# High-resolution molecular genomic autopsy reveals complex sudden unexpected death in epilepsy risk profile

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### **SUMMARY**

Advanced variant detection in genes underlying risk of sudden unexpected death in epilepsy (SUDEP) can uncover extensive epistatic complexity and improve diagnostic accuracy of epilepsy-related mortality. However, the sensitivity and clinical utility of diagnostic panels based solely on established cardiac arrhythmia genes in the molecular autopsy of SUDEP is unknown. We applied the established clinical diagnostic panels, followed by sequencing and a high density copy number variant (CNV) detection array of an additional 253 related ion channel subunit genes to analyze the overall genomic variation in a SUDEP of the 3-year-old proband with severe myoclonic epilepsy of infancy (SMEI). We uncovered complex combinations of single nucleotide polymorphisms and CNVs in genes expressed in both neurocardiac and respiratory control pathways, including SCNIA, KCNAI, RYR3, and HTR2C. Our findings demonstrate the importance of comprehensive high-resolution variant analysis in the assessment of personally relevant SUDEP risk. In this case, the combination of de novo single nucleotide polymorphisms (SNPs) and CNVs in the SCNIA and KCNAI genes, respectively, is suspected to be the principal risk factor for both epilepsy and premature death. However, consideration of the overall biologically relevant variant complexity with its extensive functional epistatic interactions reveals potential personal risk more accurately.

**KEY WORDS:** Sudden unexpected death in epilepsy, Severe myoclonic epilepsy of infancy, Epileptic encephalopathy, Dravet syndrome, Gene, Risk, Molecular autopsy.

Children with epileptic encephalopathy and uncontrolled seizures are at increased risk for sudden unexpected death in epilepsy (SUDEP).<sup>1</sup> Yet, the clinical risk factors do not provide a pathogenic mechanism, and they are not strongly predictive of the individual mortality hazard.

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Ion channel genes that modulate cardiac, autonomic, and respiratory functions are prime molecular risk factors for SUDEP. The causative mechanistic link between epilepsy, arrhythmias, sudden death, and the most common LQT gene, the potassium channel *KCNQ1*, was originally demonstrated in transgenic mice<sup>2</sup> and subsequently clinically validated.<sup>3</sup>

Because many ion channel genes that are critical for the regulation of neurocardiac and neurorespiratory pacemaking are also expressed within brain networks underlying epilepsy, a potential number of novel SUDEP candidate genes extends beyond the cardiac *LQT* genes.<sup>4</sup> For example, the voltage-gated potassium channel *KCNA is* coexpressed in brain and vagus nerve, and *Kcna1*-null mice have seizures, cardiac arrhythmias, vagal hyperexcitability, and die prematurely.<sup>5,6</sup> Similarly, the voltage-gated sodium channel *SCNIA* is dually expressed in the brain and the cardiac sinoatrial node and ventricular myocytes.<sup>7</sup> *Scn1a*-deficient

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mice also show autonomic instability and seizure-driven vagal activation preceding sudden death,<sup>7</sup> paralleling the clinical observations in children with *SCN1A* mutations and severe myoclonic epilepsy of infancy (SMEI).<sup>8</sup> However, most SMEI patients do not die suddenly, suggesting the modulating influence of other candidates in the genetic background, beginning with ion channels themselves.

We identified a SUDEP patient who displayed multiple established clinical-pathologic risk factors for SUDEP, including pharmacoresistant epileptic encephalopathy of the SMEI spectrum,<sup>9</sup> recurrent periictal respiratory compromise, and a suspected cardioautonomic clinical phenotype. In order to comprehensively assess the SUDEP risk embedded within this SMEI phenotype, we designed and performed an extensive postmortem search for deleterious variants in candidate ion channel subunit genes that regulate excitability within neural cardiorespiratory regulatory pathways.

