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Report on three additional patients and genotype-phenotype correlation in *SLC25A22*-related disorders group

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Abstract

Early infantile epileptic encephalopathy (EIEE) is a heterogeneous group of severe forms of age-related developmental and epileptic encephalopathies with onset during the first weeks or months of life. The interictal electroencephalogram (EEG) shows a "suppression burst" (SB) pattern. The prognosis is usually poor and most children die within the first two years or survive with very severe intellectual disabilities. EIEE type 3 is caused by variants affecting function, in *SLC25A22*, which is also responsible for epilepsy of infancy with migrating focal seizures (EIMFS). We report a family with a less severe phenotype of EIEE type 3. We performed exome sequencing and identified two unreported variants in *SLC25A22* in the compound heterozygous state: NM_024698.4: c.[813_814delTG];[818 G>A] (p.[Ala272Glnfs*144];[Arg273Lys]). Functional studies in cultured skin fibroblasts from a patient showed that glutamate oxidation was strongly defective, based on a literature review. We clustered the 18 published patients (including those from this family) into three groups according to the severity of the *SLC25A22*-related disorders. In an attempt to identify genotype–phenotype correlations, we compared the variants according to the location depending on the protein domains. We observed that patients with two variants located in helical transmembrane domains presented a severe phenotype, whereas patients with at least one variant outside helical transmembrane domains presented a milder phenotype. These data are suggestive of a continuum of disorders related to *SLC25A22* that could be called *SLC25A22*-related disorders. This might be a first clue to enable geneticists to outline a prognosis based on genetic molecular data regarding the *SLC25A22* gene.

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Introduction

Early infantile epileptic encephalopathy (EIEE) is a large group of the most severe forms of age-related developmental and epileptic encephalopathic diseases. It is

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characterized by erratic refractory seizures, usually myoclonic, with an onset during the first weeks or months of life associating with severe developmental delay, often evolving into West syndrome. There are about 70 types of EIEE classified based on the gene involved. In the majority of EIEE, the interictal electroencephalogram (EEG) shows typically a "suppression burst" (SB) pattern with highvoltage bursts of slow waves mixed with multifocal spikes alternating with isoelectric suppression phases [1]. In most cases, brain structural abnormalities are observed [1]. The prognosis is usually poor and most children die within the first two years of life or survive with very severe intellectual disabilities [1]. The EIEE type 3 (OMIM #609304) is characterized by myoclonic seizures, severe global hypotonia, microcephaly, SB pattern, and abnormal electroretinogram (ERG) [2]. EIEE type 3 is caused by variants affecting function in the solute carrier family 25 member 22 gene (SLC25A22, OMIM *609302) [2, 3]. SLC25A22 encodes a 323 amino acids mitochondrial glutamate/H+ symporter that is particularly expressed in the brain [4]. Initially, seven patients from three consanguineous families were reported [2, 3, 5]. These seven patients presented a similar severe phenotype: early-onset seizures, severe hypotonia, microcephaly, abnormalities on brain imaging, SB pattern, and died at 8-year-old age or were reported in a vegetative state. Recently, Reid et al. reported six additional children from three families, presenting variants in SLC25A22 and milder phenotypes than the previous reported patients [6]. In addition, SLC25A22 is also involved in epilepsy of infancy with migrating focal seizures (EIMFS, previously migrating partial seizures in infancy [7]), a severe epileptic encephalopathy with onset in infancy. It was identified in two children who had hemiconvulsive seizures with an onset in the first weeks of life and died before 4 years of age [8].

We report here three additional patients from a multiplex family with variants in *SLC25A22* and a less severe phenotype of EIEE type 3 combined with functional studies. We suggest a delineation of the *SLC25A22*-related disorders group as well as a genotype–phenotype correlation.

