BOLD signal, and hence an increased neuronal activity, in the premotor-striatal network (bilateral premotor cortex, SMA, anterior cingulate and basal ganglia) and a simultaneous reduced metabolic demand of the motor cortex and thalamus. Notably, the analysis of single-subject GSWD-related BOLD changes was consistent with the group results as most of the patients showed significant BOLD changes within the premotor-basal ganglia circuit (see Table 3). The altered metabolism of thalamus and cortico-striatal pathways found in our patients speaks to the severe abnormality...
of this functional system and leads us to hypothesize that it constitutes the central epileptogenic substrate in GLUT1DS.

As second main result, our findings shown that the BOLD dynamics related to GSWD in GLUT1DS are substantially different from the BOLD correlates of GSWD in typical genetic generalized epilepsies (Aghakhani et al., 2004; Benuzzi et al., 2012; Carney et al., 2010; Gotman et al., 2005; Hamandi et al., 2006; Li et al., 2009; Moeller et al., 2010; Vaudano et al., 2009), reinforcing the hypothesis of a specific relationship with the GLUT1DS disorder. Intriguingly, the cortical networks observed in GLUT1DS patients are very similar to the one described, by means of EEG-fMRI, in patients with ‘Doose syndrome’ or Myoclonic Astatic Epilepsy (MAE) (Moeller et al., 2014). This commonality is of particular interest since MAE has been reported as one of the possible phenotypes of GLUT1DS (Mullen et al., 2011). With respect to the BOLD findings previously reported in MAE (Moeller et al., 2014), and more in general to GGE sub-syndromes, our GLUT1 population didn't show any increase in BOLD signal at thalamic level and no decreases in the Default Mode Network (DMN) at a group level. By contrast, GLUT1DS revealed bilateral activation in the putamen and different premotor and parietal areas (Fig. 3A); this pattern in not generally reported in GGE, except for single cases description (see Carney and Jackson, 2014 as review). The consistency of 3–4 Hz GSWD associated hemodynamic changes in GGE across studies, further support that the mechanisms subserving GSWD generation in GLUT1DS involve different neural dynamics. The irregular and slower phenotype of GSWD in GLUT1DS might partially explain these BOLD differences. Nevertheless, previous evidences failed to detect specific hemodynamic pattern linked to different GGE subgroups, independently to the morphology and length of GSWD (Pugnaghi et al., 2014).

The revealed hemodynamic changes in GLUT1DS are concordant with findings observed by means of positron emission tomography (PET) imaging in humans (Akman et al., 2015; Pascual et al., 2002–2007; Suls et al., 2008). Overall considered, interictal 18F-fluorodeoxyglucose PET demonstrated a relatively homogeneous pattern with glucose hypometabolism in the mesial temporal regions, pre-postcentral gyrus, premotor cortex, inferior parietal lobe and thalamus, and a relative increased in the basal ganglia (putamen). These PET findings demonstrated to be stereotyped regardless of disease severity, or type and duration of therapy, and persist into adulthood (Pascual et al., 2002–2007). In other words, hypometabolism pattern may not

![Image](image-url)
show a progressive changes with increasing age or clinical course of seizures (Akman et al., 2015). This argument is concordant with our fMRI findings, as similar BOLD patterns were observed in adults and children and no relationship has been detected between age and IED-related hemodynamic changes. To note, we observed that BOLD signal related to epileptic discharges at the time of fMRI correlated with the level of CSF glucose at the time of diagnosis (Fig. 3B). This means that the lower the CSF glucose in early childhood, the lower are the BOLD signal changes IED-related, several years later at the time of the imaging study. Such linear correlation is particularly evident at the bilateral sensory-motor and parietal circuits which neuronal activity is highly dependent to the glucose supply especially in early infancy (Chugani, 1998). This finding supports the observations of Akman et al. (2015) who suggested “that brain glucose hypometabolism, present in infancy, leaves an imprint that persists even when the precipitating condition causing the hypoglycorrhachia is corrected”. Probably, hypoglycorrachia during a vulnerable period in early infancy can affect the maturation of the developing striatal-thalamo-cortical networks and synaptic connections, leaving a persistent alteration even when hypoglycorrachia is corrected. As far as the correlation between the GSWD-related BOLD changes and other clinical variables, no relationship was detected with the type of treatment, particularly in relation to the presence/absence of KD (KD + versus KD-). Such inconclusive result needs to be taken with caution given the low number of patients in each subgroup. A further dedicated study on a larger cohort of GLUT1DS patients with IED during fMRI scanning and different treatments (KD and/or AED) needs to be performed to test the specific effect of KD on the brain networks function in GLUT1DS.

