Unconventional intronic splice site mutation in SCN5A associates with cardiac sodium channelopathy

T Rossenbacker, E Schollen, C Kuipéri, T J L de Ravel, K Devriendt, G Matthijs, D Collen, H Heidbüchel, P Carmeliet

Background: Mutations in the cardiac sodium channel, SCN5A, have been associated with one type of long-QT syndrome, with isolated cardiac conduction defects and Brugada syndrome. The sodium channelopathies exhibit marked variation in clinical phenotypes. The mechanisms underlying the phenotypical diversity, however, remain unknown. Exonic SCN5A mutations can be detected in 20% of Brugada syndrome patients.

Results: An intronic mutation (c.4810+3_4810+6dupGGGT) in the SCN5A gene, located outside the consensus splice site, was detected in this study in a family with a highly variable clinical phenotype of Brugada syndrome and/or conduction disease and in a patient with Brugada syndrome. The mutation was not found in a control panel of 100 (200 alleles) ethnically matched normal control subjects. We provide in vivo and in vitro evidence that the mutation can disrupt the splice donor site, activate a cryptic splice site, and create a novel splice site. Notably, our data show that normal transcripts can be also derived from the mutant allele.

Conclusions: This is the first report of an unconventional intronic splice site mutation in the SCN5A gene leading to cardiac sodium channelopathy. We speculate that its phenotypical diversity might be determined by the ratio of normal/abnormal transcripts derived from the mutant allele. Mutation analysis in the variable phenotypic expression of the disease.

Mutations in coding regions of the SCN5A gene can only be found in approximately 20% of Brugada syndrome patients. It is assumed that defects in as yet unidentified genes are responsible for the remainder of Brugada syndrome cases. In this study, an unconventional intronic splice error in the SCN5A gene was investigated as the cause of a sodium channelopathy with a diverse phenotype.

METHODS

Clinical investigation

The study is based on one western European family (proband III:1) and one western European patient (patient A). Proband III:1 and patient A underwent full clinical and electrophysiological investigation, including programmed electrical stimulation (PES) with up to three ventricular extra-stimuli (ES) at two drive cycle lengths. Family members of patient III:1 were evaluated with ECG and/or with SAECG and/or with procainamide challenge (10 mg/kg IV). A PR duration of ≥200 ms or a QRS duration of ≥115 ms was considered prolonged. An AH interval ≥120 ms (in sinus rate) or a Wenckebach cycle length ≥400 ms (that is, 150/min) (baseline state) was considered indicative for atrioventricular nodal conduction slowing. An HV interval ≥60 ms was considered as proof for His-Purkinje conduction slowing.

Mutation analysis

Genomic DNA was prepared from peripheral blood lymphocytes by standard methods. The entire coding region of the SCN5A gene was amplified by PCR and analysed by denaturing high performance liquid chromatography (DHPLC; Transgenomics, Crewe, UK) and subsequent sequencing using fluorescent dye chemistry (BigDye Terminator v3.1 Cycle Sequencing Kit, Applied Biosystems, Lennik, Belgium) and an ABI-3100 genetic analyser (Applied Biosystems). Absence or presence of the mutation and single nucleotide polymorphisms (SNPs) in family members was assessed by DHPLC.

cDNA analysis

Total RNA was isolated with Trizol (Invitrogen, Merelbeke, Belgium) from lymphocytes according to the manufacturer’s protocol. cDNA was prepared using 5 µg total RNA in a total volume of 19.5 µl with SuperscriptII RNase H-reverse transcriptase and oligo-dT priming (Invitrogen). The SCN5A transcript was amplified in two successive PCR reactions with nested PCR primers: the first amplicon was 1800 bp long

Abbreviations: AV, atrioventricular; DHPLC, denaturing high performance liquid chromatography; EPS, electrophysiological study; ES, extra-stimuli; ICCD, isolated cardiac conduction defect; PES, programmed electrical stimulation; SNPs, single nucleotide polymorphisms; WT, wildtype