Materials and methods

Clinical data

Patient one is a 15-year-old girl of Caucasian origin. She is the first child of unrelated and healthy parents. She had one healthy brother and one affected brother. She was born full term after a normal pregnancy with birth weight (BW) 3865 g (70th centile), birth length (BL) 53 cm (75th centile), and birth head circumference (BHC) 35 cm (50th centile). Epilepsy started on her 3rd day of life with clonic seizures followed by generalized myoclonic seizures, and hypertonic seizures, initially refractory to treatment. Seizures were transiently controlled with two medications from 5 years: levetiracetam and clobazam, but the generalized myoclonic seizures recurred at 11 years. The interictal EEG, at 3 months, was normal and, at 1 year, did not show SB but there was slow electrogenesis and sharp waves. From the first months of life, she had severe axial and peripheral hypotonia. At 1 year, she had a kinetic and static ataxia, an important axial hypotonia and frequent dyskinesia interfering with the gripping of objects. Behavioral disturbances with inconsolable crying, sleep disturbances, and selfmutilation disturbances started on 18 months. She had severe global developmental delay from the first months of life (head holding at 8 months, siting posture at 2 years, and crawl at 8 years). Her motor development deteriorated at 11 years, and she did not crawl at 13 years. At 15 years, she had severe global developmental delay, sleep disturbances, and mostly seizures. She had no language and could not sit still. Occipito-frontal circumference (OFC) is 54 cm (20th centile). Brain MRI, at 6 months, showed an expansion of peri-cerebral fluid spaces, then at 18 months, a corticalsubcortical atrophy predominant in frontal regions. A brain MRI control, at 13 years, did not find any anomalies. Heart and kidney ultrasound and visual evoked potentials (VEP) were normal. Array CGH (Agilent; Affymetrix snp6.0) did not detect any abnormality (arr(1-22,X)x2). Metabolic assays were within normal range, including blood count, liver enzymes, lactic acid blood and very long chain fatty acids blood concentrations, analysis for congenital disorders of glycosylation, and blood amino acids chromatography.

Patient 2 is the 10-year-old brother of patient 1. He was born full term after a normal pregnancy, with BW at 4570 g (98th centile), BL at 52 cm (50th centile), and BHC at 37 cm (96th centile). Epilepsy started at 1 month with generalized myoclonic and tonic-clonic seizures, also initially refractory to treatment. His seizures were partially controlled from 3 years, but the generalized myoclonic seizures recurred at 6 years and were treated by valproate, levetiracetam, and phenobarbital at 10 years. The interictal EEG, at 1 month, was normal and, at 1 year, showed slow electrogenesis but no SB. From the first months of life, he was severe axial and peripheral hypotonia and dyskinetic movements started. Behavioral disturbances with sleep disturbances started on 3 years. He was severely globally delayed. He held his head temporarily at 18 months, but he presented a motor regression after 6 years. He neither walked nor spoke. Physical examination at 9 years revealed axial and peripheral hypotonia. He had dystonic and dyskinetic movements, scoliosis, and bilateral cryptorchidism. His OFC was 52 cm (20th centile). Brain MRI showed increased anterior peri-cerebral spaces, at 3 months, and showed cerebral atrophy, at 18 months. Heart and kidney ultrasound and VEP were normal. Array CGH (Agilent; Affymetrix snp6.0) did not detect any abnormality (arr(1-22)x2,(X,Y)x1). Metabolic assays were normal, including blood count, liver enzymes, lactic acid blood and very long chain fatty acids blood concentrations, analysis for congenital disorders of glycosylation, and blood amino acids chromatography.

Patient 3 is patients 1 and 2's first-degree cousin: their mothers are sisters and their fathers are distant cousins (Fig. 1b). She is a Caucasian 5-year-old girl, the only child of healthy unrelated parents. She was born full term after a normal pregnancy. She had a BW at 3770 g (50th centile), a BL at 51 cm (50th centile), and a BHC at 35 cm (50th centile). Her seizures started aged 6 weeks with generalized myoclonic seizures, clonic seizures, and cyanotic episodes of uncertain cause. Epilepsy was controlled by clobazam and valproate from 2 years. The interictal EEG, at 4 years, showed a poor and disorganized background and several sharp waves. She had severe global hypotonia, developmental delay, and sleep disturbances that appeared at 18 months and aggravated at 5 years, for which she has been taking melatonin from 18 months. At 5 years, she does not crawl and has no language and she starts to hold her head. Her weight was 19 kg (85th centile), her length was 110 cm (85th centile), and her OFC was 49 cm (18th centile). Brain MRI at 1 year was normal. STXBP1,

KCNQ2 targeted sequencing, and array CGH (Agilent; Affymetrix snp6.0) were normal (arr(1-22,X)x2).

