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Neurol Genet
2016;2:e115; doi: 10.1212/
NXG.0000000000000115

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Neurology.org/ng

FHF1 (FGF12) EPILEPTIC ENCEPHALOPATHY

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Voltage-gated sodium channels (Na_v s) are mainstays of neuronal function, and mutations in the genes encoding CNS Na_v s ($Na_v1.1$ [*SCN1A*], $Na_v1.2$ [*SCN2A*], $Na_v1.3$ [*SCN3A*], and $Na_v1.6$ [*SCN8A*]) are causes of some of the most common and severe genetic epilepsies and epileptic encephalopathies (EE).¹ Fibroblast-growth-factor homologous factors (FHF) compose a family of 4 proteins that interact with the C-terminal tails of Na_v s to modulate the channels' fast, and long-term, inactivations.² *FHF2* mutation is a rare cause of generalized epilepsy with febrile seizures plus (GEFS+).³ Recently, a de novo *FHF1* mutation (p.R52H) was reported in early-onset EE in 2 siblings.⁴ We report 3 patients from unrelated families with the same *FHF1* p.R52H mutation. The 5 cases together frame the FHF1 R52H EE from infancy to adulthood. As discussed below, this gain-of-function disease may be amenable to personalized therapy.

Patient 1 (P1, table) is a 3-year-old boy. Convulsive seizures began on day 2 of his life and remain intractable, with frequent status epilepticus (SE) manifesting as generalized or right-sided facial seizures. His EEGs show slow backgrounds and multifocal epileptiform discharges (figure e-1 at Neurology.org/ng). Multiple antiepileptic drug (AED) regimens failed, but he is currently at his best on a combination of the ketogenic diet and medications indicated in the table. He has severe global developmental delay, is nonverbal, has poor visual and social interaction, just started rolling slightly and sitting with support, and is tube-fed. He has generalized hypotonia with head lag and does not track. He suffers from chronic constipation. MRI at onset was normal, and at 2 years revealed widened ventricles and pericerebral CSF spaces (figure e-2A).

Patient 2 (P2) is a 16-year-old girl with intractable seizures since age 6 weeks and frequent SE. Seizures included generalized tonic-clonic (GTC) and partial seizures with left-sided facial twitching and lip-smacking (figure e-3). Phenytoin has been part of her AEDs, and relatively efficacious, since age 6 years. Since age 7, she develops 24 hours of severe ataxia after every GTC, which then gradually improves.

Vagal nerve stimulation substantially improved seizure, and ataxia, intensities, and frequencies. MRI at 16 months was unremarkable but at 8 years showed cerebellar atrophy (figure e-2B). She suffers from chronic constipation. She has severe cognitive impairment with single words, has normal motor development, and ambulates. She needs help with all activities of daily living.

Patient 3 (P3) is an 18-year-old girl with intractable epilepsy since day 2 of her life with frequent SE. Seizures include leftward head deviation followed by generalized convulsion. EEGs were slow with multifocal spikes in infancy and Lennox-Gastaut–like in early childhood. She suffers from chronic constipation, hypohydrosis, and reduced lacrimation, suggesting autonomic dysfunction. She has moderate intellectual disability and can read simple books. She ambulates with a spastic circumductive gait and has substantial heel cord tightness. Current therapies (table) include vagal nerve stimulation. MRI at present shows bilateral mesial temporal sclerosis and mild prominence of cerebellar folia, findings not present in MRIs from early childhood (figure e-2C).

All 3 patients had the same de novo *FHF1* NM_004113.5:c.155G>A, p.R52H mutation detected by whole-exome or whole-genome sequencing and confirmed by Sanger sequencing, and no other relevant de novo change. The table summarizes their clinical features and those of the recently published⁴ original sib-pair. Salient features of the latter pair include neonatal-onset intractable epilepsy, profound intellectual disability, severe feeding difficulties, MRI initially unremarkable and subsequently exhibiting cerebellar atrophy, ataxia, and death in SE.⁴

Based on 5 patients, the core FHF1 R52H disease comprises neonatal-onset persistent intractable epilepsy and moderate-to-severe intellectual disability. Radiologically, neurodegeneration, especially cerebellar, is present, which, beyond a mutational consequence appears to be aggravated by the severity and frequency of SE. It may also be exacerbated by treatment of SE. All 4 patients with cerebellar atrophy, including the original 2, were on phenytoin (table), repeatedly loaded and subsequently chronically maintained because of relative success in SE

