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Maturation of hiPSC-derived cardiomyocytes promotes adult alternative splicing of SCN5A and reveals changes in sodium current associated with cardiac arrhythmia

Giulia Campostrini (1) 1, Georgios Kosmidis 1, Dorien Ward-van Oostwaard (1) 1, Richard Paul Davis (1) 1, Loukia Yiangou (1) 1, Daniele Ottaviani (1) 2,3, Christiaan Cornelis Veerman 4, Hailiang Mei (1) 5, Valeria Viktorovna Orlova (1) 1, Arthur Arnold Maria Wilde (1) 4, Connie Rose Bezzina (1) 4, Arie Otto Verkerk (1) 4,6, Christine Lindsay Mummery (1) 1,7, and Milena Bellin (1) 1,2,3*

¹Department of Anatomy and Embryology, Leiden University Medical Center (LUMC), 2333 ZA Leiden, The Netherlands; ²Department of Biology, University of Padua, 35121 Padua, Italy; ³Veneto Institute of Molecular Medicine, 35129 Padua, Italy; ⁴Department of Clinical and Experimental Cardiology, Heart Centre, Amsterdam University Medical Centre, location AMC, University of Amsterdam, Meibergdreef 9, 1105 AZ, Amsterdam, The Netherlands; ⁵Sequencing Analysis Support Core, Leiden University Medical Center, 2333 Leiden, The Netherlands; ⁶Department of Medical Biology, Amsterdam Cardiovascular Sciences, Amsterdam UMC, University of Amsterdam, Meibergdreef 9, 1105 AZ, Amsterdam, The Netherlands; and ⁷Department of Applied Stem Cell Technologies, University of Twente, 7500 AE, Enschede, The Netherlands

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Aims

Human-induced pluripotent stem cell-cardiomyocytes (hiPSC-CMs) are widely used to study arrhythmia-associated mutations in ion channels. Among these, the cardiac sodium channel *SCN5A* undergoes foetal-to-adult isoform switching around birth. Conventional hiPSC-CM cultures, which are phenotypically foetal, have thus far been unable to capture mutations in adult gene isoforms. Here, we investigated whether tri-cellular cross-talk in a three-dimensional (3D) cardiac microtissue (MT) promoted post-natal *SCN5A* maturation in hiPSC-CMs.

Methods and results

We derived patient hiPSC-CMs carrying compound mutations in the adult *SCN5A* exon 6B and exon 4. Electrophysiological properties of patient hiPSC-CMs in monolayer were not altered by the exon 6B mutation compared with isogenic controls since it is not expressed; further, CRISPR/Cas9-mediated excision of the foetal exon 6A did not promote adult *SCN5A* expression. However, when hiPSC-CMs were matured in 3D cardiac MTs, *SCN5A* underwent isoform switch and the functional consequences of the mutation located in exon 6B were revealed. Up-regulation of the splicing factor muscleblind-like protein 1 (*MBNL1*) drove *SCN5A* post-natal maturation in microtissues since its overexpression in hiPSC-CMs was sufficient to promote exon 6B inclusion, whilst knocking-out *MBNL1* failed to foster isoform switch.

Conclusions

Our study shows that (i) the tri-cellular cardiac microtissues promote post-natal SCN5A isoform switch in hiPSC-CMs, (ii) adult splicing of SCN5A is driven by MBNL1 in these tissues, and (iii) this model can be used for examining post-natal cardiac arrhythmias due to mutations in the exon 6B.

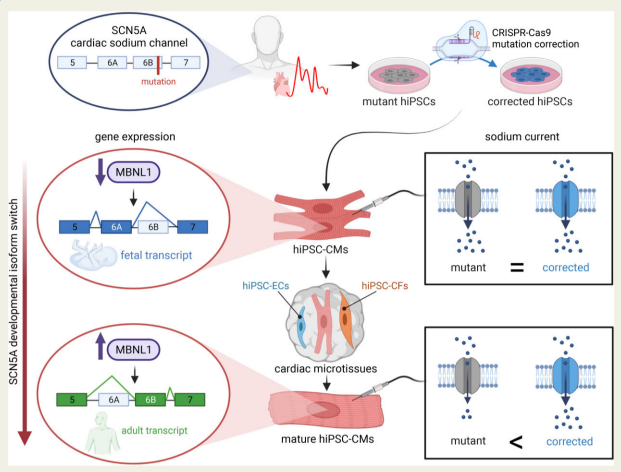
Translational perspective

The cardiac sodium channel is essential for conducting the electrical impulse in the heart. Postnatal alternative splicing regulation causes mutual exclusive inclusion of fetal or adult exons of the corresponding gene, SCN5A. Typically, immature hiPSCCMs fall short in studying the effect of mutations located in the adult exon. We describe here that an innovative tri-cellular three-dimensional cardiac microtissue culture promotes hiPSC-CMs maturation through upregulation of MBNL1, thus revealing the effect of a pathogenic genetic variant located in the SCN5A adult exon. These results help advancing the use of hiPSC-CMs in studying adult heart disease and for developing personalized medicine applications.

^{*} Corresponding author. E-mail: m.bellin@lumc.nl or milena.bellin@unipd.it

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Graphical Abstract



hiPSC-CM maturation in 3D cardiac microtissues promotes SCN5A fetal to adult isoform switch.

Keywords

human-induced pluripotent stem cell-derived cardiomyocytes • cardiac sodium channel • SCN5A • cardiac arrhythmias • cardiac microtissue

1. Introduction

Human-induced pluripotent stem cells (hiPSCs) are widely used to derive cardiomyocytes (hiPSC-CMs) and study cardiac diseases at the cellular and molecular level. 1-3 Patient-specific hiPSC-CMs capture the genetic landscape of the affected individual, allowing precise dissection of pathogenic mechanisms and testing drug responses that may ultimately lead to (personalized) treatments. Among the cardiac diseases that have been modelled using hiPSC-CMs are inherited cardiac arrhythmias caused by mutations in cardiac ion channels. ^{1,4} The SCN5A gene encodes the pore-forming α -subunit of the cardiac sodium (Na⁺) voltage-gated channel Nav1.5. This channel conducts the depolarizing Na⁺ current (I_{Na}) , which is responsible for the fast upstroke of the action potential (AP) in working cardiomyocytes; it is essential for conducting the electrical impulse in the heart. Mutations in SCN5A are associated with different cardiac diseases, including Brugada syndrome, long QT syndrome, isolated conduction defects, and dilated cardiomyopathy.⁵ Nav1.5 expression is developmentally regulated, with foetal - or adult isoforms generated by alternative splicing of the *SCN5A* exon 6 (6A foetal and 6B adult), which encodes the voltage sensor of the first channel subunit.⁶ The mechanism of *SCN5A* alternative splicing regulation and other cardiac genes during development has been studied in animal models, where the muscleblind-like RNA-binding protein 1 (Mbnl1) has been identified as key for inducing a post-natal switch.^{7–9} In humans, MBNL1 is sequestered in the heart of myotonic dystrophy patients causing re-expression of the foetal *SCN5A* exon 6A¹⁰ and the importance of MBNL proteins for late myogenic maturation has been recently demonstrated using hiPSCs.¹¹

hiPSC-CMs can recapitulate many electrical alterations due to mutations in *SCN5A* or in genes of Nav1.5 beta-subunits ^{12–22} and these are in some cases notably different from those observed in commonly used heterologous expression systems, ²³ most likely because hiPSC-CMs carry the full complement of cardiac ion channel and accessory protein genes. Moreover, gene editing in hiPSCs allows effects of specific mutations to be distinguished from potential confounding elements resulting from line-to-line variability; patient mutations can be introduced into