# **Methods**

The 11-month-old patient and his parents were recruited into the institutional review board–approved Ion Channels in Epilepsy Project at Baylor College of Medicine.<sup>10</sup> Genomic DNA prepared from blood lymphocytes was submitted for commercial diagnostic exome sequencing in five *LQT* genes—*KCNQ1*, *SCN5A*, *KCNH2*, *KCNE2*, and *ANK2* (Transgenomics); whole genome copy number variants (CNV) analysis at the Medical Genetics Laboratory at Baylor College of Medicine; exome sequencing of 237 ion channel genes<sup>10</sup>; and screening on a custom-designed Ion Channel Comparative Hybridization (ICCH) 4 × 44K microarray (Agilent Technologies, Santa Clara, CA, U.S.A.).<sup>11</sup> (See Data S1.)

# RESULTS

### Index case clinical report

The proband was a healthy, full-term Latin American male born to a G1P1 mother. At 4 months of age the child developed a prolonged, afebrile hemiclonic seizure that subsided spontaneously but was followed by cessation of respi-Cardiopulmonary resuscitation ration. (CPR) was administered by a family member and the child promptly and fully recovered. General physical and neurologic examinations, a head computerized tomography (CT) (Acquilion Toshiba 320, Toshiba America Medical Systems, Inc., Tustin, CA, U.S.A.); and electroencephalography (EEG) (24channel Nicolet/VIASYS, VIASYS Healthcare, Madison, WI, U.S.A.) were normal, and treatment was deferred. Within a month he started experiencing weekly, treatmentresistant hemiclonic seizures involving either side. Serial electroencephalography and brain magnetic resonance imaging (MRI) (Philips Achieva 1.5T, Philips, Andover, MA, U.S.A.) studies remained unremarkable. Karyotyping confirmed a normal male chromosomal pattern. Routine serum and cerebrospinal fluid (CSF) studies were repeatedly normal and a comprehensive diagnostic workup for inborn metabolic errors was noncontributory. His development remained normal. Detailed family history was positive for migraine headaches in the mother. An episode of elevated temperature of 100.8°F triggered the first generalized tonic-clonic seizure at 9 months. Treatment-resistant daily myoclonic jerks associated with loss of tone began at 11 months of age, and the EEG showed epileptiform bursts of frontocentrally dominant generalized 2-3 Hz abortive spike and slow wave activity. The monthly, prolonged partial seizures were associated with cyanosis and frequent secondary generalization. By 18 months, global developmental delay became evident and the clinical evolution led to the diagnosis of SMEI.<sup>12</sup> A cardiac murmur was noted during a follow-up visit. Routine electrocardiography (ECG) was unremarkable, but he was referred to a cardiologist for further evaluation. The proband was 3 years and 3 months old and in his usual state of health when he was found cyanotic and unresponsive in bed. Full autopsy showed only pulmonary congestion, a frequent finding in sudden death.<sup>13</sup> SUDEP was confirmed as the official cause of death.

### Integrated genomic analysis

Our initial search centered on the five principal *LQTS* genes. It showed inherited nonsynonymous single nucleotide polymorphisms (nsSNPs) of unknown clinical significance in *KCNH2* (*LQT2*), *SCN5A* (*LQT3*), and *KCNE1* (*LQT5*) (Table 1) and in the context of the *LQT* genotype (Table S1), it failed to reveal a plausible molecular diagnosis. We next evaluated the channel variant profile (channotype) of the proband through parallel Sanger sequencing of 237 ion channel genes.<sup>10</sup> This step confirmed the previously detected LQTS polymorphisms and additionally uncovered a maternally transmitted heterozygous ryanodine receptor 2 (*RYR2*) nsSNP Q2958R (rs34967813) that has been reported previously in association with catecholaminergic polymorphic ventricular tachycardia (CPVT)<sup>14</sup> (Table 1; Fig. S1A).