Patients' features from this report and the literature are summarized in the Table 1.

Exome sequencing

DNA from patient 2 and his mother and patient 3 and her father were studied using exome sequencing (Centre National de Recherche en Génomique Humaine, Institut de Biologie Francois Jacob, CEA). After complete DNA quality control for each sample (duplicate quantification, evaluating DNA integrity/quality, and checking for absence of PCR inhibitors), genomic DNA (3 µg) was captured by using an in-solution enrichment method (Human All Exon v5-50 Mb, Agilent Technologies, Santa Clara, CA, USA). Library preparation and exome enrichment (~20.000 targeted genes) was performed automatically by using NGSx (Perkin Elmer, MA, USA) and Bravo (Agilent Technologies), respectively, according to the manufacturer's instructions (SureSelect, Agilent Technologies). After normalization and quality control, exome-enriched libraries were sequenced by using the Illumina HiSeq2000 system (Illumina, CA, USA) as paired-end 100-bp reads. Samples were sequenced as pools of four samples per lane, to obtain an average coverage of 70-80×, with at least 80% of the





Fig. 1 a Visualization of variants by integrative genomics viewer (IGV). The sequence is the complementary sequence of genomic DNA. The first two lines correspond to patients 2 and 3. They shared compound heterozygous variants NM_0246989.4: c.[813_814delTG];

[818G>A]. The third line corresponds to the father of patient 2 who carries the variant c.818G>A. The fourth line corresponds to the mother of patient 3 who carries the variant c.813_814delTG. **b** Pedigree