Our fMRI results demonstrated that the disrupted metabolism of the premotor-basal ganglia circuit is responsible for the generation and maintenance of interictal GSWD. Since we did not record any clinical seizure as well as involuntary movements during the experimental sessions, we could not directly link the detected hemodynamic changes with these clinical phenomena. Previous attempts to segregate the PET patterns based on the GLUT1DS clinical phenotype shown that a prominent hypometabolism in the thalamus and mesial temporal cortex linked to the presence of epilepsy (Akman et al., 2015) while PED frequency score was positively correlated to the metabolic activity of basal ganglia (putamen), and negatively with the premotor cortex (anterior cingulate and superior frontal cortex) (Suls et al., 2008). Intriguingly, ictal SPECT during a PED episode shown hyper-perfusion over the motor cortex and putamen (Suls et al., 2008). Based on these observations, a dysfunction of the cortico-striatal pathways has been postulated to play a role in the pathophysiology of PED (Suls et al., 2008). Our fMRI maps appear to support this hypothesis as most of our patients shown PED as the main motor disorder. It is likely that an altered excitability of striatal-thalamic-(frontal)cortical pathway, which is the consequence of a defect in glucose transport across the blood-brain barrier, suberves the generation of both epileptic activity and PED. Given the lack of movement disorders during fMRI, this conclusion remains speculative and warrants further investigations to be confirmed.

The pathogenic role of low CSF glucose to determine GLUT1DS phenotype, including seizures, has been objected of intense speculations. Aghakhani, Y., Bagshaw, A.P., Bénar, C.C., Hawco, C., Andermann, F., Dubeau, F., Gotman, J., 2004. IED activation during spike and wave discharges in idiopathic generalized epilepsy. Brain 127, 1127-1144.

4.1. Study limitations

The major limitation of this study is related to the number of investigated subjects, which is justified by the rarity of this genetic disorder. This limitation might lead to caution in generalizing the results outside of the group of subjects analyzed. The fixed effect approach applied within the GLUT1DS cohort and between the GLUT1DS and GGE, allows to make a comment about the specific subjects studied (only those) but it cannot be used to obtain an inference at the population level. However, by using a conjunction approach, based on the fixed-effect model, we were able to infer that at least some proportion of the population from which the patients came show this effect (Friston et al., 1999), i.e. to extend our speculations to the general GLUT1DS population.

4.2. Conclusions

In conclusion, our study provides new findings on the epileptic networks subserving epilepsy in GLUT1 deficiency syndrome. Several questions, however, are unanswered and should be addressed by future studies investigating both microstructural abnormalities in GLUT1 patients with milder form of the disease, as well as the integrity or alteration of resting-state networks subserving motor and cognitive functions.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.nicl.2016.12.026.

Fundings

A.E. Vaudano received research grant support from the Fondazione Epilessia LICE (Lega Italiana contro L’Epilessia) Onlus, and from research grant support from the Ministry of Health (MOH), Regione Emilia Romagna, grant ID RPRUA1GR-2013-0000120. S. Olivotto received research grant support from AIEF (Associazione Italiana Epilessia Farmacoresistente) Onlus. A. Ruggieri received a PhD bursary from the University of Modena and Reggio Emilia. V. De Giorgis received a PhD bursary from the University of Pavia. P. Veggiotti received educational grant from EISAI S. Meletti received Research grant support from the Ministry of Health (MOH), grant ID NET-2013-0235513-3; from the non-profit organization CarisMO Foundation, grant ID A.010@FCRMO RINT@MELFONINFO; has received personal compensation as scientific advisory board member for UCB and EISAI. Other authors report no disclosures. The authors have stated that they had no interests that might be perceived as posing a conflict or bias.

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