<u>薛田直昌</u>	【致死性不整脈診療の最前線】 致死性不整脈診療 遺伝性心臓 伝導障害	最新医学	68	1588-1596	2013
蒔田直昌	【イオンチャネル病のすべて】 進行性心臓伝導障害	医学のあゆみ	245	802-809	2013
<u>蒔田直昌</u>	難治性不整脈の遺伝子解析	循環器専門医	21	3-8	2013
Watanabe H, Makita N, Tanabe N, Watanabe T, Aizawa Y	Electrocardiographic abnormalities and risk of complete atrioventricular block.	Int J Cardiol	155	462-464	2012
T, Nagao S, Yagihara N, Takehara N, Kawamura Y, Sato A, Okamura K, Hosaka Y, Sato M, Fukae S, Chinushi M, Oda H, Okabe M, Kimura A, Maemura K, Watanabe I, Kamakura S, Horie M, Aizawa Y, Shimizu W, Makita N	Response to Letter Regarding Article, "Electrocardiographic Characteristics and SCN5A Mutations in Idiopathic Ventricular Fibrillation Associated With Early Repolarization"		5	e60-e61	2012
Delmar M, <u>Makita N</u>	Cardiac Connexins, Mutations and Arrhythmias	Curr Opin Cardiol	27	236-241	2012
1	Membrane-Associated Protein Gene Mutations Impair Intracellular Trafficking of hNav1.5.	Circ Arrhythm Electrophysiol	5	1098-1107	2012
Shimada T, Ohkubo K, Abe K, Watanabe I, <u>Makita N</u>	A novel 5' splice site mutation of SCN5A associated with Brugada syndrome resulting in multiple cryptic transcripts	Int J Cardiol	158	441-443	2012
蒔田 直昌	特発性心室細動とJ波症候群の遺 伝子診断	CIRCULATION Up-to-Date	7	20-25	2012
蒔田 直昌	早期再分極とJ波症候群:オーバービュー	心臓	44	1226-1231	2012
	Utility of ECG-gated MDCT to differentiate patients with ARVC/D from patients with ventricular tachyarrhythmias.	J Cardiovasc Comput Tomogr	7	223-233	2013
	Mutations in the cardiac troponin T gene show various prognoses in Japanese patients with hypertrophic cardiomyopathy.	Heart Vessels	28	785-794	2013

Naganuma M, Suzuki A, <u>Hagiwara N</u>	Contributing factors to the apparent clearance of bepridil in patients with paroxysmal or persistent atrial fibrillation: analysis using population pharmacokinetics.  Estradiol promotes neural stem	Ther Drug Monit  Angiogenesis.	35	367-373 45-58	2013
K, Thorne T, Ito A, Klyachko E, Hamada H, Kessler JA, Tabata Y, Kawana M, Asahi M, <u>Hagiwara N</u> , Losordo DW	cell differentiation into endothelial lineage and angiogenesis in injured peripheral nerve.				2012
Matsuura K, Wada M, Shimizu T, Haraguchi Y, Sato F, Sugiyama K, Konishi K, Shiba Y, Ichikawa H, Tachibana A, Ikeda U, Yamato M, Hagiwara N, Okano T	Creation of human cardiac cell sheets using pluripotent stem cells.	Biochem Biophys Res Commun.	425	321-327	2012
	Generation and characterization of functional cardiomyocytes derived from human T cell-derived induced pluripotent stem cells.	PLoS One	9	e85645	2014
Ohno Y, Yuasa S, Egashira T, Seki T, Hashimoto H, Tohyama S, Saito Y, Kunitomi A, Shimoji K, Onizuka T, Kageyama T, Yae K, Tanaka T, Kaneda R, Hattori F, MurataM, Kimura K, <u>Fukuda K</u>	Cardiac Differentiation Efficiency.	Stem Cells Int	E-pub		2013
Hasegawa M, Lichtler A, Reichenberger EJ	Induced pluripotent stem cell reprogramming by integration-free sendai virus vectors from peripheral blood of patients with craniometaphyseal dysplasia.	Cell Reprogram	15	503-513	2013
	The generation of induced pluripotent stem cells from a patient with KCNH2 G603D, without LQT2 disease associated symptom	J Med Dent Sci	60	17-22	2013

Wada R, Muraoka N, Inagawa K, Yamakawa H, Miyamoto K, Sadahiro T, Umei T, Kaneda R, Suzuki T, Kamiya K, Tohyama S, Yuasa S, Kokaji K, Aeba R, Yozu R, Yamagishi H, Kitamura T, Fukuda K, Ieda M	Induction of human cardiomyocyte-like cells from fibroblasts by defined factors.	Proc Natl Acad Sci USA	110	12667-12672	2013
Egashira T, Yuasa S, Fukuda K	Novel insights into disease modeling using induced pluripotent stem cells.	Biol Pharm Bull	36	182-188	2013
福田惠一	再生医学・再生医療の最前線 iPS細胞の循環器領域への臨床 応用	日本内科学会雑誌	102	2232-2240	2013
Seki T, Yuasa S, Fukuda K	Generation of induced pluripotent stem cells from a small amount of human peripheral blood using a combination of activated T cells and Sendai virus.	Nat Protoc.	7	718-728	2012
Mitani Y, Ohta K Ichida F, Nii M, Arakaki Y, Ushinohama H, Takahashi T, Ohashi H, Yodoya N, Fujii E, Ishikura K, Tateno S, Sato S, Suzuki T, Higaki T, Iwamoto M, Yoshinaga M, Nagashima M, Sumitomo N	Circumstances and Outcomes of Out-Of-Hospital Cardiac Arrest in Elementary and Middle School Students in the Era of Public-Access Defibrillation: Implications for Emergency Preparedness in Schools.	Circ J	78	701-707	2014
Yoshikane Y, Yoshinaga M, Hamamoto K, Hirose S	A case of long QT syndrome with triple gene abnormalities: Digenic mutations in KCNH2 and SCN5A and gene variant in KCNE1.	Heart Rhythm	10	600-603	2013
Ninomiya Y, Yoshinaga M, Kucho Y, Tanaka Y	Risk factors for symptoms in long QT syndrome in a single pediatric center.	Peadiatr Int	55	277-282	2013
Yoshinaga M	Prevalence of sudden death and out-of-hospital cardiac arrest in infants, children, and adolescents; what does it imply?	Circ J	77	2475-2476	2013
吉永正夫	乳児突然死症候群とQT延長症 候群	日本小児科学会 雑誌	117	44-48	2013
吉永正夫, 長嶋正實	自動計測とマニュアル計測での QT時間の差に関する検討	心電図	32	7-35	2013

吉永正夫, 泉田直己, 岩本眞理, 牛ノ濱大 也, <u>住友直方</u> , 田打宣 生, 高橋良明, 富田 英,長嶋正實, <u>堀米仁</u> 志, 山内邦明, 日本 小児循環器学会学校 心臟検診委員会	器質的心疾患を認めない不整脈 の学校生活管理指導ガイドライ ン(2013年改訂版)		29	277-290	2013
吉永正夫	心臓突然死のリスク評価のパラメータ、HRT,HRV,QT時間の日内変動-自律神経の関与の面から-	循環器診療	18	43-47	2013
吉永正夫, 長嶋正實	Timothy症候群	医学のあゆみ	245	821-824	2013
堀米仁志, 石川康宏, 加藤愛章, 中村昭宏, 岩本眞理, <u>住友直方</u> , 吉永正夫	独立成分文政期を用いた先天性 QT延長症候群のT波の解析 - 主 成分分析との診断精度の比較 -	i e	8	14-25	2013
JF, Yu S, Horigome H,	In Utero Diagnosis of Long QT Syndrome by Magnetocardiography. Circulation.	Circulation	128	2183-2191	2013
Cuneo BF, Etheridge SP, Horigome H, Sallee D, Moon-Grady A, Weng HY, Ackerman MJ, Benson DW	Arrhythmia Phenotype During Fetal Life Suggests Long-QT Syndrome Genotype: Risk Stratification of Perinatal Long-QT Syndrome.	Circ Arrhythm Electrophysiol	6	946-951	2013
Mitchell JL, Cuneo BF, Etheridge SP, Horigome H, Weng HY, Benson DW	Fetal heart rate predictors of long QT syndrome	Circulation	126	2688-2695	2012
Mitani Y, Ohta K, Yodoya N, Otsuki S, Ohashi H, Sawada H, Nagashima M, Sumitomo N, Komada Y	Public access defibrillation improved the outcome after out-of-hospital cardiac arrest in school-age children: a nationwide, population-based Utstein registry study in Japan.	Europace	15	1256-1266	2013

