

# Cardiac Channel Molecular Autopsy: Insights From 173 Consecutive Cases of Autopsy-Negative Sudden Unexplained Death Referred for Postmortem Genetic Testing

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## Abstract

**Objective:** To perform long QT syndrome and catecholaminergic polymorphic ventricular tachycardia cardiac channel postmortem genetic testing (molecular autopsy) for a large cohort of cases of autopsy-negative sudden unexplained death (SUD).

**Methods:** From September 1, 1998, through October 31, 2010, 173 cases of SUD (106 males; mean  $\pm$  SD age, 18.4 $\pm$ 12.9 years; age range, 1-69 years; 89% white) were referred by medical examiners or coroners for a cardiac channel molecular autopsy. Using polymerase chain reaction, denaturing high-performance liquid chromatography, and DNA sequencing, a comprehensive mutational analysis of the long QT syndrome susceptibility genes (*KCNQ1*, *KCNH2*, *SCN5A*, *KCNE1*, and *KCNE2*) and a targeted analysis of the catecholaminergic polymorphic ventricular tachycardia type 1-associated gene (*RYR2*) were conducted.

**Results:** Overall, 45 putative pathogenic mutations absent in 400 to 700 controls were identified in 45 autopsy-negative SUD cases (26.0%). Females had a higher yield (26/67 [38.8%]) than males (19/106 [17.9%];  $P < .005$ ). Among SUD cases with exercise-induced death, the yield trended higher among the 1- to 10-year-olds (8/12 [66.7%]) compared with the 11- to 20-year-olds (4/27 [14.8%];  $P = .002$ ). In contrast, for those who died during a period of sleep, the 11- to 20-year-olds had a higher yield (9/25 [36.0%]) than the 1- to 10-year-olds (1/24 [4.2%];  $P = .01$ ).

**Conclusion:** Cardiac channel molecular autopsy should be considered in the evaluation of autopsy-negative SUD. Several interesting genotype-phenotype observations may provide insight into the expected yields of postmortem genetic testing for SUD and assist in selecting cases with the greatest potential for mutation discovery and directing genetic testing efforts.

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Sudden cardiac death (SCD) is a major cause of death in developed countries. An estimated 300,000 to 400,000 individuals die suddenly each year in the United States, with most deaths involving elderly people.<sup>1</sup> In comparison, sudden death in infants, children, adolescents, and young adults is relatively uncommon, with an incidence between 1.3 and 8.5 per 100,000 patient-years.<sup>2</sup> Nevertheless, an estimated 1000 to 5000 individuals between 1 and 35 years of age die suddenly each year. Fortunately, the cause and manner of death can be explained in many cases by a comprehensive medicolegal investigation that includes an autopsy.<sup>3,4</sup>

A conventional autopsy investigation may detect a noncardiac basis for the sudden death, such as pulmonary embolism, asthma, or epilepsy. However, SCD is the most common cause of sudden death in young people, with structural cardiovascular abnormalities often identifiable at autopsy.<sup>4,5</sup> Nevertheless, standard forensic autopsy

investigations often fail to reveal the underlying cause of the SCD. In fact, even after gross and histologic examination, at least 3% and perhaps as much as 53% of sudden deaths involving previously healthy children, adolescents, and young adults have no identifiable morphological abnormalities found at autopsy, remain unexplained, and are classified as autopsy-negative sudden unexplained death (SUD).<sup>3,4,6-8</sup>

Long QT syndrome (LQTS) and catecholaminergic polymorphic ventricular tachycardia (CPVT) are potentially lethal, heritable channelopathies associated with structurally normal hearts that leave no evidentiary clue to be gleaned during a comprehensive medicolegal autopsy. This absence of evidence leaves medical examiners, coroners, and forensic pathologists to only surmise that a lethal arrhythmia may have precipitated the SUD.<sup>3,9-13</sup> However, postmortem genetic testing, specifically a cardiac channel molecular autopsy, may potentially elucidate such a pathogenic mechanism and estab-

lish probable cause and manner for an individual's death.<sup>14-17</sup> In fact, recent guidelines for autopsy investigations of SUD in young people have suggested that postmortem cardiac channel genetic testing should become the new standard of care in the evaluation of SUD cases.<sup>18-20</sup>

Since the availability of preliminary case reports of molecular autopsies,<sup>15</sup> investigators have sought to determine the spectrum and prevalence of pathogenic cardiac ion channel mutations in the now 9 unique molecular autopsy series of SUD cases, providing a range of 15% to 35% mutation detection yield when testing for the 3 major LQTS susceptibil-

ity genes (*KCNQ1*, *KCNH2*, *SCN5A*) and/or the major CPVT-associated gene (*RYR2*).<sup>17,21-28</sup> However, these 9 molecular autopsy series, now totaling only 207 cases (49 of which we have reported on previously), mostly represent very small case series (<20 cases), used a largely unreliable source (formalin-fixed paraffin-embedded tissue) of high-quality DNA for comprehensive mutational analysis, or performed a limited mutational analysis (ie, only assessed a partial list of the major LQTS and CPVT genes).

Given the recent guidelines for autopsy-negative SUD investigations and the relatively small co-

**TABLE 1. Clinical Characteristics of the Autopsy-Negative SUD Cohort<sup>a</sup>**

Characteristic	All cases	LQTS-positive cases	CPVT I-positive cases	Negative cases
No. of individuals	173	25	20	128
Sex, No.				
Male	106	8	11	87
Female	67	17	9	41
Age at SUD (y), mean ± SD (range)				
Overall	18.4±12.9 (1-69)	17.6±9.4 (2-43)	16.5±10.2 (2-36)	18.9±13.9 (1-69)
Females	18.8±13 (1-52)	18.1±10.4 (4-43)	18.9±11 (5-35)	19.3±14 (1-52)
Males	18.2±13 (1-69)	16.6±7.2 (3-24)	14.5±9.6 (2-36)	18.8±14 (1-69)
Reported ethnicity, No.				
White	154	24	16	113
Black	9	1	3	6
Hispanic	6	0	1	5
Asian	3	0	0	3
Unknown	1	0	0	1
Events at SUD				
Sleep	70 (40.5)	12 (48.0)	1 (5.0)	57 (44.5)
Nonspecific	52 (30.1)	6 (24.0)	8 (40.0)	38 (29.7)
Exertion	46 (26.6)	5 (20.0)	11 (55.0)	30 (23.4)
Auditory	1 (0.6)	1 (4.0)	0	0
Emotion	4 (2.3)	1 (4.0)	0	3 (2.3)
Personal history before SUD				
Positive	42 (24.3)	8 (32.0)	7 (35.0)	27 (21.1)
Negative	124 (71.7)	15 (60.0)	13 (65.0)	96 (75.0)
Unknown	7 (4.0)	2 (8.0)	0	5 (3.9)
Family history of cardiac events				
Positive	34 (19.7)	5 (20.0)	7 (35.0)	22 (17.2)
Negative	127 (73.4)	19 (76.0)	12 (60.0)	96 (75.0)
Unknown	12 (6.9)	1 (4.0)	1 (5.0)	10 (7.8)
Family history of SCD	20 (11.5)	2 (8.0)	7 (35.0)	11 (8.6)
Personal or family history				
Seizures	12 (6.9)	1 (4.0)	2 (10.0)	9 (7.0)
Syncope	25 (14.5)	9 (36.0)	4 (20.0)	12 (9.4)

<sup>a</sup>Data are presented as No. (percentage) unless indicated otherwise. CPVT I = catecholaminergic polymorphic ventricular tachycardia type I; LQTS = long QT syndrome; SUD = sudden unexplained death.