wild-type (WT) hiPSCs or mutations in patient hiPSC lines can be corrected. 15,24 However, cardiomyocytes differentiated from hiPSCs have an immature (foetal-like) phenotype with their electrical properties, structure, and gene expression profiles resembling first or second gestational trimester foetal cardiac cells^{25,26} rather than adult cardiomyocytes. Although I_{Na} in hiPSC-CMs display half-maximal potential $(V_{1/2})$ of activation and inactivation consistent with values reported for native human ventricular myocytes, 27 the maximal AP upstroke velocity (V_{max}) is slower than in adult cardiomyocytes. ²⁸ In addition, hiPSC-CMs mostly express the Nav1.5 foetal isoform, masking effects due to mutations located in the adult exon 6B.²⁹ Previously, we and others addressed the issue of SCN5A in vitro maturation by culturing hiPSC-CMs for long periods to promote spontaneous maturation²⁹ and by using a medium inducing a metabolic switch from glycolysis to fatty acid oxidation.²⁰ We recently discovered that hiPSC-CM maturation is overall enhanced in a three-dimensional (3D) microtissue (MT) environment containing cardiac fibroblasts (CFs) and endothelial cells (ECs).³⁰ hiPSC-CMs isolated from MTs have more hyperpolarized RMP and faster AP upstroke velocity, suggesting an electrical maturation involving Nav1.5.

Here, we showed that maturation of hiPSC-CMs in MTs promoted expression of the *SCN5A* adult isoform and allowed (i) evaluation of functional effects of the p.R225W mutation, located in adult exon 6B and (ii) detection of the more severe disease phenotype associated with the compound heterozygosity for p.W156X mutation, ¹³ as observed in patients. ³¹ Using gene-corrected isogenic hiPSCs was crucial in identifying mutation-specific effects. We also found that *MBNL1* upregulation in MTs was necessary and sufficient to induce the expression of *SCN5A* exon 6B.

2. Methods

2.1 Patients' clinical history and phenotype

The clinical characteristics of the family were previously reported. ³¹ The proband patient (*Figure 1A*, II-3), born with a severe tachycardia and conduction disorder, was treated with beta-blockers and later fitted with an ICD implant. Despite the ICD, he died from intractable ventricular arrhythmias at age 21. The parents and a sister (*Figure 1A*, II-1) of the proband were asymptomatic, whereas the other sister (*Figure 1A*, II-2) carrying the same mutations of the proband was affected by intractable arrhythmia and died at age of 1.

2.2 hiPSCs generation and culture

A skin punch biopsy was obtained following written informed consent from the patient at the age of 18 and approval by the medical ethics committee of the Amsterdam Medical Center (AMC), in accordance with the Declaration of Helsinki. Dermal fibroblasts were reprogrammed using the Sendai virus by the Leiden University Medical Center (LUMC) hiPSC core facility, as described previously, ¹³ following the protocol approved by the LUMC and AMC ethical committees. Details for hiPSC culture and differentiation are reported in the Supplementary material online.

2.3 hiPSC editing using CRISPR/Cas9

For the correction of c.468G>A mutation in SCN5A exon 4 (W156X) and excision of SCN5A exon 6A specific sgRNA sequences (*Figures 1E* and *3A*, Supplementary material online, *Table S1*) were cloned into the pSpCas9(BB)-2A-Puro (PX459) v2 plasmid encoding Cas9 nuclease and puromycin resistance (Addgene #62988).³² For the correction of

c.673C > T mutation in exon 6B (p.R225W) and the excision of exon 4 of MBNL1, the Alt-R® CRISPR–Cas9 System (IDT Technologies) was used. Specific single-stranded DNA oligonucleotide (ssODN) repair template carrying the WT base and silent mutations were used for each mutation correction (see Supplementary material online, Table S1). Additional information is provided in the Supplementary material online.

2.4 Formation of 3D cardiac MTs

MTs were formed by combining defined ratios of hiPSC-CMs, -ECs, and -CFs (70:15:15), as previously described. 30,33

2.5 Electrophysiology

Beating monolayers of hiPSC-CMs were dissociated at day 21 of differentiation using Tryple Select $0.5\times$ (ThermoFisher Scientific) and plated onto Matrigel-coated glass coverslips at low density. Electrophysiological properties of hiPSC-CMs were analysed 10 days after dissociation. MTs were dissociated at day 21 of culture using collagenase II (Worthington Industries), as previously described. Cells were plated on Matrigel-coated coverslips in LI-BPEL supplemented with 50 ng/mL VEGF and 5 ng/mL bFGF, refreshed with LI-BPEL medium the next day, and analysed for electrophysiological properties 7–10 days after dissociation.

Patch-clamp experiments were performed using an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA, USA). Clampex 10.7 was used for data acquisition, voltage control, and analysis. APs were recorded in isolated single cells using the perforated patch-clamp technique in current-clamp mode and sodium current (I_{Na}) was recorded with the ruptured patch-clamp technique in voltage-clamp mode. Patch-clamp solutions and protocols are reported in the Supplementary material online.

2.6 RNA isolation, cDNA synthesis, and gene expression analysis

Total RNA was isolated either from hiPSC-CMs on day 21 of differentiation and from hiPSCs using Nucleospin kit (Machery and Nagel), or from MTs on day 21 of culture using RNeasy Micro Kit (Qiagen), according to the manufacturers' instructions. cDNA was synthesized using iScriptTM cDNA Synthesis Kit and gene expression was analysed by qPCR with IQ Syber Green Supermix (Bio-Rad), according to the manufacturer's protocol. All qPCR reactions were performed in duplicate. Gene expression levels were normalized to the *RPL37A* housekeeping gene.

For quantification of mRNA-splicing isoforms containing 6A or 6B exons, BioRad QX200TM Droplet Digital PCR (ddPCR) system was used following the manufacturer's instructions. Primer and probe sequences used for qPCR and ddPCR are reported in Supplementary material online, *Table S2 and S3*.

2.7 RNA-seq analysis

The RNA was extracted with the RNeasy micro kit (QIAGEN) as from the manufacturer's instructions, followed by paired-end sequencing at 150 bp, 40 million reads for each sample (Novogene, Cambridge, UK). Residual Illumina adapters were trimmed from the RNA sequences using cutadapt³⁴ with parameter -m 20. Reads were aligned to the human genome (build 38, Ensembl version 96) using STAR.³⁵ BAM files were used by rMATS³⁶ to detect differentially spliced exons and sashimi plots were made using the function rmats2sashimiplot (https://github.com/Xinglab/rmats2sashimiplot).