	-	~			
Lubitz SA, Lunetta KL,	Novel genetic markers associate	J Am Coll Cardiol		In press	2014
Lin H, Arking DE,	with atrial fibrillation risk in				
Trompet S, Li G,	Europeans and Japanese.				
Krijthe BP, Chasman					
DI, Barnard J, Kleber					
ME, Dörr M, Ozaki K,					
The state of the s					
Smith AV, Müller M,					
Walter S, Agarwal SK,					
Bis JC, Brody JA,					
Chen LY, Everett BM,					
Ford I, Franco OH,					
Harris TB, Hofman A,					
Kääb S, Mahida S,					
Kathiresan S, Kubo M,					
Launer LJ, MacFarlane					
PW, Magnani JW,					
McKnight B,					
McManus DD, Peters					
A, Psaty BM, Rose					
LM, Rotter JI,					
Silbernagel G, Smith					
JD, Sotoodehnia N,					
1					
Stott DJ, Taylor K,					
Tomaschitz A Tsunoda					
T, Uitterlinden AG,					
VanWagoner DR,					
Völker U, Völzke H,					
Murabito JM, Sinner					
MF, Gudnason V,					
Felix SB, März W,					
Chung M, Albert CM,					
Stricker BH, <u>Tanaka T</u> ,					
Heckbert SR, Jukema					
JW, Alonso A,					
Benjamin EJ, Ellinor					
PT					
Behr ER, Ritchie MD,	Genome wide analysis of	PLoS One	13	08180R2	2013
Tanaka T, Kääb S,	drug-induced Torsades de Pointes:				
	lack of common variants with large				
	effect sizes.				
P, Floratos A, Sinner	errect sizes.				
MF, Kannankeril PJ,					
AA. M. Wilde, Bezzina					
CR, Schulze-Bahr E,					
Zumhagen S,					
Guicheney P, Bishopric					
NH, Marshall V, Shakir				Processing	
S, Dalageorgou C,					
1 -					
Bevan S, Jamshidi Y,					
Bastiaenen R,					
Myerberg RJ, Schott					
J-J, Camm AJ,					
Steinbeck G, Norris K,					
	i e		I	1	1
Altman RB, Tatonetti					
Altman RB, Tatonetti N. Jeffery S, Kubo M.					. :
N, Jeffery S, Kubo M,					. :
N, Jeffery S, Kubo M, Nakamura Y, Shen Y,					. :
N, Jeffery S, Kubo M,					. :
N, Jeffery S, Kubo M, Nakamura Y, Shen Y,					. :
N, Jeffery S, Kubo M, Nakamura Y, Shen Y,					. :

Okada Y, Kubo M, Ohmiya H, Takahashi A, Kumasaka N, Hosono N, Maeda S, Wen W, Dorajoo R, Go MJ, Zheng W, Kato N, Wu J-Y, Lu Q, GIANT consortium, Tsunoda T, Yamamoto K, Nakamura Y, Kamatani N, Tanaka T		Nat Genet	44	302-306	2012
Morita H, Miura D, Nishii N, Nagase S,	Depolarization and Repolarization Abnormalities are Synergistically Associated with Fatal Arrhythmic Events in Patients with Brugada Syndrome.	J Am Coll Cardiol	63	In press	2014
Nakagawa K, Nagase S, <u>Morita H</u> , Ito H	Left ventricular epicardial electrogram recordings in idiopathic ventricular fibrillation with inferior and lateral early repolarization.	Heart Rhythm	11	314-317	2014
Wada T, <u>Morita H</u>	Clinical outcome and risk stratification in Brugada syndrome.	J Arrhythmia	29	100-109	2013
Morita H	Ion channel complex disease in long QT syndrome.	Heart Rhythm	10	738-739	2013
上岡 亮,森田 宏	Brugada症候群と特発性心室細動	レジデント	7	91-100	2014
	Brugda症候群	最新医学	68	1579-1587	2013
Take Y, <u>Morita H</u>	Identification of high-risk syncope related to ventricular fibrillation in patietns with Brugada syndrome	Heart Rhythm	9	752-759	2012
Morita H	The compound mutation, a model for acquire long QT syndrome	J Cardiol Cases	6	e187-e188	2012
Take Y, <u>Morita H</u>	Fragemented QRS: what is the meaning?	Indian Pacing and Electrophysiology Journal	12	213-225	2012
<u>森田 宏</u>	J波症候群及びBrugada症候群の 活動電位、心電図、電気生理学 的検査の特徴	心電図	32	S4-56-71	2012
<u>森田 宏</u>	早期再分極症候群とJ波症候群- 細胞学的成因について	心臓	44	1232-1236	2012
Kato K, <u>Makiyama T,</u> Wu J, Ding WG, Kimura H, Naiki N, Ohno S, Itoh H, Nakanishi T, Matsuura H, <u>Horie M</u>	Cardiac channelopathies associated with infantile fatal ventricular arrhythmias: From the cradle to the bench.	J Cardiovasc Electrophysiol	25	66-73	2014

Kamakura T, Makiyama T, Sasaki K, Yoshida Y, Wuriyanghai Y, Chen J, Hattori T, Ohno S, Kita T, Horie M, Yamanaka S, Kimura T	Ultrastructural maturation of human-induced pluripotent stem cell-derived cardiomyocytes in a long-term culture.	Circ J	77	1307-1314	2013
Nakajima S, <u>Makiyama</u> <u>T</u> , Hanazawa K, Kaitani K, Amano M, Hayama Y, Onishi N, Tamaki Y, Miyake M, Tamura T, Kondo H, Motooka M, Izumi C, Nakagawa Y, <u>Horie M</u>	A novel SCN5A mutation demonstrating a variety of clinical phenotypes in familial sick sinus syndrome.	Intern Med	52	1805-1808	2013
Fukuyama M, Ohno S, Wang Q, Kimura H, Makiyama T, Itoh H, Ito M, Horie M	L-type calcium channelmutations in Japanese patients with inherited arrhythmias.	Circ J	77	1799-1806	2013
Ohno S, Nagaoka I, Fukuyama M, Kimura H, Itoh H, <u>Makiyama</u> T, Shimizu A, <u>Horie M</u>	Age-dependent clinical and genetic characteristics in Japanese patients with arrhythmogenic right ventricular cardiomyopathy/dysplasia.	Circ J	77	1534-1542	2013
牧山 武	疾患特異的iPS細胞を用いた遺 伝性心疾患研究	心電図	33, Suppl	5012-5022	2013
Katsuumi G, Shimizu W, Watanabe H, Noda T, Nogami A, Ohkubo K, Makiyama T, Takehara N, Kawamura Y, Hosaka Y, Sato M, Fukae S, Chinushi M, Oda H, Okabe M, Kimura A, Maemura K, Watanabe I, Kamakura S, Horie M, Aizawa Y, Makita N, Minamino T	Efficacy of bepridil to prevent ventricular fibrillation in severe form of early repolarization syndrome.	Int J Cardiol	E-pub		2014
Watanabe H, Van der Werf C, Roses-Noguer F, Adler A, Sumitomo N, Veltmann C, Rosso R, Bhuiyan ZA, Bikker H, Kannankeril PJ, Horie M, Minamino T, Viskin S, Knollmann BC, Till J, Wilde AA	Effects of flecainide on exercise-induced ventricular arrhythmias and recurrences in genotype-negative patients with catecholaminergic polymorphic ventricular tachycardia.	Heart Rhythm	10	542-547	2013

Watanabe H. Minamino	Role of mutations in l-type calcium	Circ J	77	1689-1690	2013
T	channel genes in brugada syndrome, early repolarization syndrome, and idiopathic ventricular fibrillation associated with right bundle branch block.				
Watanabe H, Minamino T	Similarities and differences of clinical characteristics between brugda syndrome and early repolarization syndrome.	J Arrhythmia	29	134-137	2013
	不整脈症候群ならびに心臓電気 生理学における遺伝学研究の進 歩.	血管	36	63-68	2013
渡部 裕, 南野 徹	早期再分極症候群の特徴	最新医学	68	422-428	2013
Kawashiri MA, <u>Hayashi K</u> , Konno T, Fujino N, Ino H, Yamagishi M	Current perspectives in genetic cardiovascular disorders: from basic to clinical aspects.	Heart Vessels	29	129-141	2014
Fujino N, Uchiyama K, Konno T, Tsuda T,	transmembrane nonpore region of the KCNH2 gene causes severe clinical manifestations of long QT syndrome.	Heart Rhythm	1	61-67	2013
林 研至,津田豊暢,川尻剛照,山岸正和	家族性心房細動	最新医学	68	1626-1634	2013
Kamakura S	Epidemiology of Brugada syndrome in Japan and rest of the world.	J Arrhythmia	29	52-55	2013
Kamakura S	Two decades of progress in the understanding of Brugada syndrome.	J Arrhythmia	29	51	2013
鎌倉史郎	Brugada症候群	医学のあゆみ	245	782-789	2013
鎌倉史郎	Brugada波形	日本医事新報	4670	33-39	2013
I, Iwai N, <u>Miyamoto Y</u> , Kokubo Y, Okamura T,	CDH13 Gene Coding T-Cadherin Influences Variations in Plasma Adiponectin Levels in the Japanese Population.	Hum Mutat	33	402-410	2012
Aiba T, Barth AS, Hesketh GG, Hashamb	Cardiac Resynchronization Therapy Improves Altered Na Channel Gating in Canine Model ofDyssynchronous Heart Failure.	Circ Arrhythm Electrophysiol.	6	546-554	2013