The known epilepsy genes—*SCN1A*, *KCNA1*, and *SCN8A*—have also been implicated in SUDEP.<sup>5,7,15–17</sup> Proband channotype analysis<sup>10</sup> revealed an inherited common polymorphism A1067T and a de novo nsSNP, A1783V in *SCN1A* (Table 1; Fig. S1B) previously found in SMEI (http://www.molgen.vib-ua.be/SCN1AMutations/Home). This predicted deleterious de novo mutation in our case sug-

gests a contribution to the epileptic encephalopathy phenotype, yet its influence on the lethality is uncertain. We also uncovered a paternally inherited, novel, nsSNP, C1288Y, in the *RYR3* gene that is preferentially expressed in hippocampus and smooth muscle cells of the pulmonary artery,<sup>18,19</sup> and the animal models support its role in learning, cognition, and in hypoxia-induced pulmonary vasoconstriction. Therefore, a dysfunctional *RYR3* channel could contribute

#### **Genetic SUDEP Risk Profile in SMEI**

Table I. Nonsyn	nonymou	s single nucle	otide polymo compareo	orphisms in candic d to parental profi	late genes for SUDEP in les	the proband	i (IE124)
Syndrome	GENE	PROTEIN	Polyphen	SIFT	IE124p (dbSNP)	IE125m	IEI26f
Cardiac LQT Gene SNP Sequencing (gene dosage: heterozygous = 1; homozygous = 2)							
LQT2 LQT3 LQT5 CPVT	KCNH2 SCN5A KCNEI RYR2	hERG/KvII.I NavI.5 MinK RyR2	Tolerated Tolerated Tolerated	Benign Benign Benign Probably damaging	K897T (2) (rs1805123) H558R (1) (rs1805124) S38G (1) (rs1805127) Q2958R (1) (rs34967813)	K897T (2) H558R (1) S38G (2) Q2958R (1)	K897T (I)  
Human Epilepsy Gene Sequencing (gene dosage: heterozygous = 1; homozygous = 2)							
ADNFLE IGE DEND	CHRNA2 CLCN2 KCNJI I	nAChR2 CLC-2 Kir6.2	Tolerated Tolerated Tolerated	Benign Benign Benign	T125A (1) (rs891398) T668S (1) (rs9820367) K23E (1); (rs5219) V337I (1) (rs5215)	T125A (I) T668S (I) -; -	T125A (2) T668S (2) K23E (2); V337I (2)
Dravet/SMEI/GEFS+	SCNIA	Nav1.1	Tolerated; Deleterious	Benign; Probably damaging (benign)	A1067T (1) (rs2298771); A1783V (1) (rs121917980)	A1067T (I); _	A1067T (I);
Respiratory Serotonin Receptor Gene Sequencing (gene dosage: (heterozygous = 1; homozygous = 2)							
N/A N/A	HTR3C HTR3D	5-HT3C 5-HT3D	Tolerated Tolerated; Tolerated	Benign Benign; Possibly damaging (benign)	G405A (1) (rs6807362) G36A (2) (rs6443930); R260H (2) (rs6789754)	G405A (2) G36A (1); R260H (2)	_ G36A (I); R260H (I)

to the cognitive impairment and respiratory compromise of our patient, and targeted *RYR3* analysis in SMEI cohorts will be essential to validate this assumption.

Given the clinical history of recurrent, seizure-related apnea, we also analyzed genetic variation in all 18 of the known 5-HT ligand gated ion channels (*HTR1A-F*, *HTR2A-C*, *HTR3A-E*, *HTR4*, *HTR5A*, *HTR6*, and *HTR7*) and found three inherited nsSNPs (Table 1), of which only the *R260H* variant is predicted to be possibly damaging by Sorting Intolerant From Tolerant ([SIFT] sift.jcvi.org) SIFT.