et al. 2009		Total 9F/9M	4 Caucasian/ 14 Arab	14 Parental consangui- nity (78%)	15/17 before	18/18 Hypotonia (100%); 18/18 developmental delay (100%)	8/11 (72%)	9/17 presenting myoclonic seizure (53%)		5/14 presenting SB (36%)	5/11 (45%)	2/8 (25%)	11/15 abnormal innaging (73%)	5/14 (36%)
Molinari	Severe	P. 18 M	Arab	Yes	s 2 weeks	Peripheral hypotonia and no psychomo- tor acquisition	Yes	Focal seizures	Unknown	SB	Normal	Unknown	Thin CC, subtrach- noid enlarge- ment, and arrophy	Normal
2-15];]	Severe	P. 17 F	Arab	Yes	First month	Hypotonia later spastic and no psychomo- tor acquisition	Yes	Erratic myoclonic and focal seizures	Unknown	SB	Normal	Unknown	Thin CC and brain atrophy	Normal
[patients 1 nere	Severe	P. 16 M	Arab	Yes	5 days	Hypotonia, no psychomo- tor acquisition, weak suction, and incomptee archaic reflexes	Yes	Focal, secondary generaliza- tion epileptic sp- asms	No	SB and hypsar- rhythmia	Abnormal	Abolition of macular and peripheral responses and progressive alteration	Cerebellar hypoplasia, abnormal CC gyration of temporo- parietal myelinisa- temporal po- temporal po- ken	Normal
al. 2005 reported	Severe	P. 15 F	Arab	Yes	Birth	Hypotonia and later spastic	Yes	Myoclonic	Unknown	SB	Abnormal conduction and retinal function	Abnormal	Subarachnoid enlargement	Normal
olinari et 1- 3 are	Severe	P. 14 M	Arab	Yes	Birth	Hypotonia	Yes	Myoclonic	Unknown	Unknown	Unknown	Unknown	unknown	Unknown
ttients (M Patients	Severe	P. 13 F	Arab	Yes	Birth	Hypotonia	Yes	Myoclonic	Unknown	Unknown	Abnormal visual nerve- conduction velocity	Normal	Brain atrophy	Unknown
lished pa ts 4–9]).	Severe	P. 12 M	Arab	Yes	Birth	Hypotonia	Yes	Myoclonic	Unknown	SB	Abnormal	Unknown	Brain atrophy	Unknown
f the pub 7 [patien	ISHM	P.11 F	Arab	Yes	2 weeks	Hypotonia and developme- ntal delay	Unknown	Migrating partial seizures	No	Independent focal spikes in a multifocal distribution	Abnormal	Normal	Delayed myelimation patter and diffuse thiming of the CC	Normal
pdates o t al. 201	ISdW	P.10 M	Arab	Yes	1 week	Hypotonia and developme- ntal delay	Unknown	Migrating partial seizures	Partially	Delta brush pattern, positive spikes, and high- voltage focal spikes	Normal	Normal	Normal	Normal
cluding u _]]; Reid e	Moderate	P.9 F	Caucasian	No	6 weeks	Develop- mental delay, hypotonia, poor head control, and no visual fixation	Unknown	Focal seizures and asymetric cl- onic	No	No SB and semidiscon- tinuous pattern	Unknown	Unknown	Normal	Hyperproli- naemia
E phenotype, ind atients 10 and 11	Moderate	P. 8 F	Arab	Yes	7 years	Developmental delay (ten words), hypetonia, hypemetropia and astignatism	Yes	Absence seizure and febrile illness	I	Nomal	Normal	Nomal	Temporal pole myelinacion delay, prominent cerebalar folia, and small splenium and small splenium	Normal
2014 [p:	Moderate	P. 7 F	Arab	Yes	First month	Develop- mental delay and global hypotonia	Unknown	Febrile ilhess seizures and brief and tonic seizures	No	Symmetri- cal diffuse irregular slow wave activity and no SB	Unknown	Unknown	Temporal pole myelination delay, prominent cerebellar folia, and splenium	Hyperproli- naemia
<i>LC25A22</i> uri et al.	Moderate	P. 6 M	Arab	Yes	16 weeks	Axial hypotonia, develop- mental stasis, and no head control	Unknown	Flexor spasm-like and tonic- clonic seizures	Unknown	Multifocal abnormal- ities and no SB	Unknown	Unknown	Unknown	Hyperproli- næmia
ndividuals with variants in S 4 [patients 17 and 18]; Pod	Moderate	P. 5 M	Arab	Yes	16 weeks	Axial hypotonia, develop- mental stasis, and no head control	Unknown	Flexor spasm-like and tonic- clonic seizures	Unknown	Multifocal abnormal- ities and no SB	Unknown	Unknown	Unknown	Hyperproli- naemia
	Moderate	P. 4 M	Arab	Yes	6 weeks	Severe developmental delay, global hypotonia, and dystonic movement	Unknown	Myoclonic and tonic-clonic seizures	Yes	Unknown	Unknown	Unknown	Frontotem- poral hypoplasia, globalty delayed myelination, promient folia, and splenium	Hyperproli- naemia
	Moderate	P. 3 F	Caucasian	No	6 weeks 2 month-	Hypotonia and severe developme- ntal delay	No	Myoclonic and cyanotic seizures	Yes	No SB, poor and disorga- nized background, and sharp waves	Unknown	Unknown	Normal	Unknown
tures of in et al. 201	Moderate	P. 2 M	Caucasian	No	1 month	Hypotonia, severe developmental delay, and dystonic and dyskinetic movement	No	Myoclonic	Yes	No SB and electrogenesis slow down	Normal	Normal	Subarachnoid enlargement	Normal
Clinical fee 6]; Cohen	Moderate	P. 1 F	Caucasian	No	3 days	Hypotonia and severe developmenta- I delay	No	Myoclonic	Yes	No SB, electrogenesis slow down, and sharp waves	Normal	Normal	Nomal	Normal
Table 1 ([patient 1] (Group	Patient Sex	Ethnic group	Parental consanguinity	Age of onset seizure	Neurological examination	Microcephaly	Seizure type	Response to AEDs	EEG	VEP	ERG	Brain imaging	Metabolic evaluation