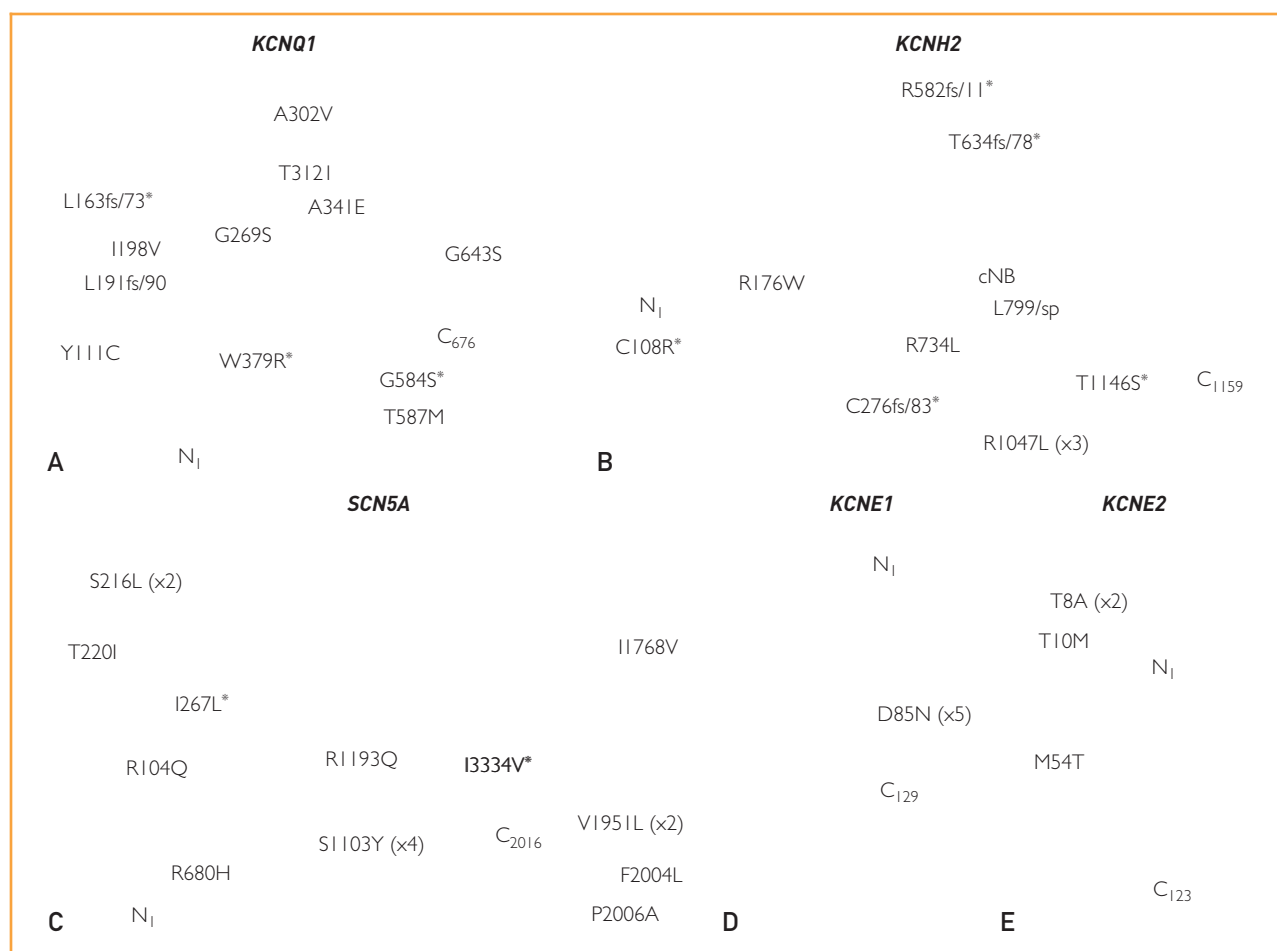
hort size of these previously published molecular autopsy series, an extensive analysis to better define the expected yield of mutation detection and perhaps offer possible genotype-phenotype correlations that may assist in guiding phenotype-directed cost-effective mutation detection efforts in future cases of SUD is much needed. We report the spectrum and prevalence of possible LQTS- and CPVT-associated channel mutations among 173 consecutive cases of SUD that were referred by medical examiners, coroners, and forensic pathologists from throughout North America during the past decade for a cardiac channel molecular au-

topsy. In an effort to guide and inform the proper use of such postmortem genetic testing, we examined the effect of different phenotypic parameters on the yield of LQTS and CPVT genetic testing from the largest molecular autopsy series on SUD published to date.

## METHODS

### Medical Examiner–Referred Cases of Autopsy-Negative SUD

From September 1, 1998, through October, 31, 2010, 173 medical examiners' cases of autopsy-negative SUD



**FIGURE 1.** Summary of cardiac channel mutations in long QT syndrome (LQTS)-associated genes identified in a series of autopsy-negative sudden unexplained death (SUD). The putative, pathogenic SUD-associated mutations (gray circles) and additional nonsynonymous, functional polymorphisms (white circles) identified in this study are depicted with their proximate location on the linear topologies (not drawn to scale) of the LQT1-associated *KCNQ1*-encoded cardiac Kv7.1/I<sub>Ks</sub> potassium channel  $\alpha$ -subunit (A), the LQT2-associated *KCNH2* encoded cardiac Kv11.1/I<sub>Kr</sub> potassium channel  $\alpha$ -subunit (B), the LQT3-associated *SCN5A* encoded cardiac Nav1.5/I<sub>Na</sub> sodium channel  $\alpha$ -subunit (C), the LQTS-associated *KCNE1*-encoded cardiac Kv7.1 potassium channel  $\beta$ -subunit (D), and the LQTS-associated *KCNE2*-encoded cardiac Kv11.1 potassium channel  $\beta$ -subunit (E). Asterisk indicates novel mutation absent in the published literature. The numbers within parentheses represent the number of times the variant was seen in cases. For example, R1047L (x3) is a functional polymorphism seen in 3 cases.

(106 males; mean  $\pm$  SD age,  $18.4 \pm 12.9$  years; range, 1-69 years) were referred to Mayo Clinic's Windland Smith Rice Sudden Death Genomics Laboratory for a cardiac channel molecular autopsy. After receipt of written consent from the decedent's appropriate next of kin for this Mayo Foundation Institutional Review Board-approved protocol, a postmortem genetic mutational analysis (ie, cardiac channel molecular autopsy) was performed. The first 49 of these 173 consecutively referred cases have been reported previously<sup>22,29</sup> and are included here in this extended molecular autopsy series. None of the decedents or their family members had an established diagnosis of LQTS, CPVT, or any other sudden death-predisposing cardiac condition at the time of either their death or at enrollment for postmortem genetic testing.

### Cardiac Channel Molecular Autopsy

Genomic DNA was extracted from frozen necropsy tissue or autopsy blood using the Puregene DNA Isolation Kit (Qiagen Inc, Valencia, CA). Comprehensive coding, open reading frame, splice-site mutational analysis of the entire coding region (60 exons) of the 3 major LQTS susceptibility genes (*KCNQ1*, *KCNH2*, and *SCN5A*), 2 minor LQTS susceptibility genes (*KCNE1* and *KCNE2*), and a targeted molecular screen of the 64 exons (3-28, 36-50, and 83-105) previously reported to encompass the 3 major mutation-clustering domains of the 105 exon CPVT type 1 (CPTV1) susceptibility gene *RYR2*<sup>30</sup> was performed using polymerase chain reaction, denaturing high-performance liquid chromatography, and direct DNA sequencing as previously described.<sup>31</sup>

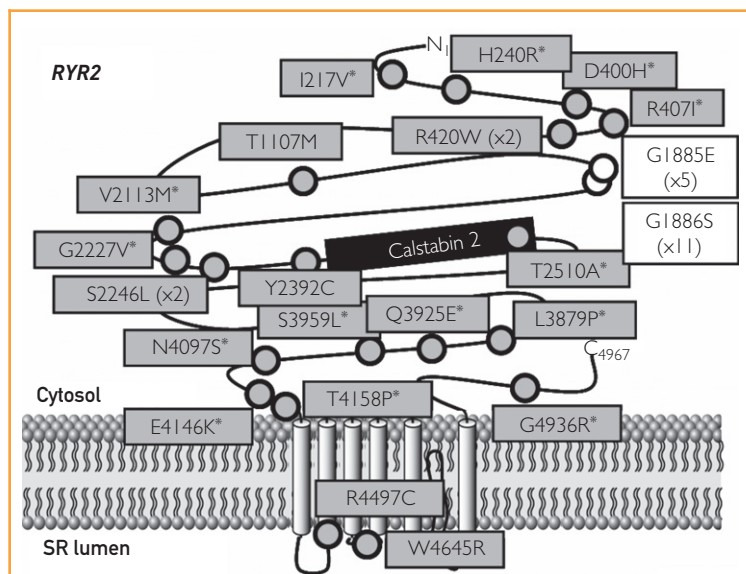
To be considered a putative, pathogenic, SUD-associated mutation, the genetic variant had to (1) be a nonsynonymous variant (synonymous single-nucleotide polymorphisms were excluded from consideration); (2) involve a highly conserved residue; and (3) be absent from either more than 2600 reference alleles previously examined for LQTS mutations or more than 800 reference alleles previously examined for *RYR2* mutations<sup>32-34</sup> and an additional 400 reference alleles derived from 100 healthy white and 100 healthy black controls. This latter control genomic DNA was obtained from the Human Genetic Cell Repository sponsored by the National Institute of General Medical Sciences and the Coriell Institute for Medical Research (Camden, NJ). Mutations were annotated using the single-letter nomenclature whereby W379R, for example, denotes a nonsynonymous single-nucleotide polymorphism producing a missense mutation involving a substitution of tryptophan (W) by arginine (R) at amino acid W379R. To assess whether mutations were sporadic or familial, direct DNA sequencing

was used to confirm the presence or absence of the putative mutations in DNA extracted from the parents of the decedents whenever possible and after written informed consent. Primer sequences and polymerase chain reaction and denaturing high-performance liquid chromatography conditions are available on request.