Given the modulating role of genetic background on clinical phenotype, we also examined the whole genome for structural gene rearrangements. The clinical aCGH screen identified eight inherited autosomal copy number changes in the proband, such as the paternally inherited duplication in *SLC6A10P*, a gene recently implicated in autistic spectrum disorder,<sup>20</sup> and the recurrent deletion at 15q11.2, which was found previously in excess in children with congenital heart defects<sup>21</sup> (Table 2). Because all eight CNVs were inherited and their pathogenic relevance to epilepsy or SUDEP was uncertain, we applied our custom high resolution custom designed Ion Channel Comparative Hybridization (ICCH) Array, which has minimal detection threshold of 50 bp and an ultra-dense coverage across the exome of 253 ion channel genes, their structurally related family members, and known accessory subunits. Eleven novel duplications in nine known SUDEP genes were confirmed by quantitative polymerase chain reaction (qPCR; Table 2). Duplication size ranged from to 60 to 3,059 bp. Four CNVs were de novo. Two rearrangements were independent gains in RYR2, and one was a duplication in GABRG3. They were restricted to introns. The single coding de novo CNV, confirmed by qPCR, was at the 3' end of exon 2 in KCNA1 (Fig. 1A–C), a gene encoding the Kv1.1 pore forming alpha subunit, the loss of function of which causes severe epilepsy and SUDEP in animal models<sup>5</sup> (Fig. 1C). Normalization with two reference genes revealed that the proband harbored five extra copies of this exonic region as compared to the diploid genomes of both parents (Fig. 1D). This gain has a direct impact on the protein coding sequence of the KCNA1 gene. It extends from the highly conserved proline hinge motif (Pro-X-Pro) to the end of the S6 transmembrane helix of the Kv1.1 subunit (Fig. 1E). The PVP motif in this membrane-spanning helix forms a flexible hinge in the

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Table 2. Copy n	umber va	ariation ir	n SUDEP	proband	(IE I 24) co char	ompare nnel co	ed to pai mparat	rental pr ive hybri	ofiles usii dization	ng a clini array	ical diagn	ostic microarray ar	nd a custom	ı high-der	isity ion
CNV no.	Chromoson	ne CytoBi	Start and (H	position Er Ig19)	1d position (Hg 19)	Gain/Lo	Num ss of pro	ber Leng bes Cl	gth of K NV O	NV* 0	Number of genes	RefSeq (HUGO) gene names	In proband (IE124)	ln mother (IE125)	In father (IEI 26)
Clinical aCGH Diagnostic Array (BCM Molecular Diagnostics Core – Director Dr. Ankia Patel)															
-	chrl	p21.1	104(	012051 14	04012498	Loss		8	447 N	(1,2,4)			λ	ч	~
2	chr4	q12	57,	746415	57988228	Gain		9 24	1813 N	(1,2,3)			×	и	Y
с С	chr8	p11.23	39	369942	39499498	Gain		8 12	9556 N	(1,2,3)	2 A	DAM5P, ADAM3A	~	Y	~
4	chrl 0	q11.22	.94	384979	4650680 I	Gain		4 12	1822 N (	l ,2,3,4)	S S	YT I 5,GPRIN2,PPYR I	Y	Y	Y
5	chrl4	q11.2	216	\$09644	22028409	Loss	-	4	8765 N	(1,2,3)			~	×	~
9	chrl4	q11.2	18	864561	19459230	Gain		3 59	4669 N	(1,2,3)	6 P	704P, OR4Q3, OR4M1,	٨	×	×
7	chrl 5	q11.2	19	108763	19464920	Loss	=	3 35	6157 N(	l ,2,3,4)	3 17	UR4N2, UR4N2, UR4N3 DC646214, CXADRP2, POTER	×	×	~
ω	chr16	p11.2	32,	481308	33528443	Gain	01	6 104	7135 N(	1,2,3,4)	5	NF267, HERC2P4, LOC729355, TP53TG3, SLC6A10P	~	c	~
CNV no.	Chrc	omosome	CytoBand	Start position (Hg18)	End posi (Hg18	s) G	ain/Loss	Number of probes	Length of CNV	Known CNV*	Number of genes	RefSeq (HUGO) gene names	In proband (IE124)	In mother (IE125)	In father (IE I 26)
Ion Channel Compar. Hybridization (ICCF Custom High Densii aCGH Array (Translational Neurogenetics in Epilepsy Laboratory - Directo Dr. Alica Goldman)	- H) ty or														
_	chr l		q43	23587908	4 235879	141 D	ain	-	60	z	-	RYR2 (intron 49/50) NM_001035.2	~	Ē	c
2	chr l		q43	23602830	4 236028(	634 G	ain	c	330	N (1,2)	-	RYR2 (intron 97/98) NM 001035.2	~	c	c
m	chr l		q43	1986462	I 19864(	666 G	ain	2	45	z	-	HTR6 (exon I 5' UTR) NM_000871.1	~	~	c
4	chr7		q36.1	I 5028590	8 150285	968 G	ain	_	60	N (1,2)	-	KCNH2 (intron 4/5) isol NM 000238.3	~	Ē	~
ß	chr l	2	p13.33	211716	3 2123	823 G	ain	Μ	6660	N (1,2)	-	CACNA / C (intron 2/3) iso   NM_   99460.2	×	۲	~
															Continued