Das S, <u>Aiba T</u> ,	Pathological role of serum- and	Circulation	126	2208-2219	2012
	glucocorticoid-regulated kinase 1	Circulation	120	2200-2219	2012
	in adverse ventricular remodeling.				
Ottaviano FG, Knight	adverse ventricular remodelling.				
AC, Graham EL,					
Boström P,Morissette					
1					
MR, del Monte F,					
Begley MJ, Cantley					
LC, Ellinor PT,					
Tomaselli GF,					
RosenzweigA					
1	Mastication and risk fordiabetes in	PLoS One	8	e64113	2013
M, Asai K,	a Japanese population: a				
Nakano-Araki I,	cross-sectional study.				
Yamaguchi A,					
Takahashi K, Sekine A,					
Matsuda F, Kosugi S,					
Nakayama T, Inagaki					
N, Bessho K:Nagahama					
Study Collaboration					
Group					
•					
Tabara Y, Takahashi Y.	Association of Longer QT Interval	Am J Hypertens	26	973-980	2013
Kohara K, Setoh K,	With Arterial Waveform and		[		
Kawaguchi T, Terao C,	i .				
-	Amplification: The Nagahama				
	Study.				
Miki T, Nakayama T,					
Matsuda F: Nagahama					
Study Group					
The state of the s					
	Replication study of 15recently	J Atheroscler	20	336-350	2013
	published Loci for body fat	Thromb			
1	distribution in the Japanese				
1	population.				
Hyogo H, Ochi H,					
Nakamura T,					
Kamohara S, Miyatake					
N, Kotani K, Itoh N,					
MineoI, Wada J,					
Yoneda M, Nakajima					
A, Funahashi T,			1		
Miyazaki S, Tokunaga					
K, MasuzakiH, Ueno T,					
Chayama K,					
Hamaguchi K, Yamada					
K, Hanafusa T, Oikawa	·				
S, Sakata T, Tanaka K,					
Matsuzawa Y, Nakao					
K, Sekine A					
1			1		

Kitamoto A, Kitamoto	NUDT3 rs206936 is associated	Endocr J	60	991-1000	2013
T, Mizusawa S,	with body massindex in obese				
Teranishi H,So R,	Japanesewomen.				
Matsuo T, Nakata Y,					
HyogoH, Ochi H,					
Nakamura T,					
Kamohara S, Miyatake					
N, Kotani K, Komatsu,					
Itoh N, Mineo I, Wada					
J, Yoneda M,					
NakajimaA, Funahashi					
T, Miyazaki S,				500 A C C C C C C C C C C C C C C C C C C	
Tokunaga K,					
MasuzakiH, Ueno T,					
Chayama K,					
Hamaguchi K, Yamada K, Hanafusa T, Oikawa					
S, Sakata T, Tanaka K,					
Matsuzawa Y, Nakao					
K, Sekine A, Hotta K					
ix, <u>bekine 11,</u> Hotta ix					
Yoshimura K,	B-type natriuretic peptide as an	Neurourology and	31	1266-1271	2012
Nakayama T, Sekine A,	independent correlate of nocturnal	urodynamics			
Matsuda F, Kosugi S,	voiding in Japanese women.				
Yamada R, Shimizu Y,					
Kanematsu A,					
Yoshimura K, Ogawa					
O; Nagahama Cohort			:		
Research Group					
TO 1 TO NOT 1		T.C. 1: 1	12	205 205	2014
Tokuyama T, Nakano	Deterioration of the circadian	J Cardiol	13	385-387	2014
	variation of heart rate variability in				
–Makita Y, Fujiwra M, Watanabe Y, Sairaku	Brugada syndrome may contribute to the pathogenesis of ventricular				
1	fibrillation.				
C, Oda N, Kihara Y	inormation.				
c, oda 11, ismaia 1					
Sairaku A, Yoshida Y,	Ablation of atrial fibrillation in	Int J Cardiol	168	5273-5276	2013
Nakano Y, Kihara Y	Brugada syndrome patients with an		33	2.5 52.6	[ ]
	implantable cardioverter				
	defibrillator to prevent				
	inappropriate shocks resulting				
	from rapid atrial fibrillation.				

JOURNAL OF THE AMERICAN COLLEGE OF CARDIOLOGY

© 2014 BY THE AMERICAN COLLEGE OF CARDIOLOGY FOUNDATION
PUBLISHED BY ELSEVIER INC.

VOL. ■, NO. ■, 2014 ISSN 0735-1097/S36.00 http://dx.doi.org/10.1016/Ljacc.2014.94.023

#### EDITORIAL COMMENT

# Importance of Clinical Analysis in the Era of New Technology in Molecular Genetic Screening\*

Q1 Wataru Shimizu, MD, РнD

or the past 2 decades, a number of inherited cardiac arrhythmia syndromes have been shown to be linked to mutations in genes encoding cardiac ion channels or other membrane components. These include congenital and acquired long-QT syndrome (LQTS), Brugada syndrome (BrS), progressive cardiac conduction defect, Lenegre disease, catecholaminergic polymorphic ventricular tachycardia (CPVT), short-QT syndrome, early repolarization syndrome, and familial atrial fibrillation (AF) (1). In congenital LQTS, 13 genotypes have been identified in approximately 75% of subjects with clinically diagnosed congenital LQTS (1,2), and genotypephenotype correlations have been investigated in detail. Thus, genetic testing is now a gold standard for diagnosing congenital LQTS, enabling risk stratification of cardiac events and better patient management (1). Mutations in the RyR2 gene or calsequestrin gene can be identified in approximately 60% of typical patients with CPVT associated with bidirectional and/or multifocal ventricular tachycardia (1,2). However, the yield associated with disease-specific genetic testing is far short of 100%, even in congenital LQTS or CPVT. Moreover, causative mutations have been identified in a small number of patients with other inherited arrhythmia syndromes (i). The yield of disease-specific genetic testing is only 20% to 30% in BrS and is still unknown in progressive cardiac conduction defect, short-QT syndrome, early repolarization syndrome, and familial AF (1,2).

SCN5A (>75% of genotyped cases); however, a worldwide cohort reported that SCN5A accounts only for 11% to 28% of clinically diagnosed patients with BrS (4). Moreover, the majority of mutations were found in a single family or a small number of families. Therefore, a genotype-phenotype correlation is not available in most cases (1,5).

The relatively lower yield of disease-specific genetic testing except for congenital LQTS or CPVT is due mainly to the technology of genetic testing. Candidate gene analysis has long been used to identify a causative mutation in a gene, which is expected to relate to the pathophysiology of each inherited arrhythmia syndrome, such as cardiac ion channel genes. However, causative mutations do not always

In BrS, the first mutation was identified in an alpha

subunit of a sodium channel gene, SCN5A, in 1998 (2).

Subsequently, genetic studies have identified 13

responsible genes on chromosomes 1, 3, 7, 10, 11, 12,

17, and 19 (1). Among 13 genotypes, more than 300

mutations have been identified in the major player,

involve genes of ion channels or membrane components. Innovative advances in molecular genetic testing are overcoming this issue with the advent of more powerful molecular genetic screening tools, including genome-wide association study (GWAS) using gene array, as well as targeted, whole-exome and whole-genome next-generation sequencing techniques.

Several recent GWASs have disclosed significant association of numerous loci in some genes with electrocardiographic markers or arrhythmia syndromes. Arking et al. (6) first identified NOS1AP (CAPON), a regulator of neuronal nitric oxide syn-

thase, as a gene that is significantly associated with

QT-interval variation in a general population derived

from 3 cohorts (6). Subsequently, 2 groups conducted

a meta-analysis of the GWAS and observed associa-

tions of single-nucleotide polymorphisms (SNPs) in

Q4

<sup>\*</sup> Editorials published in the *Journal of the American College of Cardiology* reflect the views of the authors and do not necessarily represent the views of *JACC* or the American College of Cardiology.

From the Department of Cardiovascular Medicine, Nippon Medical School, Tokyo, Japan. Dr. Shimizu is supported in part by a Research Grant for the Cardiovascular Diseases (H24-033) from the Ministry of Health, Labour and Welfare, Japan.

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207 208

209 210

211

212

213

214

215

216

Clinical Analysis and Molecular Genetic Screening

151

152 153

154

155

156

157

158

160

161

162

159 Q2

several genes in addition to NOS1AP with QT interval, suggesting that these genes are candidate genes for LQTS or sudden cardiac death (7,8). Several GWASs also identified associations of SNPs in several genes, including SCN10A, with cardiac conduction parameters, such as QRS duration and PR interval (9-11). Regarding associations with cardiac arrhythmias, some SNPs in several genes, including ZFHX3 and KCNN3, have been reported to be associated with AF (12-14). The association of a SNP in CXADR with ventricular fibrillation in acute myocardial infarction also has been reported (15). However, no responsible mutations have thus far been reported in these candidate genes in patients with clinically diagnosed inherited arrhythmia syndromes, such as congenital LQTS, familial AF, and familial conduction abnormalities.

Bezzina et al. (16) recently conducted a GWAS in 312 patients with BrS with type 1 electrocardiographic pattern and 1,115 controls. They detected 2 significant association signals at the SCN10A intronic locus (rs10428132) in chromosome 3p22 and near the HEY2 gene (rs9388451) in chromosome 6q22 with BrS. SCN10A, which encodes the sodium channel isoform Nav1.8, was originally reported as highly expressed in cardiac neurons. Recent evidence indicates that SCN10A also is expressed in the working myocardium and the specialized conduction system, indicating a possible role for Nav1.8 in cardiac electrical function. HEY2 is involved in patterning Nav1.5 (SCN5A) expression across the ventricular wall. In an experiment using HEY2 knockout mouse, Bezzina et al. (16) suggested that loss of HEY2 might affect the transmural expression gradient of sodium channel implicated in BrS.

In this issue of the *Journal*, Hu et al. (17) report on a clinical analysis and direct sequencing of *SCN10A* and all known BrS genes in 150 unrelated patients with BrS and 17 family members, as well as more than 200 ethnically matched healthy controls. They identified 17 *SCN10A* mutations in 25 of 150 patients with BrS (a yield of 16.7%). Twenty-three of the 25 (92.0%) displayed overlapping phenotypes, such as early repolarization syndrome and cardiac conduction

defect. Patients with BrS with *SCN10A* mutations were more symptomatic and displayed significantly longer PR and QRS intervals than *SCN10A*-negative patients with BrS. Heterologous coexpression of *SCN10A* mutants (R14L and R1268Q) with wild-type *SCN5A* caused 79.4% and 84.4% reductions in sodium channel current, strongly implicating *SCN10A* as a major susceptibility gene for BrS. This study provides the first major step forward in more than 16 years in the identification of new BrS susceptibility genes, advancing the yield for detection of a genotype to more than 50%.