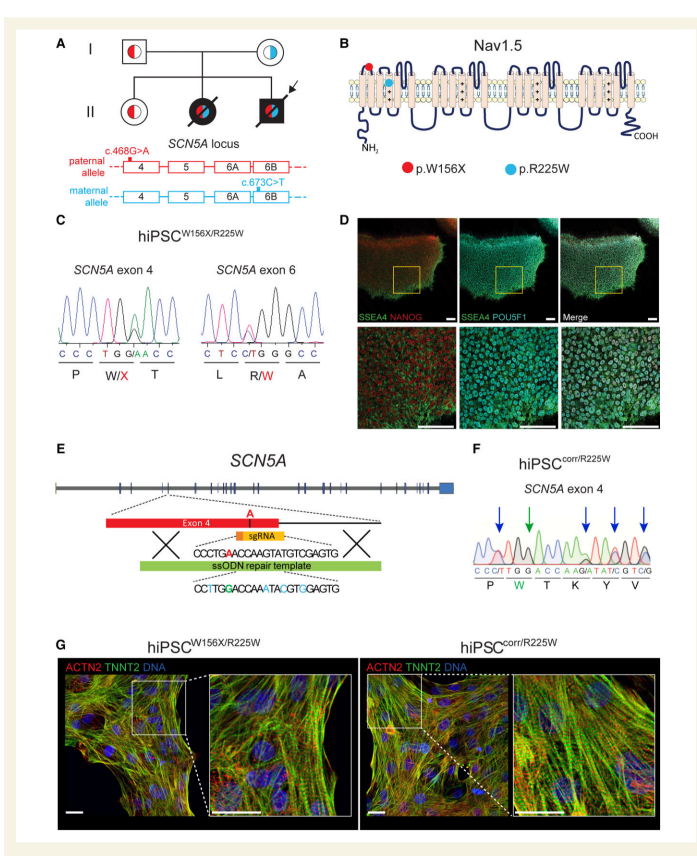


Figure 1 Generation of hiPSCs from a patient with compound heterozygous mutations in *SCN5A* and their genetic correction. (*A*) Top, family tree of the proband (arrow) showing in black filling the patients affected by cardiac conduction defects and in white unaffected individuals. Red (left) and light blue (right) colours within the family tree symbols indicate the genotype at the *SCN5A* locus, as indicated at the bottom of the figure. Red (top), paternal allele carrying the c.468G>A *SCN5A* (p.W156X) mutation in exon 4; light blue (bottom), maternal allele carrying the c.673C>T *SCN5A* (p.R225W) mutation in exon 6B. (*B*) Schematic representation of the sodium channel Nav1.5 α-subunit, encoded by *SCN5A*. The red (left) and the light blue (right) circles show the position of p.W156X and p.R225W mutations, respectively. (*C*) Sanger sequencing chromatograms showing the c.468G>A *SCN5A* (p.W156X)

Exon counts for the SCN5A and MBNL1 canonical transcripts, ENST00000413689 and ENST00000282486, respectively, were retrieved using featureCounts³⁷ with parameters -t exon and -g exon_id. Transcripts per kilobase million (TPM) was calculated considering all exons counts as library size and by using the R script convertCounts. The coverage of constitutive and alternative exons for SCN5A or MBNL1 was calculated as percentage of TPM assigned to each exon bin over the TPM mean of the whole transcript.

RNA-seq data are available in the GEO repository with accession number GSE180290. Publicly available foetal heart (FH) and adult heart (AH) RNA-seq data (GEO, accession number GSE62913³⁸) were used for Supplementary material online, *Figure S7A*.

2.8 In vitro generation of MBNL1 mRNA and transfection

MBNL1 mRNA was generated by in vitro transcription using INCOGNITO T7 5mC- and Ψ -RNA Transcription Kit, ScriptCap Cap 1 Capping System and A-Plus Poly(A) Polymerase Tailing Kit (all from Cellscript, LLC) following the manufacturer's instructions.

hiPSC-CMs were transfected with MBNL1 mRNA using Lipofectamine Stem Transfection Reagent (Invitrogen) according to the manufacturer's instructions. Details are reported in the Supplementary material online.

2.9 Statistical analysis

Data were obtained from at least three independent differentiation- or MT formation experiments and were expressed as mean \pm standard error of the mean (SEM). Data were compared using t-test or one-way ANOVA followed by Fisher LSD post hoc test. Statistical analysis was performed using OriginPro 2016 (Origin Lab), RStudio, or GraphPad Prism (GraphPad Software, Inc.).

3. Results

3.1 Generation and genetic correction of hiPSCs from a patient with compound heterozygous mutations in SCN5A

A patient (*Figure 1A*, Il-3) presented with severe cardiac conduction abnormalities and ventricular arrhythmias early after birth and was genetically characterized as carrying bi-allelic mutations in the *SCN5A* gene. These were a c.468G>A nucleotide change in exon 4 of *SCN5A* (NM_198056.2) inherited from the father, leading to the stop codon mutation p.W156X (NP_932173.1), and a c.673C>T nucleotide change in exon 6B of *SCN5A* inherited from the mother. This led to the substitution of a positively charged arginine with an aromatic tryptophan

(p.R225W) in the voltage-sensor segment of the first domain of the channel (*Figure 1A* and *B*). When the patient turned 18, we reprogrammed dermal skin fibroblasts using the Sendai virus to generate the hiPSC^{W156X/R225W} line. Sanger sequencing confirmed the presence of both *SCN5A* mutations (*Figure 1C*). After several passages, hiPSCs were negative for the Sendai virus (see Supplementary material online, Figure *S1A*), had a normal karyotype (see Supplementary material online, *Figure S1B*), expressed the pluripotency markers NANOG, SSEA4, POU5F1 (*Figure 1D*), and genome-wide gene expression analysis indicated high 'pluripotency score' and low 'novelty score' with the PluriTest algorithm³⁹ (see Supplementary material online, *Figure S1C*, left).

While the p.W156X Nav1.5 mutation was previously characterized in a heterologous system³¹ and using hiPSC-CMs derived from the proband's father (Figure 1A, I-1), 13 the phenotypic effects of the p.R255W Nav1.5 mutation was only studied in heterologous system³¹ since hiPSCs from the proband's mother were not available. To examine the effects of this mutation specifically, we corrected the c.468G>A SCN5A mutation in the hiPSCW156X/R225W line using CRISPR-Cas9 and generated the hiPSC^{corr/R225W} line (Figure 1E and F). One hiPSC clone was selected, in which Sanger sequencing confirmed the correction of the c.468G>A SCN5A mutation and the presence of the silent mutations in heterozygosity (Figure 1F). No modifications were present in the in silico-predicted two intragenic Cas9 off-target sites by Sanger sequencing (data not shown). hiPSC^{corr/R225W} had a normal karyotype (see Supplementary material online, Figure S1D) and high pluripotency score, as confirmed by PluriTest (see Supplementary material online, Figure S1C, right). The original hiPSCW156X/R225W line was heterozygous for the rs6781731 SNP located in the intron downstream exon 4. The corrected hiPSC^{corr/R225W} was also heterozygous for this SNP, confirming that the recombination involved only the paternal allele as intended (see Supplementary material online, Figure S1E).

3.2 No functional effects of exon 6B mutation in monolayer hiPSC-CM cultures expressing the foetal but not adult SCN5A isoform

We differentiated hiPSC^{W156X/R225W} and hiPSC^{corr/R225W} lines into cardiomyocytes (CMs) in monolayer (*Figure 1G*). Differentiation efficiencies were comparable between the two lines and consistently yielded >90% cardiac troponin T (TNNT2)-positive cells (see Supplementary material online, Figure S2A). The cardiac ion channel genes *SCN5A*, *NCX1*, *KCND2*, *KCNH2*, *KCNQ1*, and *KCNJ12* were expressed at similar levels in CMs from hiPSC^{W156X/R225W} and hiPSC^{corr/R225W} (see Supplementary material online, Figure S2B).

Figure 1 Continued

mutation in exon 4 (left), and the c.673C>T SCN5A (p.R225W) mutation in exon 6B (right), both present in heterozygosis in the patient-derived hiPSC^{W156X/R225W} line. (*D*) Representative immunofluorescence images of hiPSC^{W156X/R225W} undifferentiated colonies showing expression of pluripotency markers NANOG (red), SSEA4 (green), and POU5F1 (cyan). Bottom panels are an enlargement of the framed area in top panels. Scale bars: 25 μm. (*E*) Schematic showing the strategy used to correct the c.468G>A (p.W156X) mutation in SCN5A exon 4 with CRISPR/Cas9 in the paternal allele of hiPSC^{W156X/R225W}. The mutant adenine base is shown in red; in yellow the sgRNA guiding the Cas9 to the mutation; in green the ssODN used as donor template for homology-directed DNA repair. Underneath part of the ssODN sequence, showing in green the WT guanine base and in light blue the silent mutations. (*F*) Sanger sequencing chromatogram showing SCN5A exon 4 after correction of the c.468G>A (W156X) mutation in hiPSC^{corr/R225W}. The green arrow (second arrow from the left) indicates the corrected patient mutation and the blue arrows indicate the silent mutations inserted in one allele. (*G*) Representative immunofluorescence staining for ACTN2 (red) and TNNI3 (green) in hiPSC^{W156X/R225W}- and hiPSC^{corr/R225W}-CMs. Nuclei are stained with Dapi (blue). Panels on the right are enlargement of the framed area in the left panels. Scale bars: 10 μm.