Group	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate	ISHW	MPSI	Severe	Severe	Severe	Severe	Severe	Severe	Severe	
Mutation	c.818G>A; p. (Arg273Lys) c.813_814del- TG; p. (Ala272Ghf- s*144)	c.818G>A; p. (Arg273Lys) c.813_814del- TG; p. (Ala272Glnf- s*144)	c.818G>A; p. (Arg273- Lys) c.813_814- deITG; p. (Ala272GI- nfs*144)	c.166A>C; p. (Thr56Pro)	с.166А>С; р. (Ли:56Рю)	c.166A>C; p. (Thr56Pro)	c.886G>A; p. (Ala296Th- r)	c.886G>A; p.(Ala296Thr)	c.235G>A; p. (Glu79Lys) c.746T>A; p. (Val249- Glu)	c.328G>C; p. (Gly110Ar- g)	c328G>C; p. (Gly110Ar- g)	c.617C>T; p. (Pro206- Leu)	c.617C>T; p. (Pro206Leu)	c.617C>T; p. (Pro206Leu)	(Pro206Leu)	с.706СъТ; р. (Gly236Tr- р)	c.617C>T; p. (Pro206- Leu)	c.617C>T; - p. (Pro206- Leu)	
Death or vegetative state	No	No	°N	Unknown	Unknown	Unknown	No	No	No	Death at 14 months of life	Death at 4- year-old age	Unknown	Unknown	Death at 8- year-old age	Unknown	Vegetative state at 10- year-old age	Vegetative state at 6- year-old age	Vegetative 6/12 (50% state at 4- year-old age	(2)
P patient,	F female,	<i>M</i> male,	AEDs an	ntiepileptic	drugs, E.	EG electi	roenceph	alogram, VEP vis	sual evo	ked poter	ıtial, <i>ER</i> 0	3 electro	retinogran	ı, <i>SB</i> sup	pression 1	ourst, CC	corpus	callosum	

Table 1 (continued)

target nucleotides covered at 30×. Image analysis and base calling involved use of the Illumina Real Time Analysis Pipeline. Sequence-quality parameters were assessed daily during the 12 days of sequencing. The standard bioinformatics analysis of sequencing data was based on the Illumina pipeline (CASAVA1.8.2) to generate a FASTQ file for each sample.

Sanger sequencing

SLC25A22 was sequenced by direct Sanger sequencing to confirm next-generation sequencing results as described by Barat-Houari et al. (BDT3.1 technology in a ABI3130XL sequencer) [9] on a genomic DNA sample from probands and their relatives.

Cell respiration and enzyme activities [2]

Standard conditions were used to grow cultured skin fibroblasts [10]. Cells of patient 3 could grow on glutamine in the absence of glucose. Respiration and mitochondrial substrate oxidation in the presence of either glutamate or succinate were polarographically studied in fibroblast suspensions before and after permeabilization by digitonin [11]. Frozen cell pellets were thawed and used for spectrophotometric determination of respiratory chain [12] and glutamate dehydrogenase activities [13, 14].

Results

We identified using exome sequencing and confirmed by Sanger sequencing two variants in SLC25A22 shared by the three patients: NM_024698.4:c.[813_814delTG]; [818G>A], (p.[Ala272Glnfs*144];[Arg273Lys]) (Fig. 1a). Familial segregation shows that p.(Ala272Glnfs*144) is inherited from their mothers, whereas p.(Arg273Lys) is inherited from their fathers, who were found to be distantly related (Fig. 1b). The variant c. 813_814delTG is absent from the control populations database (dbSNP, ExAC, GnomAD). The frequency in GnomAD of the variant c.818G>A is $4.124e^{-6}$ (rs 1195505218). These two variants are classified "pathogenic" according to ACMG criteria (Supplementary Table 1). We have submitted the information of the c.813_814delTG and c.818G>A variants to ClinVar database in NCBI (www.ncbi.nlm.nih.gov/clinvar/; accession number: SCV000882718 and SCV000898499, respectively).