## RESULTS

### Cohort Description

Demographic characteristics for this consecutively referred SUD cohort are given in Table 1. Briefly, there were 173 cases (106 males and 67 females; mean  $\pm$  SD age,  $18.4 \pm 12.9$  years; range, 1-69 years; 89% white). The mean  $\pm$  SD ages of the males ( $18.2 \pm 13$  years; range, 1-69 years) and females ( $18.8 \pm 13$  years; range, 1-52 years) were similar. Most (161/173 [93.1%]) were 40 years or younger at their SUD. The circumstance (events or triggers) surrounding the SUD was grouped into 1 of 3 categories, with the most prevalent being death during sleep ( $n=70$ , 30 males [40.5%]), followed by a non-



**FIGURE 2.** Summary of cardiac channel mutations in the catecholaminergic polymorphic ventricular tachycardia type 1 (CPVT1)-associated *RYR2*-encoded calcium release channel (cardiac ryanodine receptor)  $\alpha$ -subunit. Asterisk indicates novel mutation. The numbers within parentheses represent the number of times the variant was seen in cases.

TABLE 2. Summary of Putative Channelopathic Mutations and Channelopathy-Positive SUD Cases<sup>a</sup>

Case No.	Sex	Age (y)	Ethnicity	Gene	Exon	Nucleotide change	Mutation (amino acid change)	Nonsynonymous polymorphisms	SUD event	Sentinel event	Family history
1	M	3	W	<i>KCNQ1</i>	1	332 A>G	Y111C		Sleep	SUD	Positive for syncope
2	F	21	W	<i>KCNQ1</i>	3	del488T	L163FS/X236 <sup>b</sup>	T8A-KCNE2	Nonspecific	Syncope <sup>c</sup>	Negative
3	M	17	W	<i>KCNQ1</i>	3	del572_576	L191FS/90		Sleep	SUD	Negative
4	M	20	W	<i>KCNQ1</i>	3	592 A>G	I198V		Sleep	SUD	Negative
5	F	43	W	<i>KCNQ1</i>	6	805 G>A	G269S		Exertion	SUD	Negative
6	M	22	W	<i>KCNQ1</i>	6	905 C>T	A302V	G1886S-RYR2	Sleep	Seizures	Negative
7	F	8	W	<i>KCNQ1</i>	7	935 C>T	T312I		Exertion	SUD	Positive for SCD <sup>c</sup>
8	M	24	W	<i>KCNQ1</i>	7	1022 C>A	A341E		Sleep	SUD	Negative
9	F	17	W	<i>KCNQ1</i>	9	1135 T>C	W379R <sup>b</sup>		Emotion	SUD	Negative
10	F	5	W	<i>KCNQ1</i>	15	1750 G>A	G584S <sup>b</sup>		Exertion	Syncope	Negative
11	F	4	W	<i>KCNQ1</i>	15	1760 C>T	T587M		Nonspecific	SUD	Negative
12	F	14	W	<i>KCNH2</i>	3	322 T>C	C108R <sup>b</sup>	V1951L-SCN5A	Nonspecific	SUD	Positive for SCD
13	F	29	W	<i>KCNH2</i>	4	del823T	C276fsX359 <sup>b</sup>		Nonspecific	SUD <sup>c</sup>	Positive for syncope
14	F	15	W	<i>KCNH2</i>	7	insGC1746_1747	R582FS/I1 <sup>b</sup>		Sleep	Syncope	Negative
15	F	16	W	<i>KCNH2</i>	7	del1902C	T634FS/78 <sup>b</sup>		Sleep	SUD	Negative
16	F	13	W	<i>KCNH2, RYR2</i>	9, 28	2398+5 G>T, 3321 C>T	L799/SP, T1107M-RYR2		Auditory	Syncope	Negative
17	F	15	W	<i>KCNH2</i>	15	3436 A>T	T1146S <sup>b</sup>		Sleep	Syncope	Negative
18	F	16	W	<i>SCN5A</i>	3	311 G>A	R104Q		Sleep	SUD	Negative
19	F	19	W	<i>SCN5A</i>	6	659 C>T	T220I		Nonspecific	SUD	Negative
20	M	12	W	<i>SCN5A</i>	7	799 A>C	I267L <sup>b</sup>		Nonspecific	SUD	Negative
21	M	23	B	<i>SCN5A</i>	14	2039 G>A	R680H	S1103Y-SCN5A	Exertion	SUD	Negative
22	F	39	W	<i>SCN5A</i>	23	4000 A>G	I3334V <sup>b</sup>		Sleep	Syncope	Positive for syncope and CA
23	F	16	W	<i>SCN5A</i>	28	5302 A>G	I1768V		Sleep	SUD	Negative
24	M	12	W	<i>KCNE2</i>	2	29 C>T	T10M	D85N-KCNE1,	Exertion	Syncope	Negative
25	F	18	W	<i>KCNE2</i>	2	161 T>C	M54T		Sleep	SUD	Negative
26	M	21	W	<i>RYR2</i>	9	649 A>G	I217V <sup>a</sup>	G1885E-RYR2, G1886S-RYR2,	Nonspecific	Syncope	Positive for SCD

(Continued on next page)

TABLE 2. Continued

Case No.	Sex	Age (y)	Ethnicity	Gene	Exon	Nucleotide change	Mutation (amino acid change)	Nonsynonymous polymorphisms	SUD event	Sentinel event	Family history
27	M	2	W	<i>RYR2</i>	10, 90	719 A>G, 12472 A>C	H240R, <sup>b</sup> T4158P <sup>b</sup>		Exertion	SUD	Negative
28	F	35	W	<i>RYR2</i>	14	1198 G>C	D400H <sup>b</sup>		Nonspecific	SUD	Positive for SCD
29	M	9	H	<i>RYR2</i>	14	1220 G>T	R407I <sup>b</sup>		Nonspecific	Syncope	Negative
30	M	17	W	<i>RYR2</i>	14	1258 C>T	R420W		Exertion	Syncope	Negative
31	F	22	W	<i>RYR2</i>	14	1258 C>T	R420W		Nonspecific	SUD	Positive for SCD
32	M	36	W	<i>RYR2</i>	41	6337 G>A	V2113M <sup>b</sup>		Exertion	SUD	Negative
33	M	16	W	<i>RYR2</i>	43	6680 G>T	G2227V <sup>b</sup>		Nonspecific	SUD	Negative
34	M	5	W	<i>RYR2</i>	44	6737 C>T	S2246L	G643S-KCNQ1, D85N-KCNE1	Exertion	SUD	Negative
35	M	4	B	<i>RYR2</i>	44	6739 C>T	S2246L		Exertion	SUD	Negative
36	F	25	W	<i>RYR2</i>	47	7175 A>G	Y2392C	R1047L-KCNH2	Exertion	SUD	Positive for SCD
37	F	17	W	<i>RYR2</i>	50	7528 A>G	T2510A <sup>b</sup>		Nonspecific	SUD	Negative
38	M	17	W	<i>RYR2</i>	86	11636 T>C	L3879P <sup>b</sup>		Exertion	Syncope	Negative
39	F	9	W	<i>RYR2</i>	87	11773 C>G	Q3925E <sup>b</sup>		Exertion	Seizures	Negative
40	F	15	W	<i>RYR2</i>	88	11876 C>T	S3959L <sup>b</sup>		Exertion	SUD	Negative
41	M	18	W	<i>RYR2</i>	90	12290 A>G	N4097S <sup>b</sup>		Nonspecific	SUD	Positive for SCD
42	M	14	W	<i>RYR2</i>	90	12436 G>A	E4146K <sup>b</sup>		Sleep	SUD	Positive for SCD
43	F	34	B	<i>RYR2</i>	93	13610 C>T	R4497C		Nonspecific	SUD	Positive for SCD
44	F	5	W	<i>RYR2</i>	96	13933 T>A	W4645R		Exertion	SVT	Negative
45	F	8	W	<i>RYR2</i>	104	14803 G>A	G4936R <sup>b</sup>		Exertion	Seizures	Negative

<sup>a</sup>B = black; CA = cardiac arrest; F = female; H = Hispanic; M = male; SCD = sudden cardiac death; SUD = sudden unexplained death; SVT = sustained ventricular tachycardia; W = white.

<sup>b</sup>Novel to this cohort.

<sup>c</sup>Postpartum period.

specific event (n=52, 27 males [30.1%]) and exertion (n=46, 30 males [26.6%]). In addition, 1 individual (0.6%) experienced an auditory-triggered sudden death, and 4 (2.3%) died suddenly during extreme emotion. The distribution of triggers or events was not statistically different between males and females, respectively (sleep, 45.3% [48/106] vs 32.8% [22/67],  $P=.11$ ; nonspecific, 25.5% [27/106] vs 37.3% [25/67],  $P=.12$ ; exertion, 28.3% [30/106] vs 23.9% [16/67],  $P=.60$ ).