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The reaction conditions were: 95 °C, 2 min; regular Taq polymerase and PCR buffer (Roche Diagnostics, 2 µl) performed in 50 µl with 1 µl cDNA in the first reaction and 2 µl of the first PCR product in the second reaction using regular Taq polymerase and PCR buffer (Roche Diagnostics, Indianapolis, IN). The reaction conditions were: 95°C, 2 min; (95°C, 20 s; 65°C 20 s (−1°C/cycle); 72°C, 1 min)×10; (95°C, 20 s; 55°C, 20 s; 72°C, 1 min)×4; 72°C, 3 min, 4°C, hold. The amplicons were analysed by direct sequencing using BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems) and analysis on ABI-3100 (Applied Biosystems).

Figure 1 Reverse transcriptase-PCR with vector primers SD6 and SA2 (see fig 3) on COS-7 cells transfected with WT and mutant pSPL3. Amplification products were characterised by direct sequencing. Lane 1: 100 bp DNA ladder (Eurogentec, Seraing, Belgium). Lanes 2 and 3: WT. Lane 4: 1 kb DNA ladder (Eurogentec). Lane 5: mutant. One WT band of 532 bp can be seen in lanes 2 and 3. Direct sequencing indicated correct splicing (see also fig 3). In the background, the sequence of transcript 3 can be detected (indicated by **). Note that PCR product was PCR purified and not prepared by gel extraction. Sequencing of the middle band detected transcript 3 (see also fig 3). Notably, the sequence of a WT transcript can be noticed in background (indicated by *). Sequencing of the lower band revealed transcript 2 (see also fig 3).

EXON TRAPPING EXPERIMENTS
Using DNA from proband III:1 as template, harbouring one wildtype (WT) and one mutant allele, a 933 (WT) and 937 (mutant) base pair (bp) SCN5A genomic fragment, containing exon 27 with 347 bp of 5' flanking sequences, was generated (forward primer F: 5'-TTG AGC CTT CCT CTT TAG ACC AA-3' and R: 5'-CCT TTT GGT GAA GGC AAA GA-3'). PCR was performed in 50 µl with 1 µl cDNA from exon 27; 72°C, 5 min; 4°C, hold. The family pedigree is shown in fig 2A and the clinical data of family members are given in table 1. After 2 years of follow up no appropriate shocks were delivered. The family pedigree is shown in fig 2A and the clinical data of family members are given in table 1. As can be appreciated, clinical phenotypes are highly variable, ranging from a benign presentation with a subtle conduction defect to a more aggressive variant of the disease.

RESULTS
Association of splice site mutation with cardiac sodium channelopathy
Female proband III:1 presented at the age of 15 with syncope evoked by atrial flutter with 1:1 conduction. Baseline ECG recording showed Brugada syndrome phenotype, first degree atrioventricular (AV) block, and intraventricular conduction disturbances (fig 2C). An electrophysiological study (EPS) noted diffuse conduction disturbances at AV nodal and ventricular level. The patient underwent successful catheter ablation of the flutter without later recurrence. PES induced polymorphic ventricular tachycardia with one and two ES and ventricular fibrillation with three ES. An implantable cardioverter defibrillator was inserted. After 2 years of follow up no appropriate shocks were delivered. The family pedigree is shown in fig 2A and the clinical data of family members are given in table 1. As can be appreciated, clinical phenotypes are highly variable, ranging from a benign presentation with a subtle conduction defect to a more aggressive variant of the disease.
During EPS, microsatellite marker data and haplotype analysis based on detection of SNPs in the SCN5A gene were indicative of a common ancestor for proband III:1 and patient A. The following SNPs were analysed: IVS2-24C/T, IVS16-6C/T, c.3183G/A, and IVS19+72T/C. The haplotype shared between proband III:1 and patient A was IVS2-24T (2%)/IVS16-6T (4%)/c.3183A (14%)/IVS19+72C. The frequencies of the specified allele in a control population of 100 healthy subjects is given between brackets. Considering the low frequency of these SNPs, the fact that each of these individuals carries the same haplotype strongly suggests a common ancestor origin. Additionally, both patients shared the same length for microsatellite markers D3S1298 (197 bp) and D3S3593 (214 bp).

