## **Risk for Life-Threatening Cardiac Events in Patients With Genotype-Confirmed Long-QT Syndrome and Normal-Range Corrected QT Intervals**

Ilan Goldenberg, MD,\* Samuel Horr, MA,\* Arthur J. Moss, MD,\* Coeli M. Lopes, PHD,† Alon Barsheshet, MD,\* Scott McNitt, MS,\* Wojciech Zareba, MD, PHD,\* Mark L. Andrews, BBA,\* Jennifer L. Robinson, MS,\* Emanuela H. Locati, MD,§ Michael J. Ackerman, MD, PHD,¶ Jesaia Benhorin, MD,|| Elizabeth S. Kaufman, MD,# Carlo Napolitano, MD,\*\*†† Pyotr G. Platonov, MD, PHD,§§ Silvia G. Priori, MD, PHD,\*\*†† Ming Qi, MD,‡ Peter J. Schwartz, MD,‡‡ Wataru Shimizu, MD, PHD,|||| Jeffrey A. Towbin, MD,¶¶ G. Michael Vincent, MD,\*\*\* Arthur A. M. Wilde, MD, PHD,## Li Zhang, MD\*\*\*

Rochester and New York, New York; Milan and Pavia, Italy; Tel Aviv, Israel; Rochester, Minnesota; Cleveland, Ohio; Lund, Sweden; Suita, Japan; Houston, Texas; Amsterdam, the Netherlands; and Salt Lake City, Utah

Objectives	This study was designed to assess the clinical course and to identify risk factors for life-threatening events in patients with long-QT syndrome (LQTS) with normal corrected QT (QTc) intervals.
Background	Current data regarding the outcome of patients with concealed LQTS are limited.
Methods	Clinical and genetic risk factors for aborted cardiac arrest (ACA) or sudden cardiac death (SCD) from birth through age 40 years were examined in 3,386 genotyped subjects from 7 multinational LQTS registries, categorized as LQTS with normal-range QTc ( $\leq$ 440 ms [n = 469]), LQTS with prolonged QTc interval ( $>$ 440 ms [n = 1,392]), and unaffected family members (genotyped negative with $\leq$ 440 ms [n = 1,525]).
Results	The cumulative probability of ACA or SCD in patients with LQTS with normal-range QTc intervals (4%) was significantly lower than in those with prolonged QTc intervals (15%) ( $p < 0.001$ ) but higher than in unaffected family members (0.4%) ( $p < 0.001$ ). Risk factors ACA or SCD in patients with normal-range QTc intervals included mutation characteristics (transmembrane-missense vs. nontransmembrane or nonmissense mutations: hazard ratio: 6.32; $p = 0.006$ ) and the LQTS genotypes (LQTS type 1:LQTS type 2, hazard ratio: 9.88; $p = 0.03$ ; LQTS type 3:LQTS type 2, hazard ratio: 8.04; $p = 0.07$ ), whereas clinical factors, including sex and QTc duration, were associated with a significant increase in the risk for ACA or SCD only in patients with prolonged QTc intervals (female age >13 years, hazard ratio: 1.90; $p = 0.002$ ; QTc duration, 8% risk increase per 10-ms increment; $p = 0.002$ ).
Conclusions	Genotype-confirmed patients with concealed LQTS make up about 25% of the at-risk LQTS population. Genetic data, including information regarding mutation characteristics and the LQTS genotype, identify increased risk for ACA or SCD in this overall lower risk LQTS subgroup. (J Am Coll Cardiol 2011;57:51-9) © 2011 by the American College of Cardiology Foundation

Medicine, New York, New York; #‡Department of Cardiology, Fondazione Policlinico S. Matteo IRCCS and University of Pavia, Pavia, Italy; §\$Department of Cardiology, Lund University, Lund, Sweden; |||Division of Cardiology, Department of Internal Medicine National Cardiovascular Center, Suita, Japan; ¶¶Department of Pediatric Cardiology, Baylor College of Medicine, Houston, Texas; ##Department of Cardiology, Academic Medical Center, Amsterdam, the Netherlands; and the \*\*\*Department of Medicine, University of Utah School of Medicine, Salt Lake City, Utah. This work was supported by research grants HL-33843 and HL-51618 from the National Institutes of Health. The authors have reported that they have no relationships to disclose.

Manuscript received May 29, 2010; revised manuscript received July 8, 2010, accepted July 12, 2010.

From the \*Cardiology Division of the Department of Medicine, University of Rochester Medical Center, Rochester, New York; †Cardiovascular Research Institute University of Rochester Medical Center, Rochester, New York; ‡Department of Pathology, University of Rochester Medical Center, Rochester, New York; \$Cardiovascular Department De Gasperis, Niguarda Hospital, Milan, Italy; ||Heart Institute, Tel Aviv Sourasky Medical Center, Tel Aviv, Israel; ¶Departments of Medicine, Pediatrics, and Molecular Pharmacology and Experimental Therapeutics/Windland Smith Rice Sudden Death Genomics Laboratory, Mayo Clinic College of Medicine, Rochester, Minnesota; #The Heart and Vascular Research Center, MetroHealth Campus, Case Western Reserve University, Cleveland, Ohio; \*\*Molecular Cardiology, Fondazione S. Maugeri, University of Pavia, Pavia, Italy; †Leon Charney Division of Cardiology, New York University School of