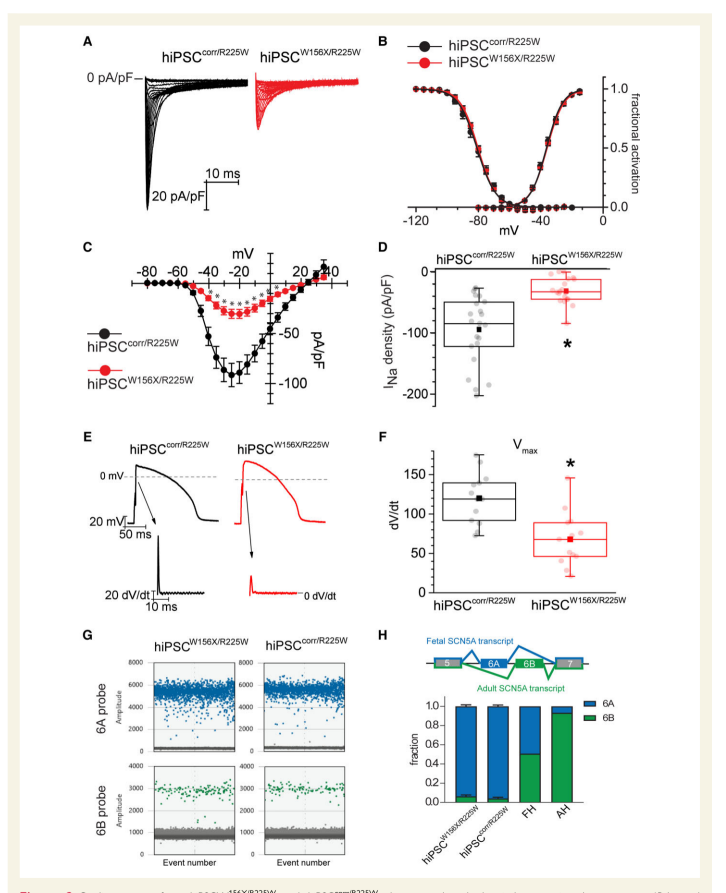


Figure 2 Cardiomyocytes from hiPSCW $^{156X/R225W}$ and hiPSC $^{corr/R225W}$ show no altered electrical properties due to exon 6B-located p.R225W SCN5A mutation and express mainly the foetal SCN5A isoform. (A) Representative I_{Na} traces recorded as indicated in hiPSC $^{corr/R225W}$ - (black) and hiPSC $^{W156X/R225W}$ - (red) CMs during voltage-clamp activation protocol (test range -80/-15 mV, 22 steps, holding

We next examined the electrophysiological properties of CMs from hiPSCW156X/R225W and hiPSCcorr/R225W lines. Figure 2A shows representative I_{Na} traces from hiPSC^{W156X/R225W}- and hiPSC^{corr/R225W}-CMs. While the voltage dependence of I_{Na} activation and inactivation was similar between the two lines (see Supplementary material online, Figure 2B), I_{Na} density was lower in hiPSC^{W156X/R225W}-CMs over the whole physiological voltage range (Figure 2C), with the mean peak current density in hiPSCW156X/R225W-CMs ($-31 \pm 4.5 \text{ pA/pF}$) significantly reduced compared with hiPSC^{corr/R225W}-CMs ($-94.2 \pm 11.4 \text{ pA/pF}$) (Figure 2D). Notably, the current density recorded in hiPSC^{corr/} $^{
m R225W}$ -CMs was comparable to that previously reported (~ -100 pA/ pF) for CMs differentiated with the same method from a WT hiPSC line (hiPSCWT/WT), whereas I_{Na} density in hiPSCW156X/R225W-CMs was similar to that recorded in hiPSC-CMs carrying only the p.W156X mutation (~ -30 pA/pF) derived from the proband's father (Figure 1A I-1).¹³

Altered I_{Na} impacts the electrical activity of cardiomyocytes, particularly the AP upstroke velocity. We, therefore, recorded APs in hiPSC^{W156X/R225W} - and hiPSC^{corr/R225W}-CMs paced at 1 Hz. To compensate for the typical depolarized RMP of foetal-like hiPSC-CMs, we used dynamic clamp to set the RMP to -86 mV. Figure 2E shows representative AP traces recorded from hiPSCW156X/R225W- and hiPSCcorr/ R225W-CMs, and their respective first derivative trace. In agreement with the differences in I_{Na} density, the maximal first derivative of AP upstroke, corresponding to the maximal upstroke velocity (V_{max}) , was significantly lower in hiPSCW156X/R225W- compared with hiPSCcorr/R225W-CMs (Figure 2F). As observed for I_{Na} , V_{max} values from hiPSC $^{corr/}$ R225W-CMs were similar to those previously reported for WT hiPSC-CMs using the same dynamic clamp technique. 41 Other AP parameters, namely RMP, AP amplitude (APA), and AP duration (APD), were not different between the two lines (see Supplementary material online, Figure S2C).

To check whether both mutations contributed to $I_{\rm Na}$ and $V_{\rm max}$, we measured exon 6A and 6B expression in CMs from hiPSC $^{\rm W156X/R225W}$ and hiPSC $^{\rm corr/R225W}$ lines. Using digital droplet PCR (ddPCR) we quantified exon 6A- and exon 6B-containing transcripts (see Supplementary material online, Figure S2D and E). Figure 2G illustrates representative ddPCR results for hiPSC $^{\rm W156X/R225W}$ - and hiPSC $^{\rm corr/R225W}$ -CMs. Exon 6B-containing SCN5A transcripts accounted for approximately 5% and were much lower than exon 6A-containing transcripts (Figure 2G and H). In FH and AH tissues, exon 6B-containing transcripts accounted for 50% and 94% of all transcripts, respectively,

in line with previous reports, ^{29,42} and thus were expressed at higher levels than in hiPSC-CMs (*Figure 2H*).

These results indicated that hiPSC $^{W156X/R225W}$ and hiPSC $^{corr/R225W}$ -CMs expressed the foetal SCN5A isoform almost exclusively and I_{Na} density and V_{max} reduction in the compound mutant was due to the p.W156X Nav1.5 mutation. The effect of p.R225W mutation could not be properly determined.

3.3 Genetic excision of exon 6A does not increase adult SCN5A isoform expression

To investigate whether the removal of exon 6A might force SCN5A exon 6B expression and thus reveal the effect of the p.R225W Nav1.5 mutation, we used CRISPR-Cas9 to excise this exon from both SCN5A alleles in the hiPSC^{W156X/R225W} line. As illustrated in Figure 3A, we designed 2 gRNAs targeted to the flanking regions of exon 6A and identified a clone in which exon 6A had been excised in both alleles. The actual excision spanned a larger intronic region than intended by the strategy design (Figure 3A and Supplementary material online, S3A). In the hiPSC lines with complete exon 6A excision (referred to as 6A-KO hiPSC W156X/ R225W), we corrected the p.W156X mutation in exon 4 (generating the 6A-KO hiPSC^{corr/R225W} line), using CRISPR-Cas9 with the same strategy as earlier (Figure 1E). The excised hiPSC lines differentiated into CMs with similar efficiencies and beating properties as the parental hiPSC line (data not shown). As expected, exon 6A was not expressed at all in excised CMs at 21 or 60 days of culture, whereas expression of exon 6B was detected (see Supplementary material online, Figure S3B). To examine SCN5A transcripts expressed by CMs from the excised hiPSC lines, we amplified by PCR a region between exon 3 and exon 8 of SCN5A (Figure 3B). Here, we only used the 6A-KO hiPSCcorr/ R225W line, since in the 6A-KO hiPSCW156X/R225W line, the stop codon p.W156X mutation in exon 4 leads to nonsense-mediated mRNA decay¹³ and could confound the results. Gel electrophoresis of PCR products revealed three distinct bands (Figure 3C). Sanger sequencing demonstrated these corresponded to three different transcript species: the highest band represented the adult isoform of SCN5A transcript including exon 6B; the middle band, a transcript in which exon 6B was skipped, and in the lowest band, both exons 5 and 6B were skipped (Figure 3C).