					Table 2	. Continu	.pər						
CNV no.	Chromosome	CytoBand	Start position (Hg18)	End position (Hg18)	Gain/Loss	Number of probes	Length of CNV	Known CNV*	Number of genes	RefSeq (HUGO) gene names	In proband (IE124)	In mother (IE 125)	In father (IE126)
9	chr12	P13.32	4892174	4892251	Gain	2	11	z	_	KCNAI (exon 2): NM_000217.2	~	۲	<u>د</u>
7	chr I 5	q12	24568642	24569014	Gain	m	372	N (I)	_	GABRB3 (exon 2-3) iso1 NM_000814.5	×	~	×
8	chr I 5	q12	25052658	25055717	Gain	7	3059	N (2)	-	GABRG3 (intron 3/4) iso1 NM 033223.4	٨	Ę	c
6	chr 19	P13.13	13178788	13179169	Gain	m	381	z	_	CACNAIA (exon 47 3' UTR) isol NM 000068.3	~	×	~
10	chr 19	p13.13	13478014	l 3478059	Gain	_	45	N (2)	-	CACNAIA (exon I) NM_000068.3	λ	≻	~
=	chr 19	p13.3	565234	565928	Gain	Ω	694	N (1,2)	_	HCN2 (intron 3/4) NM_001194.3	×	٨	~
*Known aberrations To	ronto Database of C	Genomic Varian	ts (dgv.tcag.ca)	(I = region ha	s reported gai	in; 2 = region	has reporte	d loss; 3 = 1	egion has r	eported indels; $4 = region$	n has reported	inversions).	

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transmembrane domain and is directly involved in channel function. Mutations in this region have previously been shown to cause epilepsy (V408T),<sup>22</sup> and premature C-terminal truncation or deletion of the Kv1.1 gene leads to aberrant protein expression resulting in epilepsy, ataxia, megalencephaly, and SUDEP in mice.<sup>23</sup> The repeated gain of this transmembrane helix in the Kv1.1 subunit is likely to impact protein packing and lipid membrane insertion, and thus is an attractive candidate mechanism for Kv1.1 dysfunction contributing to both the seizure and SUDEP phenotype of the proband.

# DISCUSSION

As the list of validated risk genes for SUDEP expands beyond those currently linked to cardiac-related mortality, robust diagnostic platforms must be developed for optimal assessment of integrated genetic risk.

Herein we show that constructing the genetic variation risk profile for SUDEP benefits from complementary, comprehensive, candidate ion channel gene-focused detection platforms. Both single base pair substitutions and architectural defects contribute to the risk of epilepsy and SUDEP, as evidenced by the discovery of two biologically plausible pathogenic de novo variants in known SUDEP candidates: SCN1A and KCNA1. Mutations in both genes play a critical role in autonomic destabilization described in clinical reports<sup>16,24</sup> and experimental models of SUDEP,<sup>5,7</sup> and likely contributed to lethality in our patient. Yet, the co-occurrence of epileptic encephalopathy, ictal apnea, suspected cardiac compromise, and SUDEP in this patient may not be explained solely by the molecular mechanisms elucidated through the SCN1A and KCNA1 models,<sup>5,7</sup> but may also reflect the compound effect of these mutations together with the transmitted nsSNPs and CNVs of the cardiac arrhythmia and serotonin receptor genes, RYR3 gene variant, and the 15q11.2 region variant associated with structural heart defects. Because clinical phenotypes reflect the pattern of both the individually unique (de novo) and inherited ion channel variants<sup>10</sup> (Fig. S2), resolving the full genetic context is essential for accurate assessment of risk. The integration of ion channel exome sequencing, high resolution ion channel specific CNV survey, and subsequent analysis of 54 candidate SUDEP genes in the neuro-cardiac-respiratory network in this case shows the need for multiscale channel-based risk prediction for SUDEP.