Mutation detection analysis of the SCN5A gene in the proband and in patient A only revealed a duplication of 4 bases in the splice donor of intron 27 (c.4810+3_4810+6dupGGGT) besides several additional intronic SNPs, more distantly removed from the splice site. The aberration co-segregated in affected family members (fig 2B) and was not found in a control panel of 100 (200 alleles) ethnically matched normal control subjects.

Abnormal splicing by the c.4810+3_4810+6dupGGGT allele

According to several splice prediction programs (data not shown), the splice site of intron 27 is a weak splice donor. The duplication of GGGT in a repeat of two copies of GGGT...
spanning the splice site, leaves the first six bases of the donor splice site unaffected. Because of the special nature of the duplication, we decided to test the pathogenic effect of this aberration on the correct splicing of exon 27.

Although the SCN5A gene is supposed to be uniquely expressed in heart, we were able to amplify SCN5A transcripts in lymphocytes. While in control cells only the WT transcript was amplified, a WT and an aberrant transcript (corresponding to transcript 2, fig 3) were detected in lymphocytes from proband III:1 and patient A. Unfortunately, the level of these transcripts in lymphocytes was too low to permit reliable relative quantification of each transcript.

To exclude aberrant splicing due to illegitimate transcription, we tested the effect of the mutation in an in vitro exon trapping experiment. Vectors harbouring the WT or the mutant genomic fragment, containing exon 27 and at least 300 bp flanking intronic sequences, were transfected in COS-7 cells. As indicated in figs 1 and 3, splicing in COS-7 cells, transfected with WT vector, consistently generated WT transcript, whereas splicing in cells transfected with the mutant vector produced three alternatively spliced forms (transcripts 1, 2, and 3; fig 3). In transcript 1, the 59 splice donor site is not recognised and no splicing occurs at this site, resulting in incorporation of intronic sequences in this transcript. An additional number of 95 amino acids will be translated from the intronic sequence before a stop codon terminates translation. These additional residues are completely unrelated to the native sequence of the sodium channel. Transcript 2 is spliced from a cryptic splice donor site within exon 27, resulting in a deletion of the 39-most 96 bases of exon 27. This causes an in-frame deletion of amino acids 1572–1604. Considering the extent of the deletion which covers fragments of segments 2 and 3 of domain IV, it is likely that this will have an impact on protein folding and function. In transcript 3, a novel splice donor site is used at the intronic GT, 39 of exon 27, resulting in the incorporation of four additional nucleotides. This causes a frame-shift G1604fs183 (that is, glycine as last unaffected amino acid of four additional nucleotides. This causes a frame-shift mutation report

### Table 1 Clinical data

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*At time of investigation; †1:1 antegrade conduction via fast pathway till cycle length of >1000 and 520 ms, respectively; ±left anterior hemiblock.