New molecular genetic screening technologies, such as GWAS and whole-exome and whole-genome next-generation sequencing, are promising tools for identifying new candidate genes responsible for inherited arrhythmia syndromes. However, no responsible mutations have been reported in the candidate genes identified by GWAS in patients with clinically diagnosed inherited arrhythmia syndromes. To the best of my knowledge, the SCN10A is the first gene to be suggested as a BrS susceptibility gene by both GWAS and direct sequencing techniques. Direct sequencing using the Sanger technique combined with a detailed clinical analysis, including genotype-phenotype correlation and functional expression studies, continue to play an important role in molecular genetic testing, even in the new era in which gene arrays and next-generation sequencing are available. The importance of a detailed clinical analysis including genotype-phenotype correlation as well as functional expression studies cannot be overemphasized. Even in GWAS and whole-genome or whole-exome studies, clinical misdiagnosis can contribute to confounding genetic noise. A detailed, precise clinical diagnosis is therefore a prerequisite for the identification of new potential candidate genes.

REPRINT REQUESTS AND CORRESPONDENCE: Dr. Wataru Shimizu, Nippon Medical School, Department of Cardiovascular Medicine, 1-1-5, Sendagi Bunkyo-ku, Tokyo 113-8603, Japan. E-mail: wshimizu@nms.ac.jp.

#### REFERENCES

- **1.** Shimizu W. Update of diagnosis and management in inherited cardiac arrhythmias. Circ J 2013; 77:2867–72.
- 2. Ackerman MJ, Priori SG, Willems S, et al. HRS/ EHRA expert consensus statement on the state of genetic testing for the channelopathies and cardiomyopathies this document was developed as a partnership between the Heart Rhythm Society
- (HRS) and the European Heart Rhythm Association (EHRA). Heart Rhythm 2011;8:1308-39.
- Chen Q, Kirsch GE, Zhang D, et al. Genetic basis and molecular mechanisms for idiopathic ventricular fibrillation. Nature 1998;392:293-6.
- 4. Kapplinger JD, Tester DJ, Alders M, et al. An international compendium of mutations in the SCN5A-encoded cardiac sodium channel in
- patients referred for Brugada syndrome genetic testing. Heart Rhythm 2010;7:33–46.
- **5.** Shimizu W. Clinical features of Brugada syndrome. J Arrhythmia 2013;29:65-70.
- **6.** Arking DE, Pfeufer A, Post W, et al. A common genetic variant in the NOS1 regulator NOS1AP modulates cardiac repolarization. Nat Genet 2006;38:644-51.

#### ARTIQUE IN PRESS

JACC VOL. 📕, NO. 🗐, 2014

. 2014: **3** - **3** 

Clinical Analysis and Molecular Genetic Screening

Shimizu

- 7. Newton-Cheh C, Eijgelsheim M, Rice KM, et al. Common variants at ten loci influence QT interval duration in the QTGEN Study. Nat Genet 2009;41: 399-406.
- **8.** Pfeufer A, Sanna S, Arking DE, et al. Common variants at ten loci modulate the QT interval duration in the QTSCD Study. Nat Genet 2009;41:407-14.
- **9.** Chambers JC, Zhao J, Terracciano CM, et al. Genetic variation in SCN10A influences cardiac conduction. Nat Genet 2010;42:149–52.
- **10.** Pfeufer A, van Noord C, Marciante KD, et al. Genome-wide association study of PR interval. Nat Genet 2010;42:153-9.
- **11.** Sotoodehnia N, Isaacs A, de Bakker PI, et al. Common variants in 22 loci are associated with

- QRS duration and cardiac ventricular conduction. Nat Genet 2010;42:1068-76.
- **12.** Benjamin EJ, Rice KM, Arking DE, et al. Variants in ZFHX3 are associated with atrial fibrillation in individuals of European ancestry. Nat Genet 2009:41:879–81.
- **13.** Ellinor PT, Lunetta KL, Glazer NL, et al. Common variants in KCNN3 are associated with lone atrial fibrillation. Nat Genet 2010;42:240-4.
- **14.** Ellinor PT, Lunetta KL, Albert CM, et al. Metaanalysis identifies six new susceptibility loci for atrial fibrillation. Nat Genet 2012;44:670-5.
- **15.** Bezzina CR, Pazoki R, Bardai A, et al. Genome-wide association study identifies a susceptibility locus at 21q21 for ventricular

- fibrillation in acute myocardial infarction. Nat Genet 2010:42:688-91.
- **16.** Bezzina CR, Barc J, Mizusawa Y, et al. Common variants at SCN5A-SCN1OA and HEY2 are associated with Brugada syndrome, a rare disease with high risk of sudden cardiac death. Nat Genet 2013; 45:1044-9.
- **17.** Hu D, Barajas-Martínez H, Pfeiffer R, et al. Mutations in SCN10A responsible for a large fraction of Brugada syndrome cases. J Am Coll Cardiol 2014;64:000–000.

**KEY WORDS** Brugada syndrome, direct sequencing, genetic study, GWAS, sudden death

### **Original Article**

## Genetic Characteristics of Children and Adolescents With Long-QT Syndrome Diagnosed by School-Based Electrocardiographic Screening Programs

Masao Yoshinaga, MD, PhD; Yu Kucho, MD; Jav Sarantuya, MD, PhD; Yumiko Ninomiya, MD; Hitoshi Horigome, MD, PhD; Hiroya Ushinohama, MD, PhD; Wataru Shimizu, MD, PhD; Minoru Horie, MD, PhD

Background—A school-based electrocardiographic screening program has been developed in Japan. However, few data are available on the genetic characteristics of pediatric patients with long-QT syndrome who were diagnosed by this program. Methods and Results—A total of 117 unrelated probands aged ≤18 years were the subjects who were referred to our centers for genetic testing. Of these, 69 subjects diagnosed by the program formed the screened group. A total of 48 subjects were included in the clinical group and were diagnosed with long-QT syndrome–related symptoms, familial study, or by chance. Mutations were classified as radical, of high probability of pathogenicity, or of uncertain significance. Two subjects in the clinical group died. Genotypes were identified in 50 (72%) and 23 (48%) of subjects in the screened and clinical groups, respectively. Of the KCNQ1 or KCNH2 mutations, 31 of 33 (94%) in the screened group and 15 of 16 (94%) in the clinical group were radical and of high probability of pathogenicity. Prevalence of symptoms before (9/69 versus 31/48; P<0.0001) and after (12/69 versus 17/48; P=0.03) diagnosis was significantly lower in the screened group when compared with that in the clinical group although the QTc values, family history of long-QT syndrome, sudden death, and follow-up periods were not different between the groups.

Conclusions—These data suggest that the screening program may be effective for early diagnosis of long-QT syndrome that may allow intervention before symptoms. In addition, screened patients should have follow-up equivalent to clinically identified patients. (Circ Arrhythm Electrophysiol. 2014;7:107-112.)

**Key Words:** diagnosis ■ genetic testing ■ QT interval electrocardiography ■ screening

Congenital long-QT syndrome (LQTS) is a genetic disorder characterized by delayed repolarization and by a long-QT interval on 12-lead ECGs. Although many patients do not have symptoms, the hallmark of the condition is syncope or sudden death because of torsade de pointes. <sup>1,2</sup> To date, 13 genes have been identified.<sup>3,4</sup> There have been many reports on the clinical and genetic backgrounds of patients with LQTS. However, these were mainly based on data collected from patients who had LQTS-related symptoms or familial studies and from combined adult and pediatric populations.<sup>2,5-9</sup>

#### Clinical Perspective on p 112

A nationwide school-based ECG screening program for heart diseases in first, seventh, and 10th graders in Japan has revealed children and adolescents with prolonged QT intervals. The prevalence of subjects with prolonged QT intervals was ≈1:1200 in the seventh grade. <sup>10</sup> Differences in clinical

characteristics between patients who were screened by the program and those who visited hospitals with symptoms have been previously reported.<sup>11</sup> However, before this study, few data have been reported about the genetic characteristics of pediatric patients who were diagnosed by ECG screening programs and whose genetic testing was performed.<sup>12–14</sup> In addition, because of a lack of reports containing large numbers of patients who were screened alongside genetic testing, it is unclear whether screened subjects have similar mutations of a high possibility of pathogenesis to those who have LQTS-related symptoms.

From the genetic testing viewpoint, the few percentage background rate of the rare *KCNQ1* and *KCNH2* nonsynonymous single nucleotide variants among healthy individuals has lessened the ability to distinguish rare pathogenic mutations from similarly rare, yet presumably innocuous, variants. <sup>15,16</sup> Novel mutations have been found in every study, <sup>17,18</sup> but it is difficult to perform electrophysiological studies for

Received March 7, 2013; accepted November 11, 2013.

The Data Supplement is available at http://circep.ahajournals.org/lookup/suppl/doi:10.1161/CIRCEP.113.000426/-/DC1

Correspondence to Masao Yoshinaga, MD, PhD, Department of Pediatrics, National Hospital Organization Kagoshima Medical Center, 8-1 Shiroyama-cho, Kagoshima 892-0853, Japan. E-mail m-yoshi@biscuit.ocn.ne.jp

© 2013 American Heart Association, Inc.