We present the first comprehensive genomic interrogation of ion channel candidate gene pathways to dissect and personalize SUDEP risk prediction in pediatric patients with epilepsy. This case harbored combination of de novo SNPs and CNVs in the *SCN1A* and *KCNA1* genes potentially acting as the principal risk factors for premature death. The larger complexity of the risk load was revealed by additional inherited structural rearrangements and missense

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#### Figure 1.

A de novo gain in the human epilepsy and SUDEP gene KCNA1 was identified in the proband. (A) Chromosomal location of the human KCNA1 gene using Hg18 as the reference genome. The region of the detected genomic gain is in the gray box with the probe positions located beneath the coding exon. (B) Higher magnification view of the 3' end of the KCNA1 gene showing the region of the gain relative to the Ion Channel Comparative Hybridization (ICCH) microarray probes. The qPCR primers TLK1822 and TLK1823 used to validate the CNV are shown where primer TLK1823 overlaps with the CGH probe. (C) Sybr green standard curve shown against known concentrations of human gDNA to establish qPCR assay efficiency (91.3%). All qPCR assays underwent optimization and efficiency analysis prior to validation experiments in proband genomic DNA. (D) Quantification of the gain in the KCNA1 gene showed five additional copies of this region in the proband and not in either parent. Normalization of genomic copy number was performed using two reference genes that are known to be free of copy number variants and compared to a normal diploid control. (E) Homology model of the human Kv1.1 ion channel subunit showing two opposing subunits in the tetrameric channel. The ribbon is colored the same as the 6TM schematic diagram top right for orientation. The S4 voltage sensor is gray with the positively charged arginine and lysine residues shown in red. The S6 domain is shown in red and the region of gain is in black. The S6 PVP hinge sequence is highly conserved from jellyfish to man. The residues V404 and V408 are shown in black where amino acid substitutions at these positions cause epilepsy, ataxia, and myokymia. *Epilepsia* (C) ILAE

polymorphisms within the clinically evident neurocardiac and respiratory pathways. As we continue to refine our understanding of the specific biologic pathways and genetic risk factors leading to SUDEP, comprehensive assessment of genomic variation in cardiac and respiratory networks using detailed gene profiling can enhance predictive value

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of gene testing in the routine neurologic care of individuals with epilepsy.

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# DISCLOSURE

J.R.L. is a paid consultant for Athena Diagnostics, has stock ownership in 23 and Me and Ion Torrent Systems, and is a co-inventor on multiple United States and European patents related to molecular diagnostics for inherited neuropathies, eye diseases, and bacterial genomic fingerprinting. The Department of Molecular and Human Genetics at Baylor College of Medicine derives revenue from the chromosomal microarray analysis (CMA) and clinical exome sequencing offered in the Medical Genetics Laboratory (MGL; http://www.bcm.edu/geneticlabs/). The remaining authors have no potential conflicts of interest. We confirm that we have read the Journal's position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

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# SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

**Table S1.** Clinical and functional consequences of common polymorphisms in the proband affected by SMEI and SUDEP.

**Figure S1.** The SUDEP proband inherited multiple missense variants in the candidate genes for SUDEP.

**Figure S2.** Personal channotype profiles of single nucleotide polymorphisms and copy number variants in the candidate genes for SUDEP.

Data S1. Methods.