To investigate whether the overall expression of the adult *SCN5A* isoform was increased compared with the non-excised line, we performed ddPCR analysis, normalizing the expression of exon 6B with that of the TATA-binding protein (TBP). Exon 6B expression was similar in the

Figure 2 Continued

potential = -100 mV). (*B*) Average activation (AC) and inactivation (IC) curve of I_{Na} recorded in hiPSC^{corr/R225W}- (black; AC: $V_{1/2} = -36.3 \pm 0.7$ mV, n = 21; IC: $V_{1/2} = -80.9 \pm 1.2$ mV, n = 15) and hiPSC^{W156X/R225W}- (red; AC: $V_{1/2} = -35.9 \pm 0.4$ mV, n = 17; IC: $V_{1/2} = -80.2 \pm 0.7$ mV, n = 14) CMs. P > 0.05 with Student's t-test. (*C*) Mean current–voltage (*I*–*V*) relationships of I_{Na} recorded in hiPSC^{corr/R225W} (black, n = 23) and hiPSC^{W156X/R225W}. (red, n = 21) CMs, showing a reduction in current density in hiPSC^{W156X/R225W}. Experiments >4.*P < 0.05 with two-way ANOVA repeated measures. Data in (*B*) and (*C*) are shown as mean \pm SEM. (D) Box plot of maximal peak I_{Na} density in hiPSC^{corr/R225W} (black) and hiPSC^{W156X/R225W} (red) CMs. Dots: single values, lines: median, square: mean, error bars: $1.5 \times$ inter-quartile range (IQR). *P < 0.05 with Student's t-test. (*E*) Representative APs recorded from hiPSC^{Corr/R225W}- (black) and hiPSC^{W156X/R225W} which corresponds to a slower V_{max} . (*F*) Box plot of mean V_{max} of APs recorded in hiPSC^{corr/R225W}- (black) and hiPSC^{W156X/R225W}- (red) CMs. Dots, lines, squares and error bars as in (*D*). n = 13, experiments = 4.*P < 0.05 with Student's t-test. (*G*) Representative outcome of the ddPCR assay showing exon 6A (blue) and 6B (green) expression in hiPSC^{W156X/R225W}- and hiPSC^{corr/R225W}- CMs. Each coloured dot represents a positive droplet for the fluorophore, grey dots represent negative droplets. (*H*) Top, schematic of the developmentally regulated alternative splicing of *SCN5A* exon 6: the foetal *SCN5A* transcript includes exon 6A, whereas the adult *SCN5A* transcript includes exon 6B. Bottom, bar graph showing the average fraction of *SCN5A* exon 6: the foetal *SCN5A* transcript includes exon 6A, whereas the adult *SCN5A* transcript includes exon 6B. Bottom, bar graph showing the average fraction of *SCN5A* exon 6: the foetal *SCN5A* transcript includes exon 6A, whereas t

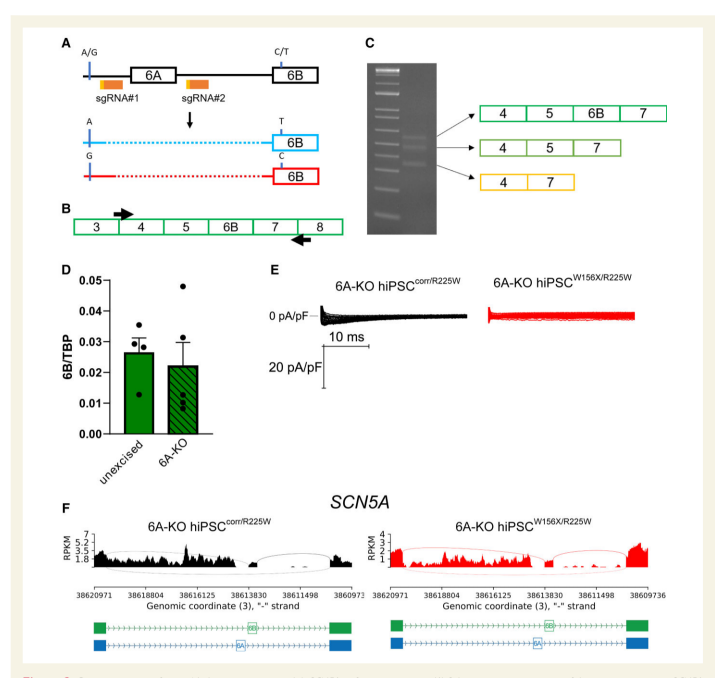


Figure 3 Genetic excision of exon 6A does not increase adult SCN5A isoform expression. (A) Schematic representation of the strategy to excise SCN5A exon 6A using CRISPR/Cas9 with two sgRNAs (sgRNA#1 and #2, orange). In red and light blue, the results of excision (dotted line) in the hiPSCs^{W156X/R225W}: the maternal allele (top, blue, with the mutated T in exon 6B and the A SNP in the intron between exon 5 and exon 6A) carried a 15 bp larger deletion upstream of exon 6A compared with the paternal allele (bottom, red, WT C in exon 6B and G SNP in 5-6A intron). (B) Schematic showing the primers (black arrows) used to amplify and sequencing cDNA from 6A-KO hiPSC^{corr/R225W}-CMs. (C) Gel electrophoresis of PCR products from (B) showing three bands corresponding to three transcript species as indicated by the arrows. (D) Bar graph of mean expression of exon 6B in unexcised and 6A-KO hiPSC-CMs. Data were normalized to TBP. n = 4. Dots: single values. (E) Representative I_{Na} traces recorded from 6A-KO hiPSC^{corr/R225W} and 6A-KO hiPSC^{W156X/R225W} CMs showing an almost complete absence of the current. (F) Sashimi plots of RNA-seq data from 6A-KO hiPSC^{corr/R225W} and 6A-KO hiPSC^{W156X/R225W} CMs showing the absence of exon 6A transcription but extensive transcription of the intronic region between exon 5 and exon 6B.

excised hiPSC-CMs and in the parental non-excised hiPSC-CMs (Figure 3D). Since exon 6B-containing transcripts were likely the only functional transcript species produced in the exon 6A-excised lines, we investigated whether this would unmask the functional phenotype of the p.R225W Nav1.5 with electrophysiology. We attempted to record $I_{\rm Na}$ in 6A-KO hiPSC hiPSC and 6A-KO hiPSC Corr/R225W CMs but $I_{\rm Na}$ was virtually absent in all (n=9) 6A-KO hiPSC hiPSC CMs

(Figure 3E), while 50% of the 6A-KO hiPSC^{corr/R225W} cells (n=10) showed very small $I_{\rm Na}$ (Figure 3E) with mean peak current density of -12.9 ± 5.5 pA/pF. Analysis of SCN5A transcript expression by RNA-seq showed that upon excision of exon 6A, not only was there spurious mRNA produced but also the intronic region between exon 5 and exon 6B was aberrantly included at a similar level to the exons (Figure 3F).