Circ Arrhythm Electrophysiol is available at http://circep.ahajournals.org

DOI: 10.1161/CIRCEP.113.000426

From the Department of Pediatrics, National Hospital Organization Kagoshima Medical Center, Kagoshima, Japan (M.Y., Y.K., Y.N.); Department of Molecular Biology and Genetics, School of Bio-medicine, Health Sciences University of Mongolia, Ulaanbaatar, Mongolia (J.S.); Department of Child Health, Faculty of Medicine, University of Tsukuba, Tsukuba, Japan (H.H.); Department of Cardiology, Fukuoka Children's Hospital and Medical Center for Infectious Diseases, Fukuoka, Japan (H.U.); Department of Cardiovascular Medicine, Nippon Medical School, Tokyo, Japan (W.S.); and Department of Cardiovascular and Respiratory Medicine, Shiga University of Medical Science, Otsu, Japan (M.H.).

each novel mutation except in large laboratories. Recently, an algorithm designed to guide the interpretation of genetic testing results for *KCNQ1* and *KCNH2* has been developed.<sup>16</sup>

Thus, the aim of the present study was to determine the genetic characteristics of pediatric patients with LQTS who were diagnosed by a school-based screening program and whose genetic testing was performed and to compare results with subjects who visited hospitals because of the presence of LQTS-related symptoms, familial history, or who were diagnosed by chance.

#### Methods

#### **Study Population**

The study population included 117 unrelated probands ≤18 years of age who were referred to the Department of Pediatrics, Kagoshima University Hospital, Japan, between November 1993 and March 2005 or to the National Hospital Organization Kagoshima Medical Center from April 2005 to December 2012 for genetic evaluation. The population included 69 subjects who were screened by a school-based ECG screening program (Table 1). In the present study, LQTS-related symptoms were defined as syncope, aborted cardiac arrest, or sudden cardiac death at <30 years old. Subjects were divided into 2 groups on the basis of index events: subjects who were diagnosed by the school-based ECG screening program (screened group) and those who visited hospitals because of the presence of symptoms and family history or who were diagnosed by chance (clinical group; Table 1).

## Diagnosis of LQTS and Screening of QT Intervals in the School-based ECG Screening Program

The present study was a retrospective study, and diagnosis of LQTS and screening for prolonged QT intervals was based on the judgment of the chief medical doctors in each hospital or doctors who

Table 1. Characteristics of Probands

Subjects	Screened Group	Clinical Group	P Value
No. of subjects	69	48	
Age at diagnosis*	10.4±3.4	$7.4 \pm 6.0$	0.04
Age at diagnosis (median and range)	12.2 (6.2–18.8)	8.9 (0-17.2)	
Sex (men/women)	36/33	27/21	0.66
Mean QT interval, ms*	466±51	442±83	0.09
Mean RR interval, ms*	887±170	802±261	0.09
QTc (Bazett), ms <sup>1/2*</sup>	496±40	502±52	0.84
History of symptoms†	9 (13%)	31 (65%)	< 0.0001
Syncope	9	28	
Aborted cardiac arrest	0	7‡	
Family history of long-QT syndrome†	27 (39%)	18 (38%)	>0.99
Family history of sudden death†	5 (7%)	7 (15%)	0.23
Follow-up periods*	4.6±4.9	5.2±5.7	0.36
Symptoms after diagnosis†	12 (17%)	17 (35%)	0.03
Syncope	12	17	
Aborted cardiac arrest	0	2§	
Sudden cardiac death	0	2#	

<sup>\*</sup>The mean value±SD.

#0f each 2 subjects with ACA or sudden cardiac death (SCD), all 4 subjects experienced syncope and ACA or SCD.

participated in the program in each area. Many Japanese cardiologists use a scoring system published in 199319 and recently for the final diagnosis of LQTS. To screen subjects with prolonged QT intervals in the program, the Japanese Society of Pediatric Cardiology and Cardiac Surgery recommended that children and adolescents be screened when they have a QTc value, using Bazett formula, of ≥450 ms at a heart rate of <75 beats per minute or a OTc≥500 ms at a hours of ≥75 beats per minute.20 Bazett formula overcorrects the QT interval at high heart rates. Pediatric cardiologists who participated in the program used age- and sex-specific criteria using an exponential formula (QT/RR<sup>0.31</sup>)<sup>21</sup> or Fridericia formula.<sup>22</sup> In the screening program, cardiologists use computer-based QTc values as a reference because all ECG machines used in Japan are generally equipped with a function for automated measurement of OT intervals. However, manual measurement using the tangent method is usually applied to obtain QT intervals in Japan.22,23

#### **Genetic Testing**

Referral for genetic testing was based on the opinion of the chief medical doctors in the present study. Pediatric cardiologists in the present study recommended genetic testing based on the following criteria: (1) for a patient in whom they had established a strong clinical index of suspicion for LQTS based on examination of the patient's clinical history, family history, and expressed ECG phenotype or; (2) for an asymptomatic patient with QT prolongation in the absence of other clinical conditions that might prolong the QT interval, as detailed in the recent consensus recommendation report.<sup>24</sup>

Genomic DNA was isolated from blood after obtaining written informed consent. Genetic screening for all exons of KCNQ1, KCNH2, SCN5A, KCNE1, KCNE2, KCNJ2, and CAV3 was reperformed for the present study using polymerase chain reaction and direct DNA sequencing. When a patient was suspected to have Timothy syndrome, which is a multisystem disorder characterized by cardiac (QT prolongation and sometimes congenital heart diseases), hand/foot, facial, and neurodevelopmental features, the exons of CACNAIC were amplified. When a patient had a prolonged QT interval and hyperaldosteronism, the exon of KCNJ5 was amplified. The exons of ANKB, SCN4B, AKAP9, and SNTA1 were not analyzed because of a lack of reported cases of these mutations in the Japanese population. Genomic DNA was isolated using a QIAamp DNA Blood Midi Kit (Qiagen, Gaithersburg, MD). Polymerase chain reaction products were purified using AMPure (Beckman Coulter, Brea, CA). After treating with BigDye Terminator version 1.1 Cycle Sequence Kit (ABI, Warrington, United Kingdom) and BigDye X Terminator, direct sequencing was performed by a genetic analyzer, ABI3130x1 Genetic Analyzer (ABI). The study was approved by the Ethics Committee of the Kagoshima University Hospital between November 1993 and March 2005 and the National Hospital Organization Kagoshima Medical Center from April 2005.

Nucleotide changes reported as single nucleotide polymorphisms  $^{18,25}$  were excluded from mutation analysis in the present study. However, amino acid changes of G643S in  $KCNQI^{26}$  and D85N in  $KCNEI^{27}$  were included in the present study because previous reports have shown that these mutations are associated with an  $\approx 30\%$  reduction in potassium channel currents.  $^{26,27}$  When multiple mutations were present, each mutation was counted in each genotype.

#### **Mutations of High Probability of Pathogenicity**

Mutations of a high probability of pathogenicity were based on data published by Giudicessi et al. <sup>16</sup> Radical mutations included splicesite, nonsense, frame-shift, and insertion/deletions. <sup>16</sup> Mutations of a high probability of pathogenicity in the present study were defined as those present in the subunit assembly domain of the C-terminal of *KCNQ1*, the Per-Arnt-Sim domain, Per-Arnt-Sim—associated C-terminal domain, and the cyclic nucleotide—binding domain of *KCNH2*. Mutations present in the transmembrane/linker/pore and C-terminal regions of *KCNQ1* and the transmembrane/linker/pore regions of *KCNH2* were also defined as those of a high probability of pathogenicity. <sup>16</sup> Remaining mutations were defined as those of uncertain significance.

<sup>†</sup>Number of subjects and percentage in parenthesis.

<sup>‡0</sup>f 7 subjects with aborted cardiac arrest (ACA), 4 experienced both syncope and ACA.

#### **Statistical Analysis**

Differences in the mean values and prevalence values were examined using the Mann–Whitney U test and Fisher exact probability test, respectively. Tukey multiple comparison test was used to assess differences in the mean QTc values among first, seventh, and 10th graders. Statistical analysis was performed using IBM SPSS Statistics version 21.0 (IBM Japan, Ltd, Tokyo, Japan). A 2-tailed P value of <0.05 was considered statistically significant.

#### Results

#### **Population**

Characteristics of the 117 subjects, including 69 screened and 48 clinical patients, are shown in Table 1. Of the 48 subjects included in the clinical group, 36 were diagnosed with LQTSrelated symptoms, 6 were diagnosed by familial study, and 6 were diagnosed by chance. Subjects who were diagnosed by chance included those who visited hospitals for medical checks and for examination of heart murmurs at 1 month (4 patients), those who had been followed with Kawasaki disease (1 patient), and as Ehlers-Donlos syndrome (1 patient). There were no differences in sex, mean QTc values, family history of LQTS, family history of sudden death, or follow-up period between the screened and clinical groups. The mean age was lower in the clinical group when compared with the screened group (P=0.04). Prevalence of subjects having LQTS-related symptoms before and after diagnosis was significantly lower in the screened group when compared with that in the clinical group (P<0.001 and P=0.03, respectively). Symptoms before and after diagnosis in the screened group were all syncope. Of 117 subjects, 2 subjects in the clinical group died. A girl had a history of aborted cardiac arrest at 2 months, and died suddenly in her sleep at 5 years of age. An 11-year-old boy had frequent symptoms and died suddenly during class. Genetic analysis failed to show the presence of any of the mutations analyzed in this study. The treatment of subjects with symptoms during follow-up period is shown in Table I in the onlineonly Data Supplement.