Abbreviations and Acronyms
ACA = aborted cardiac arrest
<b>ECG</b> = electrocardiographic
LQTS = long-QT syndrome
LQT1 = long-QT syndrome type 1
LQT2 = long-QT syndrome type 2
LQT3 = long-QT syndrome type 3
<b>QTc</b> = corrected QT interval
<b>SCD</b> = sudden cardiac death

Congenital long-QT syndrome (LQTS) is an inherited channelopathy characterized by a prolonged corrected QT interval (QTc) at rest that is associated with an increased predisposition for polymorphic ventricular arrhythmias and sudden cardiac death (SCD) in young subjects without structural heart disease (1). To date, more than 500 mutations have been identified in 12 LQTS-susceptibility genes, with the long-QT syndrome type 1 (LQT1), long-QT syndrome type 2 (LQT2), and long-QT syndrome type 3 (LQT3) genotypes constituting more than

95% of genotype-positive LQTS and approximately 75% of all LQTS (2). Risk assessment in affected patients with LQTS relies primarily on a constellation of electrocardiographic (ECG) and clinical factors, including QTc interval and age-sex interactions (3–6). In addition, there is increasing evidence that genetic information and the molecular and cellular properties of the LQTS-causative mutation may identify subjects with increased risk for cardiac events (7–10). Despite these recent advances, however, currently there are limited data regarding the clinical course and risk factors for life-threatening events in patients with LQTS with normal resting QTc values, so-called silent mutation carriers, concealed LQTS, or normal–QT interval LQTS.

## See page 60

In the present study we used combined data from 7 national LQTS registries to: 1) compare the clinical courses of patients with LQTS and normal-range QTc intervals to those of patients with prolonged QTc intervals and of genotype-negative unaffected family members; and 2) identify specific clinical and genetic risk factors for life-threatening cardiac events in patients with LQTS with normal-range QTc intervals.

## **Methods**

**Study population.** The study population comprised 3,386 genotyped subjects drawn from the Rochester, New York, enrolling center (center 1) of the International LQTS Registry (n = 2,630), the Netherlands LQTS Registry (n = 391), and the Japanese LQTS Registry (n = 205), as well as from data submitted by other investigators specifically for this collaborative mutation analysis project from Denmark (n = 90), Italy (n = 28), Israel (n = 25), and Sweden (n = 17). Patients were derived from 552 proband-identified *KCNQ1* (LQT1), *KCNH2* (LQT2), and *SCN5A* (LQT3) families. The proband in each family had otherwise unex-

plained, diagnostic QTc prolongation or experienced LQTS-related symptoms. Patients were excluded from the study if they had: 1) >1 LQTS identified mutation (n = 70); 2) Jervell and Lange-Nielsen syndrome with deafness and 2 *KCNQ1* mutations or 1 known *KCNQ1* mutation and congenital deafness (n = 2); and 3) no identified mutation on genetic testing with prolonged QTc interval (>440 ms [n = 428]).

Data collection and end point. Routine clinical and rest ECG parameters were acquired at the time of enrollment in each of the registries. Measured parameters on the first recorded electrocardiogram included QT and R-R intervals in milliseconds, with QT interval corrected for heart rate using Bazett's (11) formula. Clinical data were collected on prospectively designed forms with information on demographic characteristics, personal and family medical histories, ECG findings, therapies, and events during long-term follow-up. Data common to all LQTS registries involving genetically tested subjects were electronically merged into a common database for the present study. In addition, information regarding QT interval-prolonging medications and triggers for cardiac events was collected through a specific questionnaire for patients enrolled the U.S. portion of the registry.

The primary end point of the study was the occurrence of a first life-threatening cardiac event, comprising aborted cardiac arrest (ACA; requiring external defibrillation as part of the resuscitation or internal defibrillation in patients with implantable cardioverter-defibrillators) or LQTS-related SCD (abrupt in onset without evident cause, if witnessed, or death that was not explained by any other cause if it occurred in a nonwitnessed setting such as sleep). In the multivariate models, follow-up was censored at age 41 years to avoid the influence of coronary disease on the occurrence of cardiac events. We also evaluated a secondary end point that included the occurrence of a first cardiac event of any type during follow-up (comprising syncope [defined as transient loss of consciousness that was abrupt in onset and offset], ACA, or SCD).

Phenotype characterization. For the purpose of this study, the QTc interval was categorized as normal range ( $\leq$ 440 ms) or prolonged (>440 ms) according to accepted criteria for the phenotypic definition of LQTS (12). Using this definition, the study population were categorized into 3 genotype and QTc subgroups: 1) LQTS with normal-range QTc interval (n = 469), comprising patients identified to have LQT1 to LQT3 mutations with QTc intervals  $\leq$ 440 ms; 2) LQTS with prolonged QTc interval (n = 1,392), comprising patients with LQT1 to LQT3 mutations with QTc intervals >440 ms; and 3) unaffected family members (n = 1,525), comprising registry subjects from genotypepositive proband-identified families who were genetically tested and found to be negative for the LQTS-associated mutation, with QTc intervals  $\leq$  440 ms (i.e., genetically and phenotypically unaffected family members).

Download English Version:

## https://daneshyari.com/en/article/2950640

Download Persian Version:

https://daneshyari.com/article/2950640

Daneshyari.com