Importantly, no decedent or relative had received a clinical diagnosis of a suspected cardiac channelopathy or heritable arrhythmia syndrome (ie, LQTS or CPVT) before the SUD or at the time necropsy tissue was received to commence the molecular autopsy. However, either a positive personal

or family history of syncope, seizures, cardiac arrest, near drowning or unexplained drowning (in a family member), or prolonged QT interval was documented explicitly by the medical examiner in 70 of the 173 cases (40.5%). A personal or family history of syncope (n=25 [14.4%], personal history in 20) or seizures (n=13 [7.5%], personal history in 11) was identified. A family history of premature SCD before 50 years of age, occurring before the current patient's SUD, was reported in 20 cases (11.6%). A personal or family history of a cardiac event occurring during the postpartum period was reported in 6 cases. Status concerning personal history was not available in 7 cases (4.0%), and family history information was unavailable in 12 cases (6.9%).

**TABLE 3. Summary of Additional SUD Patients Hosting Only a Common Functional, Nonsynonymous Channel Polymorphism<sup>a</sup>**

Case No.	Sex	Age (y)	Ethnicity	Gene	Nonsynonymous polymorphism	SUD event	Sentinel event	Family history
46	M	27	W	<i>KCNE1</i>	D85N-KCNE1	Nonspecific	SUD	Positive for CA
47	M	17	W	<i>KCNE1</i>	D85N-KCNE1	Sleep	Syncope	Negative
48	M	46	W	<i>KCNE1</i>	D85N-KCNE1	Nonspecific	SUD	Negative
49	F	43	W	<i>KCNE2</i>	T8A-KCNE2	Sleep	SUD	Negative
50	F	30	W	<i>KCNH2</i>	R176W-KCNH2	Nonspecific	Syncope <sup>b</sup>	Negative
51	M	15	W	<i>KCNH2</i>	R1047L-KCNH2	Sleep	SUD	Positive for SCD
52	M	20	W	<i>KCNH2</i>	R1047L-KCNH2, S216L-SCN5A	Sleep	SUD	Negative
53	M	21	W	<i>SCN5A</i>	S216L-SCN5A	Sleep	SUD	Negative
54	M	1	B	<i>SCN5A</i>	S1103Y-SCN5A	Sleep	SUD	Negative
55	F	2	B	<i>SCN5A</i>	S1103Y-SCN5A	Sleep	SUD	Negative
56	M	15	H	<i>SCN5A</i>	S1103Y-SCN5A	Emotion	SUD	Negative
57	M	19	W	<i>SCN5A</i>	R1193Q-SCN5A	Sleep	Syncope	Negative
58	M	18	H	<i>SCN5A</i>	V1951L-SCN5A	Sleep	SUD	Negative
59	M	21	W	<i>SCN5A</i>	F2004L-SCN5A	Nonspecific	SUD	Negative
60	M	1	W	<i>SCN5A</i>	P2006A-SCN5A	Sleep	SUD	Negative
61	F	16	W	<i>RYR2</i>	G1885E-RYR2	Emotion	Seizures	Negative
62	M	38	W	<i>RYR2</i>	G1885E-RYR2	Sleep	SUD	Negative
63	M	19	B	<i>RYR2</i>	G1885E-RYR2	Nonspecific	SUD	Negative
64	M	12	W	<i>RYR2</i>	G1885E-RYR2	Exertion	Syncope	Positive for syncope
65	M	2	W	<i>RYR2</i>	G1886S-RYR2	Sleep	SUD	Negative
66	M	12	W	<i>RYR2</i>	G1886S-RYR2	Exertion	SUD	Positive for arrhythmias
67	F	12	W	<i>RYR2</i>	G1886S-RYR2	Exertion	Syncope	Negative
68	M	18	W	<i>RYR2</i>	G1886S-RYR2	Exertion	SUD	Positive for arrhythmias
69	M	18	W	<i>RYR2</i>	G1886S-RYR2	Nonspecific	SUD	Negative
70	F	4	W	<i>RYR2</i>	G1886S-RYR2	Exertion	Syncope	Negative
71	F	8	W	<i>RYR2</i>	G1886S-RYR2	Nonspecific	SUD	Negative
72	M	30	W	<i>RYR2</i>	G1886S-RYR2	Sleep	SUD	Negative
73	M	12	W	<i>RYR2</i>	G1886S-RYR2	Sleep	SUD	Negative

<sup>a</sup>B = black; CA = cardiac arrest; F = female; H = Hispanic; M = male; SCD = sudden cardiac death; SUD = sudden unexplained death; W = white.

<sup>b</sup>Postpartum period.

### Spectrum and Prevalence of Rare, Putatively Channelopathic Mutations in SUD

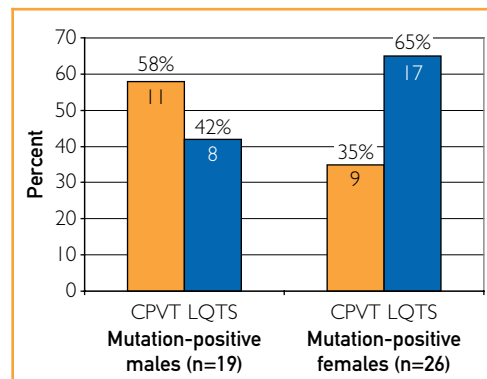
Overall, 45 rare, putative pathogenic mutations (20 *RYR2* [44.5%, 14 novel], 11 *KCNQ1* [24.5%, 4 novel], 6 *KCNH2* [13.3%, 5 novel], 6 *SCN5A* [13.3%, 2 novel], and 2 *KCNE2* [4.4%]) (Figures 1 and 2, and Table 2) were identified in 45 cases (26.0%) of autopsy-negative SUD. Most of the mutations (n=40) were missense, with only 5 (12.5%) radical mutations (4 frameshift [2 *KCNQ1*, 2 *KCNH2*] and 1 splicing error [*KCNH2*]). Two decedents hosted multiple mutations (case 16, L799sp-*KCNH2*, T1107M-*RyR2*; case 27, H240R-*RyR2*, T4158P-*RyR2*). Mutations involving LQTS-associated genes (*KCNQ1*, *KCNH2*, *SCN5A*, and *KCNE2*) were identified in 25 cases (14.5%) overall compared with 21 cases (12.1%) hosting mutations involving the CPVT-associated *RYR2* gene.

Among the 11 mutation-positive SUDS cases (of 45) for which the families chose to participate in genetic pedigree analysis, the SUD-associated mutation was established as a familial mutation in every family (4 *RYR2*, 4 *KCNQ1*, 1 *KCNH2*, 1 *SCN5A*, and 1 *KCNE2*) despite having no family history of cardiac events in 10 of 11 cases (Table 2). One of the decedents had a family history of SCD in a paternal uncle. The SUD was the sentinel event in 8 of the 11 families. Among this small subset of participating families, 3 of the decedents had a personal history of syncope.

Besides these aforementioned rare putative channelopathic mutations, several common nonsynonymous, functional polymorphisms, previously associated with LQTS, Brugada syndrome, drug-induced torsades de pointes, or ventricular arrhythmias or SCD, were also identified (Figure 1 and Tables 2 and 3), including G1886S-*RyR2*<sup>35,36</sup> (11 cases), G1885E-*RyR2*<sup>36</sup> (5 cases), D85N-*KCNE1*<sup>37</sup> (5 cases, all males), S1103Y-*SCN5A*<sup>38,39</sup> (4 cases), R1047L-*KCNH2*<sup>40</sup> (3 cases), V1951L-*SCN5A*<sup>41</sup> (2 cases), S216L-*SCN5A*<sup>41,42</sup> (2 cases), T8A-*KCNE2*<sup>43</sup> (2 cases), R176W-*KCNH2*<sup>44</sup> (1 case), G643S-*KCNQ1*<sup>45</sup> (1 case), R1193Q-*SCN5A*<sup>46,47</sup> (1 case), F2004L-*SCN5A*<sup>41</sup> (1 case), and P2006A-*SCN5A*<sup>41,48</sup> (1 case). Common polymorphisms with a heterozygous frequency greater than 20%, including K897T-*KCNH2*, H558R-*SCN5A*, G38S-*KCNE1*, and Q2958R-*RyR2*, and less common polymorphisms lacking functional data to suggest proarrhythmic susceptibility (DUP 52-54 API-*KCNQ1* [1 case], K393N-*KCNQ1* [1 case], P448R-*KCNQ1* [1 case], V648I-*KCNQ1* [1 case], R148W-*KCNH2* [1 case], R18W-*SCN5A* [1 case], S1787N-*SCN5A* [1 case], A1136V-*RyR2* [3 cases], and R4037C-*RyR2* [1 case]) have been excluded from Figure 1 and Tables 2 and 3.