**DISCUSSION**

In this report, an unconventional splice site mutation in SCN5A is described as the cause of a sodium channelopathy. Ten base pairs (–4 to +6) around the 59 exon-intron boundary are essential for normal splicing and constitute the donor splice site. Sequence alterations at these locations are potentially disease causing. Previous splice site mutations, associated with ICCD or Brugada syndrome, have been predicted to cause exon skipping by disruption of the splice site.4–11–14 Exon skipping causes either a frameshift and a premature stop codon or a deletion of the region encoded by the exon. As evidenced by an in vitro approach, the mutation described here can simultaneously disrupt the splice donor site (mutant transcript 1), can activate a cryptic splice site (mutant transcript 2), and can create a novel donor site (mutant transcript 3). To study the consequences of the splice mutation on the generation of SCN5A transcripts in a human cell line, we used lymphocytes as heart tissue of affected patients was unavailable. Mutant transcript 2 could be isolated in proband III:1 and in patient A but not in control patients. We are well aware of the problem of illegitimate transcription, which could lead to aberrant splice products of any gene in any cell type. However: (i) the same splice error was seen in both patients; (ii) the splice error was not detected in control patients; (iii) data were reproducible in subsequent experiments; and (iv) the splicing abnormality, detected in the patient’s lymphocytes, was also found in the exon trapping experiments. The fact that mutant transcript 1 and 3 were found in transfected COS-7 cells but not in lymphocytes may be explained by (i) differences in the spliceosome between human lymphocytes and COS-7 cells and (ii) by the absence of the native acceptor splice site of intron 27 in the experiments with COS-7 cells.

The duplication leaves the consensus donor splice site unaffected and, thus, correct “WT-like” splicing of the mutant allele could, theoretically, still occur. This question can, however, not be easily addressed by analysing lymphocytes from affected heterozygous individuals, as they already carry a single WT allele. However, exon trapping experiments with the isolated mutant allele revealed that a WT transcript was indeed generated from the mutant allele. It can thus be speculated that the ratio of normal/abnormal transcripts and,

**Normal splicing by the c.4810+3_4810+6dupGGGT allele**

As the duplication leaves the original consensus donor splice site unaffected, we also assessed whether correct “WT-like” splicing of the mutant allele still occurred. Since the WT transcript is only four nucleotides shorter than mutant transcript 3, it cannot be easily separated by conventional agarose gel electrophoresis. A low signal of mutant transcript 3 in the sequence analysis (fig 1) suggested, however, the presence of WT transcript in cells transfected with the mutant construct. We therefore subcloned the PCR amplification products with an approximate size of 530–540 bp (fig 1) and sequenced 59 clones. Out of those 59 clones, two clones contained the WT transcript (data not shown), indicating that the mutant allele is indeed capable of encoding a WT channel, though at a reduced level.

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consequently, the ratio of functional/non-functional channels, might influence the phenotypic expression of the disease in affected family members. At least theoretically, it is conceivable that the presence of some WT channels in affected individuals might result in a milder phenotype, whereas the complete absence of WT channels might be expected to yield the most severe clinical phenotype.

In conclusion, the c.4810+3_4810+6dupGGGT SCN5A is a novel intronic mutation that leads to aberrant splicing. Lymphocytes could be used to assess the functional effect of the mutation and constitute an interesting approach for the study of other SCN5A splice site mutations. The mutation is located outside the consensus splice site. Our findings point to the role of intronic nucleotides other than at

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**Figure 3** (A) WT plasmid generates WT transcripts. (B) Mutant plasmid generates three aberrant transcripts (1–3). WT (A) and mutant (B) exon 27 and part of flanking introns (347 and 315 bp) were subcloned in exon trapping vector pSPL3 and transfected in COS-7 cells. Transcripts are derived from the pairing of vector and cloned splice signals. In transcript 1, the 5′ splice site is not recognised and no splicing occurs at this site. In transcript 2, a cryptic splice site in exon 27 becomes activated which causes a deletion of 96 bases in exon 27. In transcript 3, a novel splice site is used, that is, second intronic GT which leads to incorporation of four extra bases in the transcript. Mutation is indicated in italic. SD6 and SA2 are vector primers used for PCR (see fig 1).
invariant donor sites. Whether a given ratio of normal/abnormal transcripts might contribute to the variation in phenotypic characteristics in different family members, awaits further confirmation by investigation of heart tissue of affected patients.

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Competing interests: none declared

Ethics approval: The study was performed in accordance with recommendations of the Ethics Committee of the University Hospital Leuven.

Written informed consent to participate in the study was obtained from all subjects or their legal representatives.

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