We investigated the oxidation of glutamate in cultured skin fibroblasts from patient 3 (Fig. 2) [2]. Under standard conditions, polarographic studies showed normal cell respiration by intact cells, indicating that glutamate does not represent a major respiratory substrate. However, after



Fig. 2 Respiration and mitochondrial substrate oxidation in cultured skin fibroblasts from one patient and one control. After polarographic measurement of intact cell respiration, digitonin-permeabilized cells were loaded with adenosine diphosphate (ADP) and amino oxyacetate. This compound inhibits the aspartate-amino transferase enzyme activity. A subsequent addition of glutamate allowed estimation of mitochondrial glutamate oxidation under phosphorylation conditions (i.e., presence of ADP). Note the lack of glutamate-triggered oxygen uptake in the patient's cells. A similar succinate oxidation rate was measured in control's and patient's cells. The dotted line corresponds to the speed of consumption of O2 by cells. After addition of glutamate, the speed rises by 41% in the control's cells and decreases by 12% in the patient's cells. Numbers along the traces are nmol O2 consumed per minute per milligram of protein. AOA = amino oxyacetate; Asp = aspartate; Glut = glutamate; α -KG = α -ketoglutarate; OAA = oxaloacetate. Experimental conditions were as described in the "Materials and methods" section

permeabilization of cell membranes by digitonin in the presence of amino oxyacetate, a specific inhibitor of aminoaspartate transferase, the patient's cells failed to oxidize glutamate normally in comparison with controls, whereas oxidation of another substrate (succinate) was normal in the patient's cells. These results provide evidence that the patient's cells show a clear defect in mitochondrial glutamate metabolism, which is consistent with the involvement of the variants in *SLC25A22* in the disease of the children.

Discussion

EIEE type 3 is a rare and severe genetic disorder due to variants affecting the functioning of protein *SLC25A22*. The SLC25A22 protein is highly expressed in brain, particularly in areas dedicated to motor coordination (red nuclei, olivary complexes, pontocerebellar nuclei, and substantia nigra). SLC25A22 is more abundant in astrocytes, where it controls glutamate uptake [15, 16]. The protein is localized in the inner mitochondrial membrane and catalyzes glutamate/H+ symporter for the transport of glutamate into the mitochondria. The absence of functional SLC25A22 protein would lead to the dysregulation of extracellular glutamate



Fig. 3 Visualization of variants in SLC25A22. **a** Linear visualization of patient's variants. Htz = heterozygote; Hmz = homozygote. **b** 2-D visualization of variants in SLC25A22 protein

and activation of the extrasynaptic glutamate receptors [16, 17]. This could impair the cAMP response element binding protein shut-off pathway that would be responsible for long-lasting protection against apoptosis to neural death induction [18], and consequently, inactivation of the prosurvival extracellular signal regulated kinase [19] and neuronal excitability [2].

The first patients reported in the literature either died during the first year of life or survived in a persistent vegetative state [2, 3, 5]. However, recently, Reid et al. described six additional patients with variants in *SLC25A22* and a milder phenotype [6]. To date, six different variants affecting function of protein have been described (Fig. 3a).

SLC25A22 is also involved in EIMFS [8], a severe developmental and epileptic encephalopathy with onset in early infancy and characterized by migrating partial seizures on EEG. The two patients, described by Poduri et al., presented a similar phenotype to EIEE type 3: they developed seizures between 1 and 2 weeks of life, hypotonia, severe developmental delay, and died before 4-year-old age. The brain MRI of the second patient revealed a delayed myelination pattern. The only difference between phenotypes concerns the type of seizures with partial seizures for EIMFS.

The patients described here present a EIEE type 3 phenotype with a milder course of the disease, like patients reported previously by Reid et al. [6], (absence of SB, some acquisitions, no microcephaly, normal VEP, and still alive at 15-year-old age for the oldest) (Table 1). The functional studies allowed us to confirm the involvement of *SLC25A22* gene in our family.

The identification of the genetic variations involved in the disease of the patients allowed us to propose a targeted research screening of the causative genetic variations for the direct family members and a complete screening of the *SLC25A22* gene for unrelated members of future unions.

We clustered the 18 patients and the variants they carried into three groups according to the severity of the *SLC25A22*-related disorders. In an attempt to identify genotype–phenotype correlations, we compared the variants according to the location depending on the protein domains (Table 2 and Fig. 3b).