Most of these polymorphisms do not appear to be overrepresented within this SUD cohort. However, the common proarrhythmic and sudden death-associated *SCN5A* polymorphism S1103Y<sup>38</sup> with an expected heterozygote frequency of 13% among healthy black controls was present in 3 of 9 black (33.3%) and 1 of 6 Hispanic (16.7%) SUD patients. The drug-induced LQT- and arrhythmia-associated D85N-*KCNE1* polymorphism was identified in 5 of 154 (3.2%, all male) white SUD patients compared with the expected prevalence of 1.0% in the general white population.<sup>33</sup> Interestingly, although only 40.5% (70/173) of the overall cohort died during sleep, 77.8% (7/9) putative, pathogenic mutation-negative SUD patients, who were identified as having an *SCN5A* functional polymorphism (S216L, S1103Y, R1193Q, V1951L, F2004L, and P2006A), died during sleep ( $P=.04$ ).

Importantly, although these proarrhythmic channel polymorphisms may have contributed to the patient's death, we excluded them from the calculations of SUD molecular autopsy yield and from the genotype-phenotype correlative studies detailed below. Accordingly, the 26.0% prevalence of channelopathy-mediated sudden death in this autopsy-negative SUD cohort could be underestimated.



**FIGURE 3.** The percent distribution of mutations identified in the catecholaminergic polymorphic ventricular tachycardia (CPVT)-associated *RYR2* gene compared with mutations identified in the long QT syndrome (LQTS)-associated genes for the 19 mutation-positive males and the 26 mutation-positive females. The numbers in the bars represent the numbers of cases with a mutation. For example, 11 of 19 (57.9%) mutation-positive males had a CPVT-associated *RYR2* mutation compared with 8 males (42.1%) who had a mutation in an LQTS-associated gene (*KCNQ1*, *KCNH2*, *SCN5A*, *KCNE1*, or *KCNE2*).



### Genotype-Phenotype Correlations

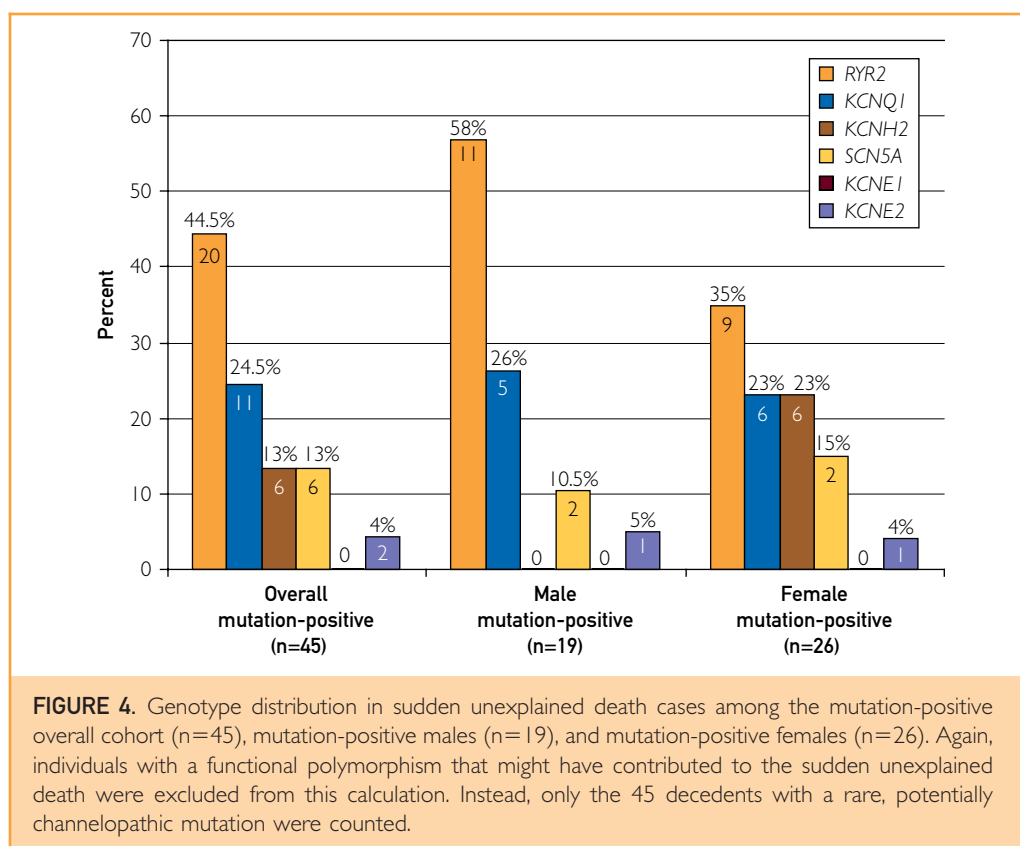
**Effect of Sex and Age on Yield.** Female SUD patients (26/67, 38.8%) were more likely to host a channel mutation than male patients (19/106, 17.9%;  $P=.004$ ). Moreover, mutation-positive females were more likely to host an LQTS-associated mutation rather than a CPVT1-associated *RYR2* mutation (65.4% [17/26] LQTS vs 34.6% [9/26] CPVT), whereas mutation-positive males were more likely CPVT1 positive (57.9% [11/19]) than LQTS mutation positive (42.1% [8/19], Figure 3). In fact, the distribution of genes hosting mutations was different between females and males (Figure 4). Specifically, although 57.9% (11/19) of the mutations seen in males were in *RYR2*, only 34.6% (9/26) of the mutations in females involved *RYR2*. In females, 23.1% (6/26) of the mutations identified involved *KCNH2*, whereas 0 of 19 male SUD patients had a *KCNH2* mutation ( $P=.03$ ).

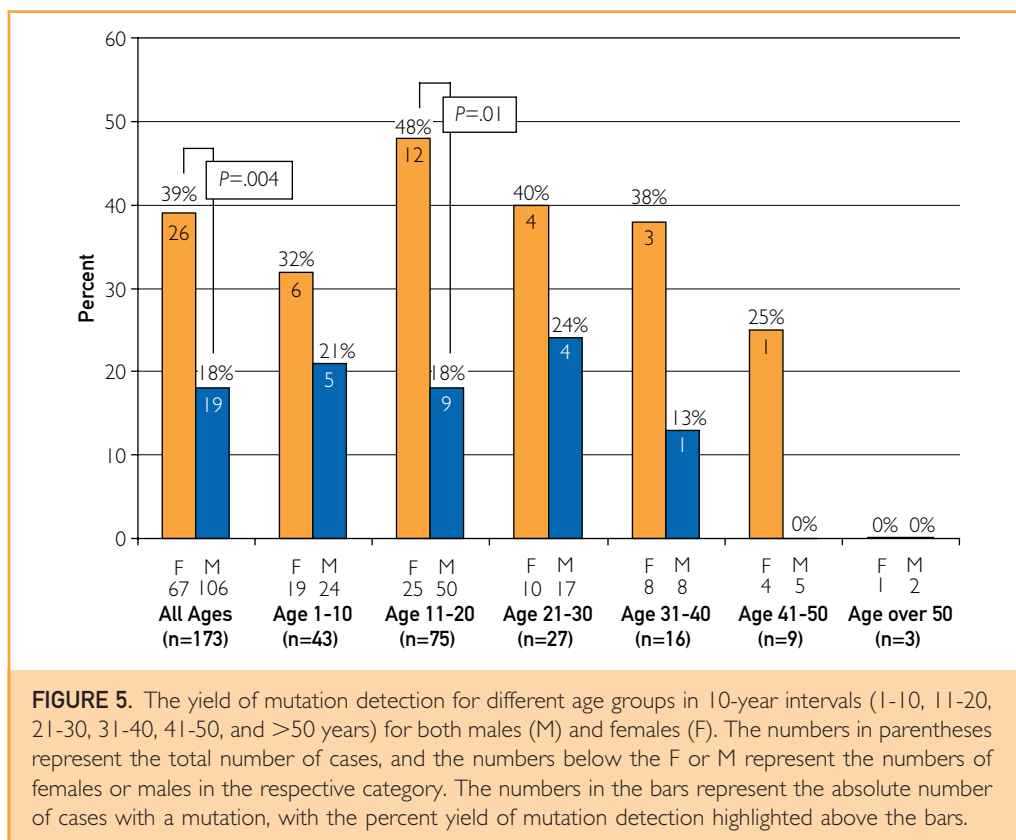
The yield of the molecular autopsy was not different between SUD patients who were 1 to 20 years of age (32/118, 27.1%) compared with decedents older than 20 years (13/55, 23.6%;  $P=.71$ ; Figure 5). Albeit small in number, none of the 3 SUD cases (1 female and 2 male) who were older than 50 years at the time of death were mutation positive. At all age groups, a trend

was seen toward a higher mutation detection yield in females, with the highest differential occurring in the 11- to 20-year range (12/25, 48.0% in females vs 9/50, 18.0% in males;  $P=.01$ ).