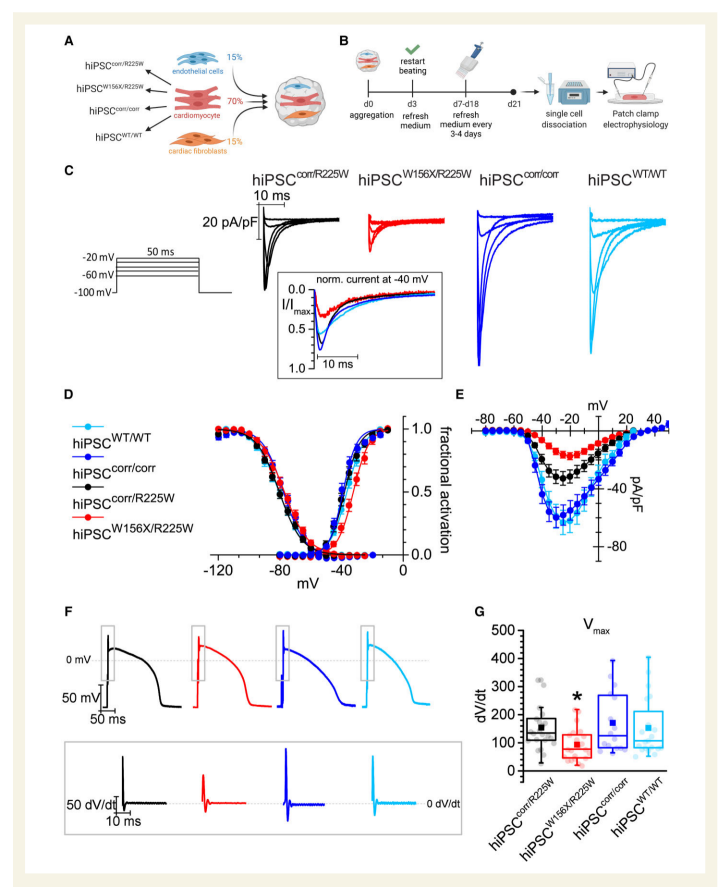


Figure 4 Electrical maturation of hiPSC-CMs in 3D cardiac microtissues with hiPSC-derived non-myocytes reveals functional effects of p.R225W SCN5A mutation. (A) Schematic of the microtissue formation using hiPSC-CMs from four lines (hiPSC^{corr/R225W}, hiPSC^{W156X/R225W}, hiPSC^{corr/corr},

Taken together, these results demonstrated that excision of SCN5A exon 6A impairs the splicing mechanism, leading to abnormally spliced transcripts of unknown significance and the inclusion of intronic regions in the transcript population. Importantly, 6B-containing transcript levels were not sufficient to generate detectable $I_{\rm Na}$ for comparison of the two mutant lines.

3.4 Electrical maturation of hiPSC-CMs in 3D cardiac MTs with hiPSC-derived cardiac non-myocytes reveals functional effects of p.R225W mutant Nav1.5

We next tested whether hiPSC-CM maturation in cardiac MTs containing hiPSC-derived CFs and cardiac ECs, 30,33 could promote SCN5A exon 6B expression and thus reveal the effect of the p.R225W mutation. The cardiac MTs were generated using defined ratios of the three cell types as shown in Figure 4A and dissociated into single cells after 21 days of culture to analyse I_{Na} by patch clamp (Figure 4B). Since I_{Na} had not been characterized in hiPSC-CMs derived in this way before, we used hiPSC-CMs from hiPSC^{WT/WT} as a process control. ^{13,43} In addition, we corrected the p.R225W mutation in hiPSC^{corr/R225W}, generating the isogenic control hiPSC^{corr/corr} (see Supplementary material online, Figure \$4A). Correction of the mutation in hiPSC^{corr/corr} was confirmed by Sanger sequencing (see Supplementary material online, Figure S4B). The line had a normal karyotype and differentiated efficiently into cardiomyocytes (see Supplementary material online, Figure S4C and D). Four groups of MTs were analysed, containing CMs from hiPSC W156X/R225W. hiPSC^{corr/R225W}, hiPSC^{corr/corr} (Supplementary material online, videos S1-S3), and hiPSCWT/WT. Representative traces of normalized I_{Na} recorded in CMs dissociated from MTs are shown in Figure 4C. The inset shows respective current trace at -40 mV normalized on peak current, showing smaller activation for hiPSCW156X/R225W compared with the other lines, indicating that a smaller fraction of I_{Na} channels opened at the given voltage. Indeed, the activation curve of $hiPSC^{W156X/R225W}$ showed a significant rightward shift compared with the other lines (Figure 4D). I_{Na} current density showed instead a genotype-specific reduction, with I_{Na} significantly lower along the physiological voltage range in hiPSCW156X/R225W-CMs compared with all the other lines, and significantly lower in hiPSC $^{\rm corr/R225W}\text{-}$ compared with hiPSC $^{\rm corr/}$ corr and hiPSCWT/WT-CMs (Figure 4E).

Finally, we examined the effect of the altered I_{Na} on the AP. Figure 4F shows representative AP traces and the derivative of the AP upstroke

recorded in hiPSC-CMs dissociated from MTs and paced at 1 Hz. $V_{\rm max}$ was significantly smaller in hiPSC^{W156X/R225W}-CMs compared with CMs from the other lines (*Figure 4G*), showing that the compound mutations have an additive effect on $I_{\rm Na}$ density, thus impacting on CM electrical activity. Other AP parameters such as spontaneous resting membrane potential, APA and APD were instead similar in the different lines (see Supplementary material online, Figure S5).

3.5 Alternative adult SCN5A splicing is promoted in hiPSC-CMs exposed to the MT microenvironment

To obtain a more general overview of gene expression, we performed bulk RNA sequencing in hiPSCW156X/R225W- and hiPSCcorr/R225W-2D CMs and 3D MTs after 21 days of culture. Gene ontology (GO) analysis showed that genes related to GO terms of heart development and contraction had a similar expression in the two hiPSC lines, both in 2D CMs and in MTs (see Supplementary material online, Figure S6A), indicating that the cells had reached similar levels of differentiation and maturation. Our single-cell (sc) RNA-seq datasets from MTs and hiPSC-CMs previously published from the same WT line used here confirmed that SCN5A is only expressed in the hiPSC-CM population (Supplementary material online, Figure S6B).³⁰ Analysis by ddPCR showed that MTs from hiPSC^{W156X/R225W}, hiPSC^{corr/R225W}, and hiPSC^{WT/WT} expressed similar fractions of SCN5A exon 6B (Figure 5A), which accounted for around 27% of total SCN5A transcripts (Figure 5B). Notably, the fraction of exon 6B-containing transcripts was 5-6 times higher in MTs from hiPSC^{W156X/R225W} and hiPSC^{corr/R225W} compared with their respective hiPSC-CMs derived in monolayer (Figure 5C). RNA-seq analysis confirmed that exon 6B expression was increased in MTs compared with monolayer hiPSC-CMs in both lines (Figure 5D). Overall SCN5A expression was also higher in MTs compared with monolayer hiPSC-CMs in both lines (Figure 5E), in agreement with scRNA-seq in hiPSC WT/WT (see Supplementary material online, Figure S6C) and confirmed by qPCR (see Supplementary material online, Figure S6D, left). The total expression of exon 6 (6A+6B) was similar to upstream and downstream exons; however, the fraction of exon 6B in MTs was higher than in monolayer hiPSC-CMs in both lines (Figure 5E). Exon 6B fraction was nonetheless lower in MTs compared with FH and AH (see Supplementary material online, Figure S7A). Interestingly, SCN5A expression was not increased in MTs containing 6A-KO hiPSC-CMs and the expression of exons downstream of exon 6B was drastically