Table	Phenotypic features of 5 patients with FHF1 R52H epileptic encephalopathy				
	O1	O2	P1	P2	P3
Current age and sex	Died age 7 y, female	Died age 3.5 y, male	3 y, male	16 y, female (DDD patient 251978)	18 y, female
Age at first seizure	2 wk	4 wk	2 d	6 wk	2 d
Status epilepticus	Frequent	Frequent	Frequent	Frequent	Frequent
Seizure types	Tonic seizures	Tonic seizures	GTC; right facial twitching	Myoclonic; GTC; partial motor seizures, lip-smacking, left facial twitching	Partial motor seizures with retained consciousness; left versive seizures followed by GTC
Interictal EEG findings	High-voltage slow activity with multifocal epileptiform discharges	High-voltage slow activity with multifocal epileptiform discharges	High-voltage slow activity with multifocal epileptiform discharges	Slow background with bilateral, right more than left, epileptiform discharges	Slow background with bilateral, right more than left, epileptiform discharges
Ictal EEG findings	Generalized high-voltage spike, sharp wave and spike waves followed by long suppression of background		Onset of high-amplitude 1.5-2 Hz rhythmic activity over right posterior head region with secondary generalization		
Current AED	AED regimen included phenytoin	AED regimen included phenytoin	Levetiracetam, phenobarbital	Phenytoin, perampanel, VNS	Phenytoin, pregabalin, perampanel, VNS
Developmental delay/intellectual disability	Severe psychomotor retardation, nonverbal	Severe psychomotor retardation	Severe global developmental delay, nonverbal	Severe global developmental delay and intellectual disability, single words	Moderate intellectual disability, possible autism spectrum disorder
Cerebellar involvement	Cerebellar atrophy, ataxia	Cerebellar atrophy, ataxia	To date uninvolved	Episodic cerebellar ataxia from age 7 y, cerebellar atrophy on MRI from age 8	Mild cerebellar atrophy appearing in adolescence
Other neurologic abnormalities	Cerebral visual impairment	Poor visual contact	Cortical visual impairment		Heel cord tightness with spastic circumductive gait
Neurologic examination	Hypotonia, microcephaly	Hypotonia, microcephaly	Diffuse hypotonia with head lag		Normal tone, slightly hyperreflexic
Concomitant morbidities	Feeding difficulties	Feeding difficulties	Constipation and vomiting; feeding difficulties that necessitate tube feeding	Severe chronic constipation	Signs of autonomic dysfunction: hypohidrosis, reduced lacrimation; chronic constipation
MRI findings	Cerebellar atrophy at age 6 y	Cerebellar atrophy at age 3 y	Cortical atrophy between 2 mo and 2 y	Cerebellar atrophy at age 8 y	Bilateral mesial temporal sclerosis, more marked on the right; prominence of cerebellar folia
Current therapies	Unspecified but includes phenytoin (G. Buyse, MD, PhD, personal communication, April 5, 2016)	Unspecified but includes phenytoin (G. Buyse, MD, PhD, personal communication, April 5, 2016)	Ketogenic diet, levetiracetam, phenobarbital	VNS, phenytoin, perampanel	VNS, phenytoin, perampanel, pregabalin

Abbreviations: AED = antiepileptic drug; O = original 2 patients from reference 4; P = present 3 patients (P2 is from the DDD Study, ID# DDD-NGS259178); GTC = generalized tonic-clonic; VNS = vagal nerve stimulation; DDD = Deciphering Developmental Disorders.

management and prevention. It may be cautious to use alternative medications, where possible, until this putative phenytoin cerebellar iatrogenicity is clarified.

Siekierska et al.⁴ demonstrated in vitro and in vivo that R52H is a toxic gain-of-function mutation. To date, thousands of patients with EE have undergone exome sequencing. Our rare finding of an FHF1 mutation suggests that FHF1 R52H might be a mutation-specific disease. It is possible that future FHF1 mutations with similar effects will be identified, but these would be expected to act through similar

gain-of-function mechanisms. As such, methods to downregulate the R52H and related alleles, e.g., with allele-specific antisense oligonucleotides, could prove therapeutic in this catastrophic encephalopathy.