#### **Mutations Determined in the Present Study**

The yield of genetic testing in the present study by QTc values using Bazett formula is shown in Table II in the Data Supplement. The data show that there was no difference in yield between subjects with a QTc<500 ms and those with a OTc≥500 ms in both screened and clinical groups in the present study. Of 50 subjects who were screened and whose mutations were identified, 29, 18, and 3 subjects were screened in the first, seventh, and 10th grade, respectively. Their QTc values using Bazett formula were 491±35, 503±43, and 500±49 ms, respectively. There were no differences in QTc values among the screened periods. Of 117 subjects, mutations were found in 50 of 69 (72%) screened and 23 of 48 (48%) clinical subjects (Table III in the Data Supplement). The prevalence of LQT1, LQT2, and LQT3 between the 2 groups was not different. LQTS-related mutations in the present study are summarized in Table IV in the Data Supplement.

#### **Genetic Characteristics of Subjects**

All mutations found in KCNQ1 from 18 mutations in the screened and 9 mutations in the clinical groups were located

in regions of a high probability of pathogenicity (Table 2; Figure 1A and 1B). In the screened group, 8 mutations were located in the transmembrane/linker/pore regions and 10 were present in the C-terminal regions (Figure 1A). Three mutations were radical and 1 mutation was present in the subunit assembly domain. About the association between locations of mutation and the presence or absence of LQTS-related symptoms, 14 (78%) of 18 mutations were associated with the presence of symptoms in probands and family members, including 4 (22%) with family history of sudden death in the screened group. Eight of 9 mutations in the clinical group were associated with the presence of symptoms, and the remaining mutation was found in a subject who was diagnosed by a familial study.

Among mutations yielded in *KCNH2*, 13 (87%) of 15 mutations in the screened group and 6 (86%) of 7 mutations in the clinical group were located in regions of a high probability of pathogenicity (Table 2; Figure 2A and 2B). In the screened group, 1 mutation was both radical and present in the cyclic nucleotide–binding domain. Another 5 mutations were radical, and 1 each was present in Per-Arnt-Sim and Per-Arnt-Sim–associated C-terminal regions. However, only 4 (31%) of 13 mutations were associated with the presence of LQTS-related symptoms in probands or family members in the screened group. In the clinical group, 6 (86%) of 7 mutations were associated with the presence of symptoms in probands and family members, and the remaining mutation was found in a subject by ECG screening during a medical check-up at 1 month.

#### Discussion

Mutations in subjects with LQTS who were diagnosed by school-based ECG screening programs were mostly of high possibility of pathogenicity, similar to clinical subjects. Clinical background, such as QTc values, family history of LQTS, or sudden death, and follow-up periods, was not different between the 2 groups. However, prevalence of symptoms before and after diagnosis in the screened group was significantly lower when compared with the clinical group.

Table 2. Number of Patients With Mutations at High Risk in Each Group

Genes	Mutations	Screened Group	Clinical Group	<i>P</i> Value
KCNQ1*	Radical mutation†	4	1	>0.99
	High probability‡	14	8	0.59
	Variants of uncertain significance	0	0	>0.99
KCNH2†	Radical mutation† and high probability‡	1	1	0.53
	Radical mutation†	5	1	0.66
	High probability‡	7	4	0.73
	Variants of uncertain significance	2	1	>0.99

<sup>\*</sup>Variants of uncertain significance include mutations other than radical or of high probability of pathogenicity.

Downloaded from http://circep.ahajournals.org/ by MASAO YOSHINAGA on March 6, 2014

<sup>†</sup>Radical mutations include splice-site, nonsense, frame-shift, and insertion/

<sup>‡</sup>Mutations of high probability include subunit assembly domain, transmembrane/linker/pore, and C-terminal regions of *KCNQ1*, and the Per-Arnt-Sim (PAS) domain, PAS-associated C-terminal domain, the cyclic nucleotide–binding domain, transmembrane/linker/pore region of *KCNH2*.

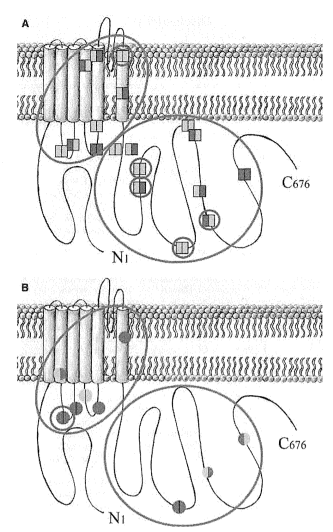
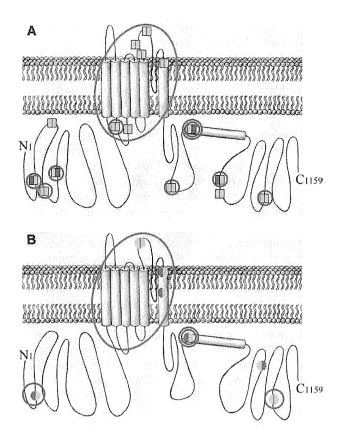


Figure 1. Topological depiction of KCNQ1 in the present study in the screened (A) and clinical (B) groups. Mutations found in the screened group are shown as boxes (A) and those in the clinical group as circles (B). Each box or circle is divided into 2 parts: left and right sides. Each part represents the presence or absence of long-QT syndrome-related symptoms in probands (left) and family members (right), respectively. Green, brown, and red colors symbolize no symptoms, syncope or aborted cardiac arrest, and sudden death, respectively. Bold red circles surrounding mutations represent radical mutations. A bold blue circle represents subunit assembly domain. Two big purple circles symbolize locations of transmembrane/linker/pore and C-terminal regions of KCNQ1.

These data suggest that screening programs may be effective for early diagnosis of LQTS and prevention of symptoms, and that screened patients should be followed similar to clinical patients.

Clinical and genetic backgrounds of patients with LQTS have been reported widely for infants, children, adolescents, and adults. These data were mostly based on symptomatic probands and family members. Few data are available on the genetic background of subjects who were diagnosed by ECG screening programs. Schwartz et al<sup>12</sup> reported that LQTS-related mutations were identified in 16 neonates of 43 080 who underwent neonatal ECG screening; 8 *KCNQ1*, 5 *KCNH2*, and 1 each of *KCNE1* and *KCNE2*. One infant had a digenic mutation of *KCNQ1* and *KCNH2*.



**Figure 2.** Topological depiction of *KCNH2* in the present study in the screened (**A**) and clinical (**B**) groups. Explanations of symbols and shapes are the same as in Figure 1. Bold blue circles surrounding mutations in this figure represent Per-Arnt-Sim (PAC), PAC-associated C-terminal, and cyclic nucleotide-binding domains, respectively, from the **left** side. A big purple circle symbolizes locations of transmembrane/linker/pore regions.

A school-based ECG screening program for heart diseases was initiated in 1994 for first, seventh, and 10th graders in Japan. The program screened subjects with QT prolongation. However, few studies have confirmed the genetic background in these screened subjects. <sup>13,14</sup> Hayashi et al<sup>13</sup> reported that mutations were identified in 3 subjects with high or intermediate probabilities of LQTS using Schwartz criteria from 7961 school children; all 3 mutations were present in *KCNH2*. Yasuda et al<sup>14</sup> reported that *KCNQ1* mutations were found in 8 of 13 pediatric patients and that 7 of 8 patients were diagnosed by the ECG screening program.

In the present study, a relatively large number of subjects, who were diagnosed by ECG screening programs accompanied by genetic testing, were included. The clinical backgrounds of the screened subjects, such as QTc values, family history of LQTS, or sudden death, were similar to clinical subjects. All 16 mutations in the *KCNQ1* gene in the screened group were radical or of high probability of pathogenicity similar to the clinical group. The ratio of mutations of radical and of high probability of pathogenicity in the *KCNH2* gene in the screened group (13/15; 87%) was remarkably similar to that in the clinical group (6/7; 86%). These data suggest that pediatricians, who asked for genetic testing in the present study, chose patients with similar clinical backgrounds in both groups, and that demand for genetic testing was more

Downloaded from http://circep.ahajournals.org/ by MASAO YOSHINAGA on March 6, 2014

prevalent in screened patients when compared with clinical patients when ECG screening was developed in Japan.

Conversely, prevalence of symptoms before and after diagnosis was significantly lower in the screened group when compared with that in the clinical group. A low prevalence of symptoms before diagnosis suggests that the ECG screening program is effective for early diagnosis of LQTS. The reason for low prevalence of symptoms after diagnosis in the screened group is uncertain. Doctors may recommend pediatric patients with LQTS and their parents adopt changes to their lifestyles, for example, not doing vigorous exercise, not swimming a lap, and not diving, <sup>28</sup> in both the screened and clinical subjects. The precise reason remains to be clarified.

The reason for no difference in the prevalence of family history between the screened and clinical groups is unclear. The authors posit that even now in Japan the general population may not be familiar with LQTS, and that the parents in the present study did not think that syncope in their children was a serious condition. In addition, they may have been unaware that LQTS is an inherited disease. The reason of the high prevalence of family history of LQTS in the screened group is also unclear. The authors speculate that doctors did not ask the parents (grandparents of the probands in the present study) to perform familial studies 2 or 3 decades ago, when parents of the probands of the present study and their family members experienced symptoms at younger ages; however, no data were obtained addressing this hypothesis from the families.