**Effect of Event or Trigger on Yield.** The mutation detection yield was associated with the circumstance (events or triggers) surrounding the death, with the highest yield occurring in those with a documented exertional trigger (34.8% [16/46]) compared with a nonspecific trigger (26.9% [14/52]) or death during a period of sleep (18.6% [13/70], Figure 6). This trend held true for females and males, with females consistently having a higher yield than males within the 3 event or trigger categories (exertion, 50.0% [8/16] vs 26.7% [8/30],  $P=.19$ ; nonspecific trigger, 36.0% [9/25] vs 18.5% [5/27],  $P=.21$ ; sleep, 31.8% [7/22] vs 12.5% [6/48],  $P=.09$ ). Again, however, because of the relatively small sizes in each of these event or trigger subsets, these sex and event or trigger trends did not achieve statistical significance.

Although cases of exercise-related SUD had the highest mutation detection rate overall, an interesting age effect emerged when examining the yield in specific triggered event categories in different age





groups. Overall, SUD patients aged 1 to 10 years with an exertion-induced death had a mutation detection yield (8/12, 66.7%) that was significantly higher than that of 11- to 20-year-olds with exertion-induced death (4/27, 14.8%;  $P=.002$ ). In contrast, for those with death during a period of sleep, the 11- to 20-year-olds had a higher yield (9/25, 36.0%) than the 1- to 10-year-olds (1/24, 4.2%;  $P=.01$ ; Figure 7).

The mutation-positive gene distribution among females (62.5% [5/8] *RYR2* and 37.5% [3/8] *KCNQ1*) and males (75.0% [6/8] *RYR2*, 12.5% [1/8] *SCN5A*, and 12.5% [1/8] *KCNE2*) with exertion was similar and somewhat anticipated based on previous studies showing *KCNQ1* LQTS susceptibility mutations and *RYR2* CPVT susceptibility mutations most often associated with exertion. Interestingly, although the gene distribution among females (42.9% [3/7] *SCN5A*, 42.9% [3/7] *KCNH2*, and 14.3% [1/7] *KCNE2*) with SUD during sleep correlated with previous observations that suggested that *KCNQ1* and *RYR2* mutations are infrequently associated with periods of rest and that *SCN5A* mutations are associated more frequently with cardiac events during rest, the male gene distribution was strikingly different from the female distribution

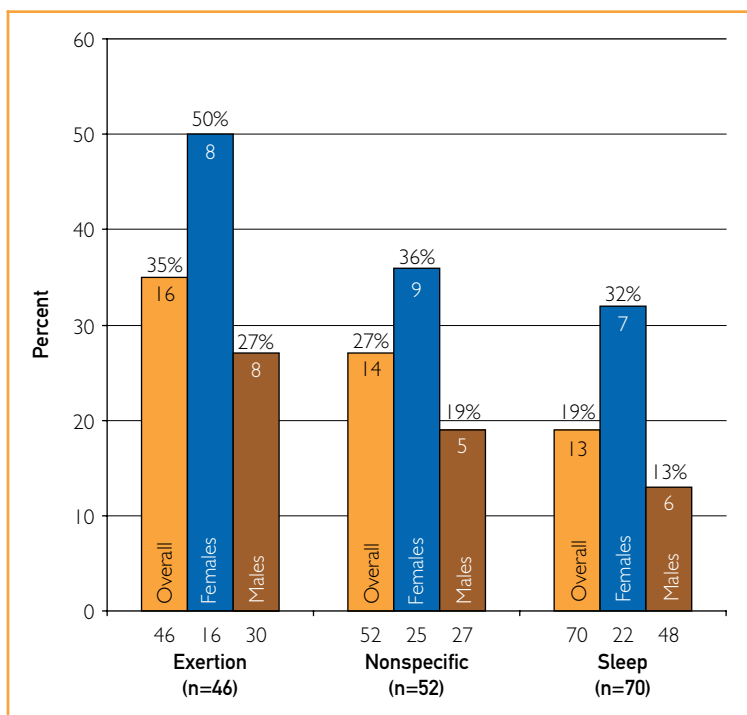
and unanticipated (83.3% [5/6] *KCNQ1* and 16.7% [1/6] *RYR2*, Figure 8).

#### Effect of Personal History and Positive Family History on Yield.

As anticipated, the mutation detection yield was significantly higher in patients with SUD who had either a positive personal or family history of cardiac events when compared with those in whom no personal or family history was elicited or documented (37.1% [26/70] vs 19.1% [17/89];  $P=.01$ ; Figure 9). Moreover, nearly half (45.0% [9/20]) of patients with SUD who had a positive family history of premature SCD and who were younger than 50 years had a positive cardiac channel molecular autopsy with identification of a rare, potentially channelopathic mutation.

#### DISCUSSION

Postmortem genetic testing (the cardiac channel molecular autopsy) has not yet been transformed fully from a research-based effort into a routine, standard part of the conventional autopsy when the coroner, medical examiner, or forensic pathologist is faced with a case of SUD in the young. However, recently, Basso and colleagues,<sup>18</sup> on behalf of the



**FIGURE 6.** The different yield of mutation detection among the 3 main categories of sudden unexplained death–associated triggers or events (exertion, nonspecific, and sleep) for the overall cohort, females, and males at all ages. The numbers below the bars represent the number of cases in each category. The numbers within the bars represent the actual number of cases with a mutation, with the percent yield of mutation detection highlighted above the bars.

Association for European Cardiovascular Pathology, have strongly recommended postmortem genetic analysis in both structural and nonstructural genetically determined heart disease as part of the requirements for the adequate postmortem assessment of SCD. In 2008, members of the Trans-Tasman Response AGAinst sudden Death in the Young (TRAGADY), endorsed by the Royal College of Pathologists of Australasia, put forward guidelines to ensure standardization of autopsy practice in young sudden unexpected deaths, ancillary testing, and retention of appropriate material for all cases of SCD to be used for genetic testing. In 2011, the Heart Rhythm Society and the European Heart Rhythm Association provided an expert consensus statement on the state of genetic testing, providing consensus-based guidelines and recommendations for postmortem genetic testing in cases of SUD.<sup>20</sup> Accordingly, the key recommendations of the Heart Rhythm Society/European Heart Rhythm Association guidelines state that “in the setting of autopsy negative SUD, comprehensive or targeted (*KCNQ1*, *KCNH2*, *SCN5A*, *RYR2*) ion channel genetic testing

may be considered in an attempt to establish probable cause and manner of death and to facilitate the identification of potentially at-risk relatives and is recommended if circumstantial evidence points toward a clinical diagnosis of LQTS or CPVT specifically.”

Unfortunately, for several reasons, it has been extremely difficult for the medical examiner, coroner, or forensic pathologist to provide this level of care. Perhaps chief among the reasons is that insurance companies or other third-party payers largely do not accept any responsibility for providing coverage for the molecular autopsy of a deceased person, regardless of the implications to surviving relatives. This position, not surprisingly, limits postmortem genetic testing to only families who can afford the out-of-pocket expense for commercially available genetic testing because few medical examiner offices have the internal budget to cover such a molecular autopsy. Alternatively, the medical examiner, coroner, or forensic pathologist and the grieving family are left with the option of enrolling the deceased’s sample into research-based genetic testing, where although the price is right (ie, free), the process can be painfully slow.<sup>49</sup> Given the expensive and time-consuming nature of postmortem genetic testing, it is currently necessary for the medical examiner, coroner, or forensic pathologist to be case selective in pursuing a molecular autopsy.<sup>50</sup>

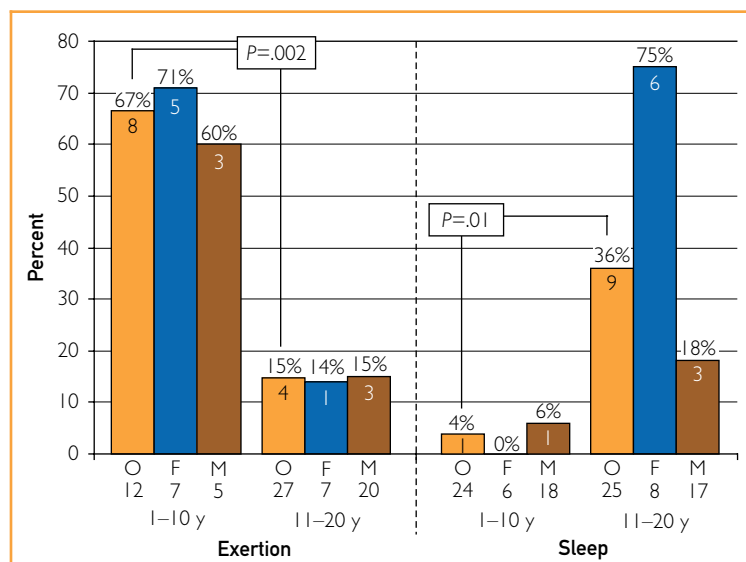
In 2007, we completed our original molecular autopsy series of 49 medical examiner–referred cases of autopsy-negative SUD.<sup>23</sup> Comprehensive mutational analysis of all 60 translated exons in the LQTS-associated genes, *KCNQ1*, *KCNH2*, *SCN5A*, *KCNE1*, and *KCNE2*, along with targeted analysis of the CPVT1-associated *RYR2*-encoded cardiac ryanodine receptor revealed that more than one-third of SUD cases hosted a seemingly pathogenic cardiac channel mutation.<sup>22</sup> We have extended this cohort to now include more than 170 cases of SUD to provide a more extensive analysis to better define the expected yield of mutation detection and offer possible genotype-phenotype correlations that may assist in guiding phenotype-directed mutation detection efforts in future cases of SUD.