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Acknowledgment: The study has UK Research Ethics Committee's approval (10/H0305/83, granted by the Cambridge South REC, and GEN/284/12 granted by the Republic of Ireland REC). The research team acknowledges the support of the National Institute for Health Research through the Comprehensive Clinical Research Network, the Health Innovation Challenge Fund (grant HICF-1009-003), and the Wellcome Trust Sanger Institute (grant WT098051). BAM holds the University of Toronto Michael Bahen Chair in Epilepsy Research. The authors thank the Next Generation Sequencing/bioinformatics teams at McGill University, the Genome Quebec Innovation Center, the Réseau de Médecine Génétique Appliquée, and Drs. Ledia Brunga (DNA sample management); Dipayan Mitra (MRI); Ms. Gail Charlton (EEG); and Ms. Sylvia Dobrzeniecka (sequencing) for their efforts.

Study funding: This work was supported by Genome Canada, Genome Quebec, the Ontario Brain Institute, and the DDD Study. The DDD Study presents independent research commissioned by the Health Innovation Challenge Fund (grant HICF-1009-003), a parallel funding partnership between the Wellcome Trust and the Department of Health, and the Wellcome Trust Sanger Institute (grant WT098051). The views expressed in this publication are those of the author(s) and not necessarily those of the Wellcome Trust or the Department of Health.

Disclosure: Dr. Al-Mehmadi, Dr. Splitt, and Dr. Ramesh report no disclosures. Dr. DeBrosse receives funding from NIH 2U54 NS078059-04 and NIH U54NS078059-04; receives philanthropic funding from families of individuals with pyruvate dehydrogenase complex deficiency; has received funding for a trip from the Alternating Hemiplegia of Childhood Foundation; and has received royalty payments from Decision Support in Medicine, LLC. Dr. Wallis

reports no disclosures. Dr. Xia has been an employee of Baylor Genetics LLC. Dr. Yang reports no disclosures. Dr. Rosenfeld has served on the editorial boards of Prenatal Diagnosis and Molecular Syndromology; has been an employee of Signature Genomic Laboratories, PerkinElmer, Inc.; and has received research support from NIH/National Human Genome Research Institute. Dr. Cosette receives funding from UCB Pharmaceuticals, Genome Quebec, Genome Canada, CIHR, and the Savoy Foundation. Dr. Michaud receives funding from Genome Quebec and Genome Canada. Dr. Hamdan reports no disclosures. Dr. Campeau serves on the editorial boards of Scientific Reports and the Journal of Pediatric Genetics; has acted as a consultant for Alexion Pharmaceuticals and Biomarin; holds stock options in Illumina, Inc.; and receives funding from the CHU Sainte-Justine Research Center, the Canadian Institutes of Health Research, the Fonds de Recherche en Santé Québec, the Réseau de Santé Buccodentaire et Oseuse, and the Fondation du Grand Défi Pierre Lavoie. Dr. Minassian receives funding from Genome Canada and the Ontario Brain Institute; holds patents for diagnostic testing of the following genes: EPM2A, EPM2B, MECP2, and VMA21; and has received license fee payments and royalty payments for the aforementioned patents. Go to Neurology.org/ng for full disclosure forms. The Article Processing Charge was paid by the authors.

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Received May 4, 2016. Accepted in final form August 29, 2016.

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1. Steinlein OK. Mechanisms underlying epilepsies associated with sodium channel mutations. *Prog Brain Res* 2014;213:97–111.
2. Goldfarb M. Voltage-gated sodium channel-associated proteins and alternative mechanisms of inactivation and block. *Cell Mol Life Sci* 2012;69:1067–1076.
3. Puranam RS, He XP, Yao L, et al. Disruption of Fgf13 causes synaptic excitatory-inhibitory imbalance and genetic epilepsy and febrile seizures plus. *J Neurosci* 2015;35:8866–8881.
4. Siekierska A, Isrie M, Liu Y, et al. Gain-of-function FHF1 mutation causes early-onset epileptic encephalopathy with cerebellar atrophy. *Neurology* 2016;86:2162–2170.

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Neurol Genet 2016;2;

DOI 10.1212/NXG.0000000000000115

This information is current as of October 28, 2016

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