Figure 4 Continued

hiPSC^{WT/WT}) and ECs and CFs from hiPSC^{WT/WT}. The percentage of each cell type is indicated. (*B*) Schematic of 21-day MT culture protocol until single-cell dissociation for analysis. (*C*) Representative I_{Na} traces recorded in hiPSC^{corr/R225W}-, hiPSC^{W156X/R225W}-, hiPSC^{corr/corr}-, and hiPSC^{WT/WT}-CMs dissociated from MTs, corresponding to the voltage steps reported on the right. In the inset, the current at -40 mV normalized to the peak current, to compare the fraction of open channels around the $V_{1/2}$ of activation. (*D*) Activation (AC) and inactivation (IC) curve of I_{Na} recorded in hiPSC^{corr/R225W}- (black; AC: $V_{1/2} = -38.1 \pm 1.2$ mV; IC: $V_{1/2} = -80.6 \pm 1.1$ mV; n = 16), hiPSC^{W156X/R225W}- (red; AC: $V_{1/2} = -32.4 \pm 1.2$ mV*, n = 11; IC: $V_{1/2} = -77.3 \pm 1.4$ mV, n = 9), hiPSC^{corr/corr}- (blue; AC: $V_{1/2} = -39.3 \pm 0.9$ mV, n = 17; IC: $V_{1/2} = -77.9 \pm 1.2$ mV; n = 16), and hiPSC^{WT/WT}- (light blue; AC: $V_{1/2} = -36.5 \pm 0.9$ mV, v = 16), and hiPSC^{W156X/R225W}-CMs. v = 160.5 vs. the other lines with one-way ANOVA and Fisher's post hoc test. Experiments v = 3. (E) Mean voltage-density plot of v = 160.5 m hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- (light blue) CMs, showing a reduction in current density in hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- and hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (red), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (red), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (red), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (red), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (black), hiPSC^{W156X/R225W}- (red), hiP

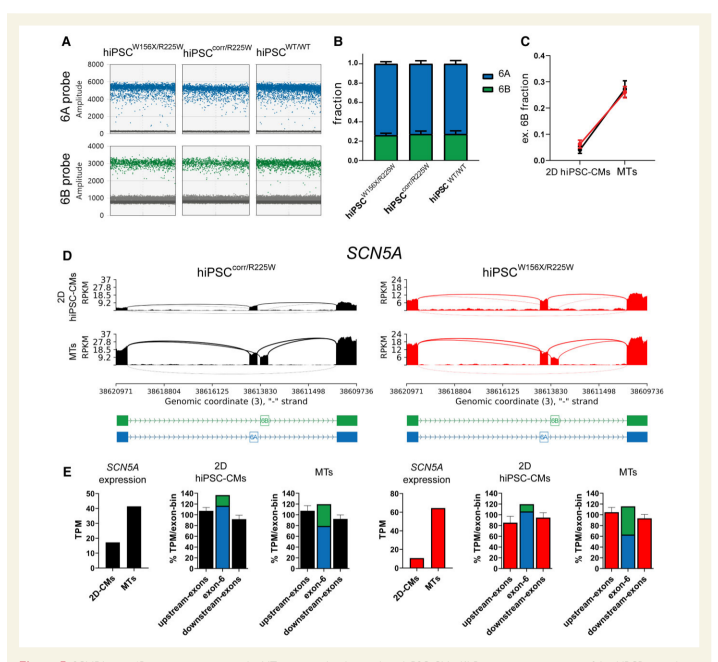


Figure 5 SCN5A exon 6B expression is increased in MTs compared with monolayer hiPSC-CMs. (A) Representative outcome of the ddPCR assay showing exon 6A (blue) and 6B (green) expression in hiPSC^{W156X/R225W}-, hiPSC^{corr/R225W}-, and hiPSC^{WT/WT}-CMs. (B) Bar graph of the average fraction of SCN5A exon 6A and 6B expression in hiPSC^{W156X/R225W}-, hiPSC^{corr/R225W}- and hiPSC^{WT/WT}-MTs (n=3). (C) Plot showing the fraction of SCN5A exon 6B expression in monolayer (2D) hiPSC-CMs compared with MTs for hiPSC^{corr/R225W} (black) and hiPSC^{W156X/R225W} (red), indicating a strong increase of the relative expression of exon 6B in MTs. (D) Sashimi plots of RNA-seq data from 2D hiPSC-CMs and MTs for hiPSC^{corr/R225W} and hiPSC^{W156X/R225W} showing increased expression of SCN5A exon 6B in MTs in both lines. (E) Bar graphs from RNA-seq data (TPM) from hiPSC^{corr/R225W} (left, black) and hiPSC^{W156X/R225W} (right, red) showing increased expression of SCN5A and higher fraction of exon 6B-including transcripts in MTs compared with 2D hiPSC-CMs.

reduced, confirming that 6A excision impaired splicing in the entire gene (see Supplementary material online, Figure S7B).

3.6 The alternative splicing regulator MBNL1 promotes *SCN5A* exon 6B inclusion during hiPSC-CM maturation

The RNA-binding protein Mbnl1 has been previously shown to promote exon 6B- and inhibit exon 6A expression in the mouse heart.^{7,9} We,

therefore, investigated whether MBNL1 is involved in hiPSC-CM alternative splicing.

RNA-seq analysis showed that *MBNL1* was up-regulated in MTs assembled using hiPSC^{W156X/R225W}- and hiPSC^{corr/R225W}-CMs compared with 2D hiPSC-CMs cultures (*Figure 6A and B*). Although we found MBNL2 was also up-regulated in MTs (see Supplementary material online, Figure *S6D*), scRNA-seq from our dataset previously published showed that *MBNL1* but not *MBNL2* was up-regulated in the CM population of MTs compared with monolayer hiPSC-CMs (see