There are some limitations of the current study. First, we did not discuss subjects with the SCN5A gene. One fourth of pediatric patients with LQTS had the SCN5A gene. We need similar algorithms designed to guide the interpretation of genetic testing results for the SCN5A mutation and to determine the possibility of pathogenesis in patients with SCN5A in the future. Second, the clinical group showed a low rate (48%) of genotypic determination. We could not find mutations in 2 cases of death in the present study. The reasons for this are unclear. One potential reason was that we did not screen copy number variations in genes associated with LQTS.<sup>29,30</sup> Eddy et al<sup>29</sup> and Barc et al<sup>30</sup> reported that 3 of 26 (12%) and 3 of 93 (3%) unrelated mutation-negative probands showed copy number variations, indicating that some mutation-negative patients may have copy number variations. Another reason may be that numerous previously undetected mutations exist in symptomatic patients.

In conclusion, mutations in subjects with LQTS who were diagnosed by screening programs had a high probability of pathogenicity similar to clinical subjects. Clinical backgrounds were not different although the prevalence of symptoms before and after diagnosis in the screened group was significantly lower when compared with that in the clinical group. These data suggest that the school-based screening program may be effective for early diagnosis of LQTS and prevention of symptoms, and that screened patients should have follow-up equivalent to clinical patients.

#### **Sources of Funding**

This study was partly supported by a Health and Labour Sciences Research Grant from the Ministry of Health, Labour and Welfare of Japan (Research on Intractable Diseases [H22-032] and [H24-033]).

#### **Disclosures**

None.

#### References

- Schwartz PJ, Crotti L, Insolia R. Long-QT syndrome: from genetics to management. Circ Arrhythm Electrophysiol. 2012;5:868–877.
- Priori SG, Schwartz PJ, Napolitano C, Bloise R, Ronchetti E, Grillo M, Vicentini A, Spazzolini C, Nastoli J, Bottelli G, Folli R, Cappelletti D. Risk stratification in the long-QT syndrome. N Engl J Med. 2003;348:1866–1874.
- Ackerman MJ, Mohler PJ. Defining a new paradigm for human arrhythmia syndromes: phenotypic manifestations of gene mutations in ion channeland transporter-associated proteins. Circ Res. 2010;107:457–465.
- 4. Yang Y, Yang Y, Liang B, Liu J, Li J, Grunnet M, Olesen SP, Rasmussen HB, Ellinor PT, Gao L, Lin X, Li L, Wang L, Xiao J, Liu Y, Liu Y, Zhang S, Liang D, Peng L, Jespersen T, Chen YH. Identification of a Kir3.4 mutation in congenital long QT syndrome. Am J Hum Genet. 2010;86:872–880.
- Spazzolini C, Mullally J, Moss AJ, Schwartz PJ, McNitt S, Ouellet G, Fugate T, Goldenberg I, Jons C, Zareba W, Robinson JL, Ackerman MJ, Benhorin J, Crotti L, Kaufman ES, Locati EH, Qi M, Napolitano C, Priori SG, Towbin JA, Vincent GM. Clinical implications for patients with long QT syndrome who experience a cardiac event during infancy. *J Am Coll Cardiol*. 2009;54:832–837.
- Goldenberg I, Moss AJ, Peterson DR, McNitt S, Zareba W, Andrews ML, Robinson JL, Locati EH, Ackerman MJ, Benhorin J, Kaufman ES, Napolitano C, Priori SG, Qi M, Schwartz PJ, Towbin JA, Vincent GM, Zhang L. Risk factors for aborted cardiac arrest and sudden cardiac death in children with the congenital long-QT syndrome. *Circulation*. 2008;117:2184–2191.
- Liu JF, Jons C, Moss AJ, McNitt S, Peterson DR, Qi M, Zareba W, Robinson JL, Barsheshet A, Ackerman MJ, Benhorin J, Kaufman ES, Locati EH, Napolitano C, Priori SG, Schwartz PJ, Towbin J, Vincent M, Zhang L, Goldenberg I; International Long QT Syndrome Registry. Risk factors for recurrent syncope and subsequent fatal or near-fatal events in children and adolescents with long QT syndrome. J Am Coll Cardiol. 2011;57:941–950
- Hobbs JB, Peterson DR, Moss AJ, McNitt S, Zareba W, Goldenberg I, Qi M, Robinson JL, Sauer AJ, Ackerman MJ, Benhorin J, Kaufman ES, Locati EH, Napolitano C, Priori SG, Towbin JA, Vincent GM, Zhang L. Risk of aborted cardiac arrest or sudden cardiac death during adolescence in the long-QT syndrome. *JAMA*. 2006;296:1249–1254.
- Sauer AJ, Moss AJ, McNitt S, Peterson DR, Zareba W, Robinson JL, Qi M, Goldenberg I, Hobbs JB, Ackerman MJ, Benhorin J, Hall WJ, Kaufman ES, Locati EH, Napolitano C, Priori SG, Schwartz PJ, Towbin JA, Vincent GM, Zhang L. Long QT syndrome in adults. *J Am Coll Cardiol*. 2007;49:329–337.
- Fukushige T, Yoshinaga M, Shimago A, Nishi J, Kono Y, Nomura Y, Miyata K, Imamura M, Shibata T, Nagashima M, Niimura I. Effect of age and overweight on the QT interval and the prevalence of long QT syndrome in children. Am J Cardiol. 2002;89:395–398.
- 11. Yoshinaga M, Nagashima M, Shibata T, Niimura I, Kitada M, Yasuda T, Iwamoto M, Kamimura J, Iino M, Horigome H, Seguchi M, Aiba S, Izumida N, Kimura T, Ushinohama H, Nishi J, Kono Y, Nomura Y, Miyata K. Who is at risk for cardiac events in young patients with long QT syndrome? Circ J. 2003;67:1007–1012.
- Schwartz PJ, Stramba-Badiale M, Crotti L, Pedrazzini M, Besana A, Bosi G, Gabbarini F, Goulene K, Insolia R, Mannarino S, Mosca F, Nespoli L, Rimini A, Rosati E, Salice P, Spazzolini C. Prevalence of the congenital long-QT syndrome. *Circulation*. 2009;120:1761–1767.
- 13. Hayashi K, Fujino N, Uchiyama K, Ino H, Sakata K, Konno T, Masuta E, Funada A, Sakamoto Y, Tsubokawa T, Nakashima K, Liu L, Higashida H, Hiramaru Y, Shimizu M, Yamagishi M. Long QT syndrome and associated gene mutation carriers in Japanese children: results from ECG screening examinations. Clin Sci (Lond). 2009;117:415–424.
- Yasuda K, Hayashi G, Horie A, Taketani T, Yamaguchi S. Clinical and electrophysiological features of Japanese pediatric long QT syndrome patients with KCNQ1 mutations. *Pediatr Int*. 2008;50:611–614.
- Kapa S, Tester DJ, Salisbury BA, Harris-Kerr C, Pungliya MS, Alders M, Wilde AA, Ackerman MJ. Genetic testing for long-QT syndrome: distinguishing pathogenic mutations from benign variants. *Circulation*. 2009;120:1752–1760.
- Giudicessi JR, Kapplinger JD, Tester DJ, Alders M, Salisbury BA, Wilde AA, Ackerman MJ. Phylogenetic and physicochemical analyses enhance

Downloaded from http://circep.ahajournals.org/ by MASAO YOSHINAGA on March 6, 2014

- the classification of rare nonsynonymous single nucleotide variants in type 1 and 2 long-QT syndrome. *Circ Cardiovasc Genet*. 2012;5:519–528.
- Tester DJ, Will ML, Haglund CM, Ackerman MJ. Compendium of cardiac channel mutations in 541 consecutive unrelated patients referred for long QT syndrome genetic testing. *Heart Rhythm*. 2005;2:507–517.
- Kapplinger JD, Tester DJ, Salisbury BA, Carr JL, Harris-Kerr C, Pollevick GD, Wilde AA, Ackerman MJ. Spectrum and prevalence of mutations from the first 2,500 consecutive unrelated patients referred for the FAMILION long OT syndrome genetic test. *Heart Rhythm*. 2009;6:1297–1303.
- Schwartz PJ, Moss AJ, Vincent GM, Crampton RS. Diagnostic criteria for the long QT syndrome. An update. Circulation. 1993;88:782–784.
- Baba K, Asai T, Kitada M, Kiyosawa N, Nagashima M, Haneda N, Baba K, Harada K, Honda S, Matsuoka S, Ura K. School-based cardiovascular screening. Guidelines for the interpretation of electrocardiograms in the screening program. *Pediatr Cardiol Card Surg*. 2006;22:503–513 (in Japanese). https://center6.umin.ac.jp/oasis/pccs/journal/journal/pdf/20062204/503.pdf. Accessed December 14, 2012.
- Aihoshi S, Yoshinaga M, Tomari T, Nakamura M, Nomura Y, Oku S, Haraguchi T, Osawa N, Miyata K. Correction of the QT interval in children. *Jpn Circ J*. 1995;59:190–197.
- Hazeki D, Yoshinaga M, Takahashi H, Tanaka Y, Haraguchi Y, Abe M, Koga M, Fukushige T, Nagashima M. Cut-offs for screening prolonged QT intervals from Fridericia's formula in children and adolescents. Circ J. 2010:74:1663–1669.
- 23. Sumitomo N. Correction of the QT interval in children.  $Circ\ J$ . 2010;74:1534–1535.
- 24. Ackerman MJ, Priori SG, Willems S, Berul C, Brugada R, Calkins H, Camm AJ, Ellinor PT, Gollob M, Hamilton R, Hershberger RE, Judge DP, Le Marec H, McKenna WJ, Schulze-Bahr E, Semsarian C, Towbin JA, Watkins H, Wilde A, Wolpert C, Zipes DP. HRS/EHRA expert consensus statement on the state of genetic testing for the channelopathies and cardiomyopathies. *Europace*. 2011;13:1077–1109.

