SNP S1103Y in the cardiac sodium channel gene SCN5A is associated with cardiac arrhythmias and sudden death in a white family

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Cardiac arrhythmias cause 400 000 sudden deaths annually in the United States alone. Mutations in the cardiac sodium channel gene SCN5A on chromosome 3p21 cause cardiac arrhythmias and sudden death. In this study, we define an SCN5A mutation, S1103Y, in a white family associated with syncope, ventricular fibrillation, and sudden death. A very recent study reported the same mutation in 13.2% of African Americans, but not in the white population. Our study shows that mutation S1103Y does exist in the white population, and it is associated with a considerable risk of syncope, ventricular arrhythmia, ventricular fibrillation, and sudden death in this population.

SCN5A encodes the cardiac sodium channel, a member of the voltage gated sodium channel family. Although SCN5A is primarily expressed in heart, its expression has been detected in brain regions including neurons. The SCN5A protein contains 2016 amino acids and has a calculated molecular weight of 227 kDa. It consists of four homologous domains (DI-DIV) joined by short linking intracellular domains (DI-DIV). Each transmembrane domain contains six segments (S1-S6). Mutations in SCN5A cause syncpe, seizures, and sudden death triggered by lethal cardiac arrhythmias associated with long QT syndrome (LQTS), idiopathic ventricular fibrillation including Brugada syndrome, and cardiac conduction disease.

A single nucleotide polymorphism (SNP) in SCN5A, S1103Y, was identified in 13.2% of African Americans and was found to increase the risk of acquired arrhythmia in this population (note that this mutation occurred at codon 1103, but was designated S1102Y in their report). However, Splawski et al did not find S1103Y SNP in 511 white people. In this study, we report the cosegregation of SCN5A mutation S1103Y with arrhythmia and sudden death in a white family.

METHODS

Identification of LQTS patients and isolation of genomic DNA

Patients with LQTS and arrhythmias were identified at the Department of Cardiovascular Medicine at the Cleveland Clinic Foundation. Diagnosis of LQTS was as described previously. Informed consent was obtained from all participants or their guardians, in accordance with standards established by the Cleveland Clinic Foundation Institutional Review Board on Human Subjects. Genomic DNA was prepared from the patients’ whole blood with the DNA Isolation Kit for Mammalian Blood (Roche Diagnostic Co).

Mutation analysis

Mutation analysis was carried out using single strand conformational polymorphism (SSCP) analysis as described previously. The aberrant SSCP conformer was cut directly from dried gels, rehydrated in water, reamplified by PCR, and sequenced by an ABI-3100 Genetic Analyzer.

Ancestry analysis

Ancestry analysis (ANCESTRY 1.0) was performed for three family members in kindred QW1446 (II.1, II.3, and II.5 in fig 1A) by DNAPrint Genomics (Sarasota, FL). ANCESTRY 1.0 facilitates the estimation of biogeographical ancestry on three axes, Native American, West African, and European. In this analysis each subject was genotyped for 31 Ancestry Informative Markers. Using maximum likelihood, the individual ancestral proportions of these three populations in the resulting multilocus genotype was computed.

RESULTS AND DISCUSSION

Using SSCP analysis, we identified an anomalous SSCP conformer in a white family (kindred QW1446) (fig 1A). Kindred QW1446 is associated with a history of ventricular tachycardia (fig 1B), ventricular fibrillation, syncope, sudden death, and LQTS. To confirm that kindred QW1446 has a European descent, the biogeographical ancestry proportions of three of the sibs (II.1, II.3, II.5 in fig 1A) from the second generation were estimated. The maximum likelihood point estimates for each of these sibs were 100% European, 0% African, and 0% Native American. These results strongly suggest that kindred QW1446 is a white family.

DNA sequence analysis of the normal and the aberrant SSCP conformers showed that the abnormal SSCP conformer contained a single base substitution (C to A) at the second nucleotide of codon S1103Y of SCN5A (GenBank accession No M77235) (fig 1C). This substitution resulted in a non-conservative replacement of serine with tyrosine (S1103Y) in the intracellular linker between transmembrane domains II and III of the cardiac sodium channel (fig 1D). The S1103Y mutation was found in only the three affected subjects in kindred QW1446, but not in the normal family members. These genetic data suggest that S1103Y is a pathogenic mutation. Mutation S1103Y was not detected in 200 white controls. These data suggest that although mutation S1103Y exists in the white population, its prevalence rate is low in the normal white population.
Figure 1  Mutation S1103Y in SCN5A associated with arrhythmia and sudden death in kindred QW1446. (A) The pedigree of QW1446 is shown at the top. Normal subjects are represented by open circles (females) or squares (males), and the affected subjects are represented by solid symbols. Results of SSCP analysis are shown below the pedigree structure. The arrow indicates the abnormal SSCP band. The affected subjects carried the abnormal SSCP band but the normal family members did not. (B) Polymorphic ventricular tachycardia (arrows) recorded in II.6 by Holter monitoring. (C) Sequence analysis of normal (wild type) and abnormal (S1103Y) SSCP conformers. The mutation spot is indicated. (D) The substitution of C by A at the second position of codon 1103 of SCN5A led to substitution of a serine residue by a tyrosine residue (S1103Y). Mutation S1103Y is located in the intracellular loop between domains II and III of the cardiac sodium channel SCN5A.
Identification of SCN5A mutation S1103Y in the white population

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