In this expanded molecular analysis of 173 SUD cases, we provide molecular evidence that suggests that just more than one-fourth (26.0%) of these SUD cases host putative pathogenic mutations in critical ion channel genes associated with the potentially lethal arrhythmia syndromes LQTS and CPVT. Besides these 26.0% of cases that host putative pathogenic mutations, another 16.2% of this cohort hosted nonsynonymous, amino acid–altering, functional polymorphisms previously associated with LQTS, Brugada syndrome, drug-induced torsades

de pointes, or ventricular arrhythmias or SCD polymorphisms. Thus, the overall contribution of channelopathies or sudden death–predisposing or proarrhythmic, functional channel polymorphisms is likely underestimated in this study. For example, we excluded 10 SUD patients (5.8%) who were positive for D85N-KCNE1, R1047L-KCNH2, or S1103Y-SCN5A. These proarrhythmic polymorphisms have been associated convincingly with increased risk for sudden death, particularly in the setting of drugs with an unwanted QT-prolonging effect. Despite our efforts, we were unable to elicit a reliable history of prevent medication exposures to determine whether a case for drug exposure plus functional polymorphism could be made. Nevertheless and perhaps more importantly than this overall point estimate for the yield of such postmortem genetic testing in the setting of an SUD, this cohort has provided us with several interesting genotype-phenotype observations that may provide insight into the expected yields of postmortem genetic testing for SUD and assist in selecting cases with the greatest potential for mutation discovery and directing genetic testing efforts.

Interestingly, although males represented two-thirds of the cohort, females were more likely to host a channel mutation than males, especially if the SUD was during adolescence (48.0% females vs 18.0% males). Moreover, there was a cardiac channelopathy–sex effect, with two-thirds of the mutation-positive females having an LQTS-associated mutation compared with only one-third having a CPVT-associated *RYR2* mutation, whereas mutation-positive males most often had a CPVT (approximately 60%) mutation rather than an LQTS mutation (approximately 40%). On average, males who had mutations in LQTS- or CPVT-associated genes were younger than females who were genotype positive. This sex predilection is consistent with previous work by Priori et al,<sup>51</sup> in which among their cohort of 30 CPVT index cases, most of the *RYR2* mutations were identified in young males (11/13 [85%]) compared with females (3/17 [18%]), and male sex in *RYR2*-mediated CPVT was associated with a relative risk of 4.2 (95% confidence interval, 1.2–15) of developing syncope compared with females. Accordingly, one might a priori expect a higher yield of LQTS-associated mutation detection in a female adolescent or young adult compared with a higher expected yield of CPVT-associated mutations among prepubertal boys.

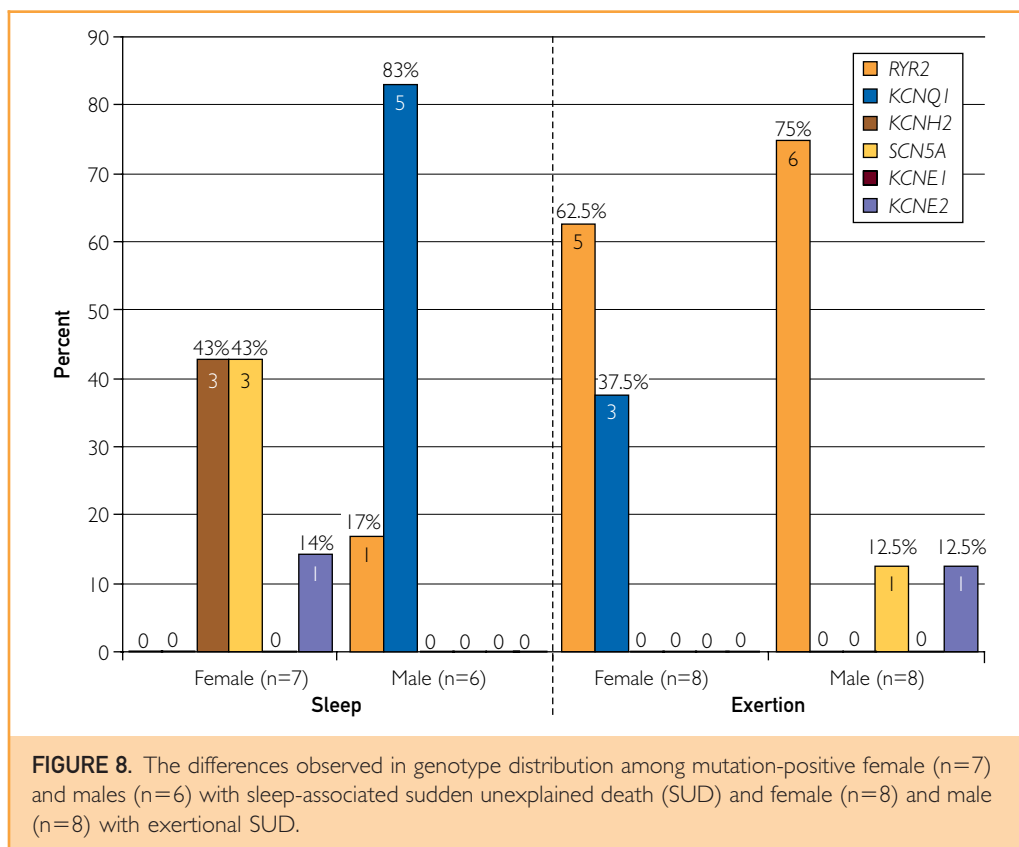
The circumstances surrounding the death also had an effect on the mutation detection yield, in which decedents had the highest mutation detection rate when the death was associated with exertion compared with a nonspecific triggered death or death during sleep. This trend held true for females and males, with females consistently having higher yields than males within the 3 event or trigger cate-



**FIGURE 7.** The difference in mutation detection between children aged 1 to 10 years and adolescents aged 11 to 20 years with either exertion- or sleep-associated sudden death. The numbers below O, F, and M represent the number of cases in each category. The numbers within the bars represent the absolute number of cases with a mutation, with the percent yield of mutation detection highlighted above the bars. F = females; M = males; O = overall.

gories. Although half of all females and 26.7% of males with exercise-associated sudden death were identified as mutation positive, only 31.8% and 12.5%, respectively, had mutations when the death occurred during sleep. However, an interesting age effect was revealed when examining the effect on yield when comparing exercise-induced death with death during sleep. For example, 71.4% of the 1- to 10-year-old girls with exercise-induced death were found to be mutation positive compared with 0% of the 1- to 10-year-old girls found dead in bed. Conversely, although only 14.3% (1 of 7) of 11- to 15-year-old girls with exercise-induced death were mutation positive, three-fourths of the adolescent girls with death during sleep were identified as mutation positive. Although failing to reach statistical significance, this trend was also observed among males.