- 25. Genetic mutation and inherited arrhythmias. The Molecular Cardiology Laboratories of the IRCCS Fondazione Salvatore Maugeri (Pavia, Italy) and the Cardiovascular Genetics Program of the New York University School of Medicine (New York, USA). http://www.fsm.it/cardmoc/. Accessed December 14, 2012.
- 26. Kubota T, Horie M, Takano M, Yoshida H, Takenaka K, Watanabe E, Tsuchiya T, Otani H, Sasayama S. Evidence for a single nucleotide polymorphism in the KCNQ1 potassium channel that underlies susceptibility to life-threatening arrhythmias. *J Cardiovasc Electrophysiol*. 2001;12:1223–1229.
- Nishio Y, Makiyama T, Itoh H, Sakaguchi T, Ohno S, Gong YZ, Yamamoto S, Ozawa T, Ding WG, Toyoda F, Kawamura M, Akao M, Matsuura H, Kimura T, Kita T, Horie M. D85N, a KCNE1 polymorphism, is a disease-causing gene variant in long QT syndrome. *J Am Coll Cardiol*. 2009:54:812–819.
- 28. Maron BJ, Chaitman BR, Ackerman MJ, Bayés de Luna A, Corrado D, Crosson JE, Deal BJ, Driscoll DJ, Estes NA 3rd, Araújo CG, Liang DH, Mitten MJ, Myerburg RJ, Pelliccia A, Thompson PD, Towbin JA, Van Camp SP; Working Groups of the American Heart Association Committee on Exercise, Cardiac Rehabilitation, and Prevention; Councils on Clinical Cardiology and Cardiovascular Disease in the Young. Recommendations for physical activity and recreational sports participation for young patients with genetic cardiovascular diseases. Circulation. 2004;109:2807–2816.
- Eddy CA, MacCormick JM, Chung SK, Crawford JR, Love DR, Rees MI, Skinner JR, Shelling AN. Identification of large gene deletions and duplications in KCNQ1 and KCNH2 in patients with long QT syndrome. Heart Rhythm. 2008;5:1275–1281.
- Barc J, Briec F, Schmitt S, Kyndt F, Le Cunff M, Baron E, Vieyres C, Sacher F, Redon R, Le Caignec C, Le Marec H, Probst V, Schott JJ. Screening for copy number variation in genes associated with the long QT syndrome: clinical relevance. J Am Coll Cardiol. 2011;57:40–47.

#### **CLINICAL PERSPECTIVE**

This study aimed to determine the genetic characteristics of 69 pediatric patients with long-QT syndrome who were diagnosed by a school-based screening program (screened group) and in whom genetic testing was performed. The screened group was compared with 48 subjects who visited hospitals because of the presence of long-QT syndrome—related symptoms, familial history, or who were diagnosed by chance (clinical group). A recently developed algorithm, designed to guide the interpretation of genetic testing results for *KCNQ1* and *KCNH2*, enabled us to classify the mutations as probably pathogenic or variant of uncertain significance. Using the algorithm, the authors found that of mutations yielded in *KCNQ1* or *KCNH2*, 31 of 33 (94%) mutations in the screened group and 15 of 16 (94%) mutations in the clinical group were radical and of high probability of pathogenicity. They also found that prevalence of symptoms before (*P*<0.0001) and after (*P*=0.03) diagnosis was significantly lower in the screened group when compared with that in the clinical group although the QTc values, family history of long-QT syndrome, sudden death, and follow-up periods were not different between the groups. Demand for genetic testing is now more prevalent in screened patients when compared with clinical patients because ECG screening was developed in Japan. This study may help to clarify the benefits of ECG screening. In addition, this study provides valuable genetic information and confirms that patients identified by ECG screening have the condition and are similar in many ways to those identified via a clinical setting.

# A Kir3.4 mutation causes Andersen-Tawil syndrome by an inhibitory effect on Kir2.1

Yosuke Kokunai, MD, PhD\* Tomohiko Nakata, MD\* Mitsuru Furuta, MD\* Souhei Sakata, PhD Hiromi Kimura, MD, PhD Takeshi Aiba, MD, PhD Masao Yoshinaga, MD, PhD Yusuke Osaki, MD Masayuki Nakamori, MD, PhD Hideki Itoh, MD, PhD Takako Sato, MD, PhD Tomoya Kubota, MD, PhD Kazushige Kadota, MD, PhD Katsuro Shindo, MD, PhD Hideki Mochizuki, MD, PhD

Wataru Shimizu, MD, PhD

Minoru Horie, MD, PhD Yasushi Okamura, MD,

PhD

Kinji Ohno, MD, PhD Masanori P. Takahashi, MD, PhD

Correspondence to Dr. Takahashi: mtakahas@neurol.med.osaka-u. ac.jp

#### Supplemental data at Neurology.org

#### **ABSTRACT**

Objective: To identify other causative genes for Andersen-Tawil syndrome, which is characterized by a triad of periodic paralysis, cardiac arrhythmia, and dysmorphic features. Andersen-Tawil syndrome is caused in a majority of cases by mutations in KCNJ2, which encodes the Kir2.1 subunit of the inwardly rectifying potassium channel.

Methods: The proband exhibited episodic flaccid weakness and a characteristic TU-wave pattern, both suggestive of Andersen-Tawil syndrome, but did not harbor KCNJ2 mutations. We performed exome capture resequencing by restricting the analysis to genes that encode ion channels/associated proteins. The expression of gene products in heart and skeletal muscle tissues was examined by immunoblotting. The functional consequences of the mutation were investigated using a heterologous expression system in Xenopus oocytes, focusing on the interaction with the Kir2.1 subunit.

Results: We identified a mutation in the KCNJ5 gene, which encodes the G-protein-activated inwardly rectifying potassium channel 4 (Kir3.4). Immunoblotting demonstrated significant expression of the Kir3.4 protein in human heart and skeletal muscles. The coexpression of Kir2.1 and mutant Kir3.4 in Xenopus oocytes reduced the inwardly rectifying current significantly compared with that observed in the presence of wild-type Kir3.4.

Conclusions: We propose that KCNJ5 is a second gene causing Andersen-Tawil syndrome. The inhibitory effects of mutant Kir3.4 on inwardly rectifying potassium channels may account for the clinical presentation in both skeletal and heart muscles. Neurology® 2014;82:1-7

#### **GLOSSARY**

cRNA = complementary RNA; LQT = long QT; SNP = single nucleotide polymorphism; SNV = single nucleotide variant.

Periodic paralysis is a heterogeneous disorder caused by mutations in several ion channel genes, including sodium, calcium, and potassium channels.<sup>1-3</sup> Andersen-Tawil syndrome is a form of periodic paralysis that is characterized by a triad of periodic muscle weakness, cardiac arrhythmia, and dysmorphic features. 4.5 Although dominantly inherited, its phenotypes are highly variable and its penetrance is low.<sup>6,7</sup> The syndrome has been proposed as LQT7; however, the ECG features are distinct from those of classic forms of long QT (LQT) syndrome, i.e., characteristic TU patterns, including enlarged U waves, a wide TU junction, and a prolonged terminal T-wave downslope.<sup>6,8</sup>

KCN/2 mutation, which encodes the Kir2.1 subunit, causes Andersen-Tawil syndrome. Kir2.1 is predominantly expressed in the brain, heart, and skeletal muscles and forms an inwardly rectifying potassium channel via the homo- or heteromeric assembly of 4 Kir2.x subunits.<sup>10</sup> Most KCNJ2 mutations cause loss of function or dominant-negative suppression of the inwardly rectifying

From the Department of Neurology (Y.K., M.F., M.N., T.K., H.M., M.P.T.), and Laboratory of Integrative Physiology, Department of Physiology (S.S., Y. Okamura), Osaka University Graduate School of Medicine, Suita, Osaka; Division of Neurogenetics (T.N., K.O.), Center for Neurological Diseases and Cancer, Nagoya University Graduate School of Medicine, Nagoya, Aichi; Department of Cardiovascular and Respiratory Medicine (H.K., H.I., M.H.), Shiga University of Medical Science, Otsu, Shiga; Division of Arrhythmia and Electrophysiology (T.A., W.S.), Department of Cardiovascular Medicine, National Cerebral and Cardiovascular Center, Suita, Osaka; Department of Pediatrics (M.Y.), National Hospital Organization Kagoshima Medical Center, Kagoshima; Department of Neurology (Y. Osaki, K.S.), Kurashiki Central Hospital, Kurashiki, Okayama; Department of Legal Medicine (T.S.), Osaka Medical College, Takatsuki, Osaka: Department of Cardiology (K.K.), Kurashiki Central Hospital, Kurashiki, Okayama; and Department of Cardiovascular Medicine (W.S.), Nippon Medical School, Bunkyo, Tokyo, Japan. Y.K. is currently affiliated with the Department of Neurology, Osaka General Medical Center, Sumiyoshi, Osaka, Japan; and T.K. is currently affiliated with the Department of Biochemistry and Molecular Biology, Division of Biological Sciences, The University of Chicago, IL. Go to Neurology,org for full disclosures. Funding information and disclosures deemed relevant by the authors, if any, are provided at the end of the article.

© 2014 American Academy of Neurology

1

<sup>\*</sup>These 3 authors contributed equally to this work.