The variants c.166A>C; p.(Thr56Pro), c.235G>A; p. (Glu79Lys), c.746T>A; p.(Val249Glu), c.813_814delTG; p.(Ala272Glnfs*144), c.818G>A; p.(Arg273Lys), and c.886G>A; p.(Ala296Thr), involved in a milder phenotype (patients 1–9 in Table 1), were located in exons 4, 5, 9, and

Common symtoms	Severe phenotype n=7	EIMFS n=2	Moderate phenotype n=9
Early-onset seizures	SB	MPS	Absence of SB
Myoclonic seizures/MPS	Untractable epilepsy	Untractable epilepsy	Tractable epilepsy
Severe hypotonia	No psychomotor acquisition	No psychomotor acquisition	Some acquisition
Severe developmental delay	Microcephaly	Unknown	No microcephaly
	Abnormal VEP/ERG	Normal VEP/REG	Normal VEP/ERG
	Died or in vegetative state	Died	Alive
	c.617C>T; p.(Pro206Leu) c.706G>T; p.(Gly236Trp)	c.328G>C; p.(Gly110Arg)	c.818G>A; p.(Arg273Lys) c.813_814delTG; p.(Ala272GInfs*144) c.166A>C; p.(Thr56Pro) c.886G>A; p.(Ala296Thr) c.235G>A; p.(Glu79Lys) c.746T>A; p.(Val249Glu)
(NG_023407.1)	Exon 8	Exon 6	Exons 4, 5, 9 and 10
	Patients 12 to 18	Patients 10 and 11	Patients 1 to 9

Table 2 Severe versus mild phenotype comparison



EIEE early infantile epilepsy encephalopathy, *EIMFS* epilepsy of infancy with migrating focal seizures, *SB* suppression burst, *MPS* migrating partial seizures, *VEP* visual evoked potential, *ERG* electroretinogram

10 (NG 023407.1), and were not in helical transmembrane domains except for the two of them: p.(Glu79Lys) and p. (Ala296Thr). However, in patient 9, the p.(Glu79Lys) variant was associated with the p.(Val249Glu) variant (c. [235G>A]; [746T>A]), which did not affect the helical domain (patient 9). The p.(Ala296Thr) variant was in the homozygous state in patients 7 and 8. These patients had atypical phenotypes with febrile and absence seizures. This variant was present once in the heterozygous state in GnomAD and ExaC database. In silico pathogenicity prediction by SIFT and PolyPhen2 was "damaging", but functional studies were not performed. This variant involved the last helical transmembrane domain (Fig. 3b). We speculate that its consequences on the threedimensional conformation and channel function are less severe than those of the other helical transmembrane domain variants. This should be tested by functional studies. The variants c.617C>T; p.(Pro206Leu) and c.706G>T; p.(Gly236Trp), involved in the severe phenotype (patients 12-18 in Table 1), were located in exon 8 (NG_023407.1), and affected the helical transmembrane domains, essential for the protein conformation [20]. The variant c.328G>C; p.(Gly110Arg) found in patients with severe EIMFS (patients 10 and 11 in Table 1) was located in exon 6 (NG 023407.1) and affected also a helical transmembrane domain. This could directly compromise the channel function and explain the more severe phenotype. The criteria of pathogenicity for each variant according to ACMG criteria are summarized in Supplementary Table 1.

In summary, we found that patients with two variants affecting function of protein localized in helical transmembrane domains were likely to present with severe phenotypes, whereas patients with at least one variant outside the helical domains presented milder phenotypes. This could explain the phenotypic difference among patients (severe versus mild, Table 2 and Fig. 3).

In conclusion, we defined three groups in *SLC25A22*related disorders: EIEE type 3 with severe outcome, EIMFS with severe outcome, and EIEE type 3 with milder outcome, and suggest an association between the location of the variants and the variability of the severity of the disease. Additional studies of patients with EIEE type 3 and EIMFS will be required to further delineate the full phenotypic range associated with *SLC25A22* variants and genotype–phenotype correlations.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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