Previously, specific genotype-phenotype associations in LQTS have been established, suggesting relatively gene-specific triggers.<sup>52</sup> Exertion-induced cardiac events are associated strongly with mutations in *KCNQ1* (LQT1), whereas auditory triggers and events occurring during the postpartum period most often occur in patients with LQT2 (*KCNH2*). Although exertion- or emotional stress–induced events are most common in LQT1 (*KCNQ1*), events occurring during periods of sleep or rest are most common in LQT3 (*SCN5A*). CPVT1 (*RYR2*) classically



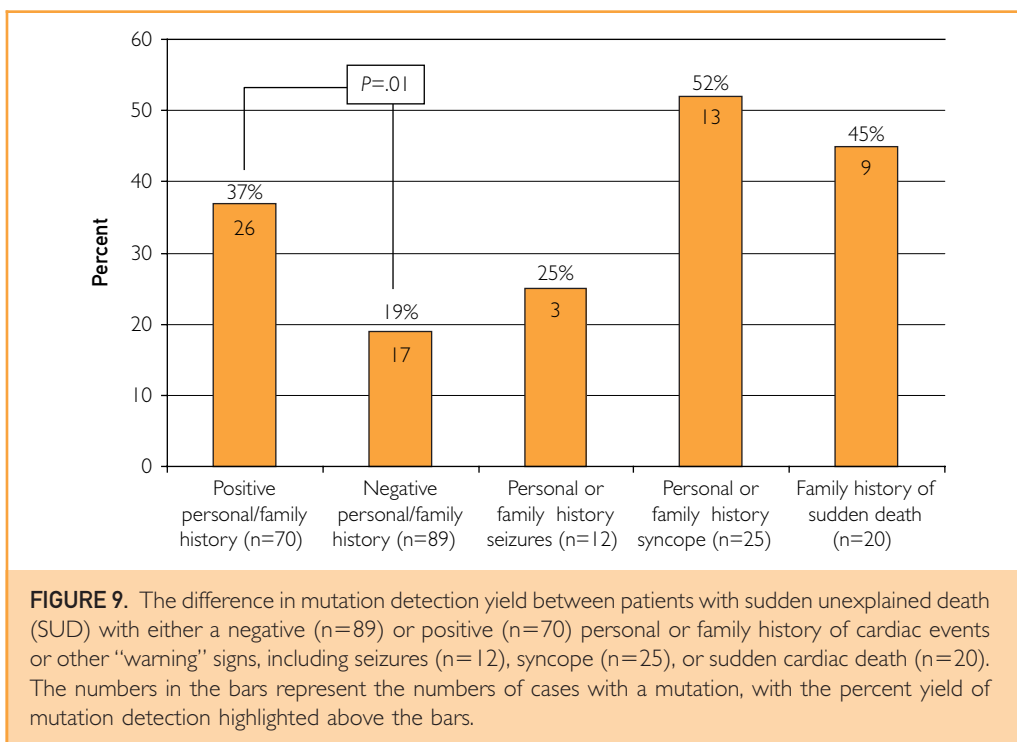
manifests with exercise-induced syncope or sudden death<sup>51</sup> and rarely manifests during periods of rest.

These particularly well-validated genotype-phenotype associations have led some investigators to selectively direct genetic testing efforts based on trigger-associated events, particularly in LQTS cases.<sup>53</sup> For example, if an LQTS individual presents with exertion-induced syncope or cardiac arrest, then a genetic test for *KCNQ1* (LQT1) mutations should be performed. If the syncope or cardiac arrest occurred during periods of rest or sleep, then a test for *SCN5A* (LQT3) should be performed. Although the observed mutation-positive gene distribution in this SUD cohort matched the anticipated results for those males and females with exercise-induced death (ie, overwhelmingly *RYR2* or *KCNQ1* mutations) and females with death during sleep (ie, mostly *SCN5A* and no *RYR2* or *KCNQ1* mutations), strikingly and unexpectedly, all of the mutation-positive males with death during sleep had mutations in either *KCNQ1* (83.3%) or *RYR2* (16.7%).

Both LQTS and CPVT are hereditary disorders that manifest within families with potential warning signs, such as syncope, seizures, survived cardiac arrest, near drowning, or prolonged QT interval, that may lead to a clinical diagnosis of the disorder and circumvent the sudden death of a prophylacti-

cally treated family member. However, tragically these particular warning signs often go unheeded or unrecognized as evident by the remarkable presence of a positive family history of cardiac events in 40.5% of our SUD cohort, including nearly 11.6% of the patients with a positive family history of a prior sudden death. Yet, no decedent or relative had received a clinical diagnosis of a suspected cardiac channelopathy or heritable arrhythmia syndrome (ie, LQTS or CPVT) before the SUD of the current patient in our study. As expected, SUD cases with such a positive personal or family history had a significantly higher mutation detection yield (37.1%) than those with no personal or family history (19.1%) of cardiac events. In fact, among the subset of patients with a positive family history of otherwise unexplained sudden death before the age of 50 years, nearly half were positive for a putative channelopathic mutation.

Interestingly, there was a trend toward a higher mutation detection yield in SUD patients with a personal or family history of syncope compared with a history of seizures. Although not as common as syncope, seizure-like symptoms are experienced by some patients with LQTS or CPVT. Our data suggest that SUD in an individual with a personal or family history of seizure activity may in fact be a result of a



neurologic disorder, akin to SUD in epilepsy (SUDEP), rather than as a result of a primarily cardiogenic disorder, such as LQTS or CPVT.

Although mutations in LQTS and CPVT genes have been illustrated as the pathogenic basis from some SUDEP cases, most SUDEP appears to not be related to mutations in major genes associated with these 2 disorders. In 2010, Tu et al<sup>74</sup> analyzed a cohort of 68 SUDEP cases for mutations in 3 major LQTS genes (*KCNQ1*, *KCNH2*, and *SCN5A*) and only identified 2 patients (3%) with putative pathogenic mutations (R176W-KCNH2 and P1090L-SCN5A), which were absent in their control population. However, both of these genetic variants have been observed in other control populations and represent functionally significant polymorphisms that confer an increased risk for arrhythmia.<sup>32,33</sup> Together, these data suggest that sudden death cases with a personal or family history of seizures may warrant mutational analysis of ion channel encoding genes, such as *SCN1A* and *KCNA1*, which are abundantly expressed in the brain and have been implicated as candidates for SUDEP,<sup>55,56</sup> rather than starting with an initial interrogation of LQTS- and CPVT-associated genes. Whether our genotype-negative SUD cases with a personal or family history of seizure activity have mutations in these SUDEP candidate genes is currently unknown. In contrast, a personal or family history of unexplained syncope, especially exercise-induced syncope, may reflect a

strong indicator for LQTS and CPVT genetic testing for the SUD case.

Although a positive personal or family history is associated with a 2-fold greater likelihood of a positive postmortem genetic test result than when there is no history, a negative personal or family history alone should not preclude genetic testing with the intent to identify mutations that may be familial and therefore provide an increased risk for sudden death of an unsuspecting surviving family member of the decedent. In fact, among the 11 mutation-positive SUD families that chose to participate in genetic pedigree analysis in our study, all 11 SUD cases were identified as having a familial mutation, rather than a sporadic one, despite having a negative personal or family history of cardiac events in 72.7% (8/11) of these families, enabling personal risk stratification and prophylactic therapies for the mutation-positive family members who are still living.

The major limitation of this study pertains to the issue of mutation calling.<sup>34</sup> Although most mutations identified in this study are either radical (ie, frameshift or splicing errors) or localize to critical ion channel domains (ie, transmembrane region or channel pore regions), with a predicted probability of pathogenicity exceeding 90%, most mutations identified in this study, in particular those that are novel, have not been characterized functionally. Heterologous expression studies for every ion channel variant identified to assess in vitro functionality are not feasible practically.

Instead, strict absence from our internal set of ethnic-matched reference alleles was required. Nevertheless, it is possible that some of these variants listed as putative pathogenic mutations may be innocuous functionally. Therein lies the difficulty in determining the “actionableness” of these rare, non-functionally characterized genetic variants for surviving family members when the probability of pathogenicity for that given variant is not 100%. However, this issue of mutation calling is present in pre-mortem genetic testing as well and underscores the reality that, whether the patient is dead or alive, a genetic test result must be interpreted with great caution.

### CONCLUSION

More than one-fourth of autopsy-negative SUD cases may stem from either LQTS- or CPVT-associated channelopathic mutations. A cardiac channel molecular autopsy should be considered as a standard part of the evaluation of autopsy-negative SUD, especially among children with exercise-induced SUD, adolescent girls, and those with a positive personal or family history of cardiac events. A 2-tiered approach starting with the LQTS-susceptibility genes for female patients with SUD and the CPVT susceptibility genes for males with SUD may be prudent.

**Abbreviations and Acronyms:** CPVT = catecholaminergic polymorphic ventricular tachycardia; LQTS = long QT syndrome; SCD = sudden cardiac death; SUD = sudden unexplained death; SUDEP = sudden unexplained death in epilepsy

**Grant Support:** This work was supported by the Mayo Clinic Windland Smith Rice Comprehensive Sudden Cardiac Death Program, the Sheikh Zayed Saif Mohammed Al Nahyan Fund in Pediatric Cardiology Research, the Dr. Scholl Fund, the Hannah M. Wemke Memorial Fund, and the National Institutes of Health, grant number HD42569.

**Potential Competing Interests:** M.J.A. is a consultant for Biotronik, Boston Scientific, Medtronic, St. Jude Medical Inc, and Transgenomic. Intellectual property derived from M.J.A.'s research program resulted in license agreements in 2004 between Mayo Clinic Health Solutions (formerly Mayo Medical Ventures) and PGx-Health (formerly Genaisance Pharmaceuticals, recently acquired by Transgenomic). D.J.T., A.M.-D., M.L.W., and C.M.H. have no conflicts to disclose. M.J.A. had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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