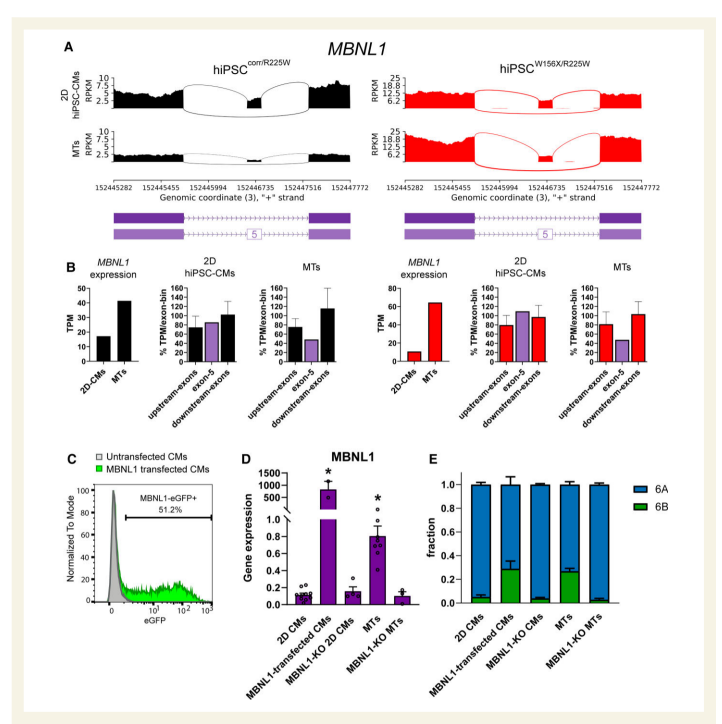


Figure 6 *MBNL1* is up-regulated in MTs and promotes exon 6B expression. (*A*) Sashimi plots of RNA-seq data from 2D hiPSC-CMs and MTs for hiPSC^{corr/R225W} and hiPSC^{W156X/R225W} showing decreased expression of *MBNL1* exon 6 in MTs in both lines. (*B*) Bar graphs showing expression based on RNA-seq data (TPM) from 2D CMs and MTs from hiPSC^{corr/R225W} (left, black) and hiPSC^{W156X/R225W} (right, red); *MBNL1* was up-regulated and a lower fraction of exon 5-including transcripts (purple) in MTs compared with 2D hiPSC-CMs. (*C*) FACS analysis showing the percentage of eGFP-expressing cells (MBNL1-eGFP+) in untransfected (grey) and *MBNL1*-transfected (green) hiPSC-CMs. (*D*) *MBNL1* expression analysis by qPCR in 2D CMs, *MBNL1*-transfected CMs, MBNL1-KO 2D CMs, MTs, and MBNL1-KO MTs. *P < 0.05, One-way ANOVA compared with 2D CMs. n > 3. Dots: single values. (*E*) Fraction of exon 6A (blue) and exon 6B (green) *SCN5A* analysed by ddPCR in the cells from (*C*), as indicated. n > 3.

Supplementary material online, Figure S6C), suggesting a major role of MBNL1 specifically in CMs. In addition, MTs displayed skipping of MBNL1 exon 5, which is promoted by MBNL1 itself in adult CMs⁴⁴ (Figure 6B) and thus suggesting that MBNL1 is functional in MTs.

To test whether MBNL1 is necessary and sufficient to promote exon 6B inclusion, we overexpressed or knocked out MBNL1 in hiPSC-CMs.

First, we transiently transfected hiPSC-CMs with an mRNA construct encoding *MBNL1* and eGFP. Transfection efficiency reached around 50%, as shown by FACS analysis of eGFP+ cells (*Figure 6C*) and overexpression of *MBNL1* in transfected hiPSC-CMs compared with untransfected cells was confirmed by qPCR (*Figure 6D*). ddPCR showed that the exon 6B fraction was increased in MBNL1-transfected hiPSC-CMs

to levels comparable with those in MTs (*Figure 6E*). We then knocked out *MBNL1* in hiPSCs (*MBNL1*-KO hiPSCs) (see Supplementary material online, Figure S8). Cardiac *in vitro* differentiation was not affected in *MBNL1*-KO hiPSCs, but *MBNL1* expression in 2D hiPSC-CMs was negligible and did not differ between unexcised and MBNL1-KO lines (*Figure 6D*). When included in MTs, *MBNL1*-KO hiPSC-CMs did not up-regulate *MBNL1*, differently from the unexcised line (*Figure 6D*). Moreover, exon 6B fraction in MBNL1-KO MTs was comparable to the 2D cultures by ddPCR analysis, and significantly lower than in MTs composed by unexcised hiPSC-CMs (*Figure 6E*).

Taken together these results demonstrated that 3D cardiac MT constructs composed of hiPSC-cardiac ECs and -CFs promoted alternative splicing maturation in hiPSC-CMs such that they displayed increased expression of SCN5A exon 6B and its regulator MBNL1. This was sufficient to reveal the functional defects caused by the p.R225W mutation on $I_{\rm Na}$ and to dissect the different contributions of p.W156X and p.R225W Nav1.5 mutations to the disease in the patient.

4. Discussion

The mammalian heart undergoes extensive functional modification after birth, regulated by changes in gene expression at both transcriptional and post-transcriptional level.⁴⁵ Alternative splicing is an important mechanism in the post-natal transition and applies to many cardiac genes, 8,46 including the cardiac sodium channel Nav1.5.6,29 The switch between the foetal SCN5A isoform containing exon 6A and the adult SCN5A isoform containing exon 6B is completed postnatally in mice and humans. 42,47 Immature hiPSC-CMs predominantly express the foetal SCN5A transcript. Although this does not preclude evaluating the functional effects of SCN5A mutations using hiPSC-CMs in general, it does represent a hurdle for those mutations located in exon 6B which is expressed only postnatally. Here, we studied a patient who developed severe cardiac conduction defects early after birth and carried compound mutations in Nav1.5 (p.W156X and p.R225W), one of which (p.R225W) located in the adult exon 6B. Clinical characterization of the family indicated that the cardiac disorder was present only in individuals with compound heterozygosity for the two mutations, with individuals carrying only one of the two Nav1.5 mutations being asymptomatic.³¹ This suggests an additive effect of the two mutations. To dissect the contribution of each mutation, we generated a hiPSC line from the patient and an isogenic corrected line carrying only the mutation in SCN5A exon 6B (hiPSC^{corr/R225VV}). However, when we recorded I_{Na} in hiPSC^{corr/R225W}-CMs, current density was similar to that measured in WT hiPSC-CMs, whereas the compound mutant showed a reduction of I_{Na} density similar to that previously observed in hiPSC-CMs with only p.W156X mutation.¹³ The lack of effect of p.R225W contrasts with the strong reduction in current density shown in Xenopus laevis oocytes overexpressing the same p.R225W Nav1.5 mutation³¹; this could be partly due to differences between heterologous expression systems and human cells. Interestingly, the single mutation effect in patients was negligible, suggesting that there might be differences between in vitro systems and the patient phenotype. Analysis of SCN5A exon expression revealed that only a small fraction of exon 6B-containing transcripts was expressed in hiPSC-CMs, in line with previous reports in independent hiPSC lines.²⁹ This explained the functional results and confirmed that the immature hiPSC-CM phenotype precludes analysis of mutations located in the adult SCN5A isoform.

Remarkably, while genetic excision of Scn5a exon 6B resulted in selective re-expression of foetal Nav1.5 in mice, 48 our complementary approach of SCN5A exon 6A excision in hiPSCs did not lead to selective expression of the adult Nav1.5. Indeed, 6A-KO hiPSC-CMs expressed similar (low) amounts of SCN5A exon 6B compared with the original non-excised lines and presented negligible I_{Na} . This demonstrates that the exon 6B expression levels in standard monolayer hiPSC-CM cultures are not sufficient to generate I_{Na} and that I_{Na} measured in non-excised hiPSC-CMs is mainly conducted by the foetal channel. The extremely low (or negligible) I_{Na} in 6A-KO hiPSC-CMs is likely a consequence of very few functional channels formed, because 6A excision interferes with the splicing machinery, as shown by the inclusion of intronic regions and exclusion of exon 5. This indicates that some regulatory sequences important for correct splicing might be present either within the intronic region removed by the excision, or in exon 6B itself, driving its own exclusion during splicing. Further studies with consecutive removal of different sequences within the 6A region are needed to elucidate the exact mechanisms underlying the regulation of SCN5A splicing. However, the switch between exon 6A and exon 6B is likely to need not only dedicated regulatory sequences, but also specific factors which may be lacking in immature CMs.

We and others developed different systems to improve the maturation of hiPSC-CMs using more physiological environments involving 3D engineering and/or co-culture with other (cardiac) cell types. 49,50 We previously showed that hiPSC-CMs in our cardiac tri-cell-type MTs undergo maturation at the functional, structural, metabolic, and gene expression levels³⁰; however, we did not examine whether it involved alternative splicing transition. Both ddPCR and RNA-seg revealed that in cardiac MTs there is a significant increase in the fraction of SCN5A exon 6B- vs. exon 6A-containing transcripts. A similar switch between SCN5A exon 6A and exon 6B was also previously promoted by culturing hiPSC-CMs in monolayer for extended periods.²⁹ Of note, cardiac MT maturation is less time-consuming than prolonged monolayer culture (40 days vs. more than 60), and the number and ratio of cells in MTs remain unchanged during the 21 days of culture required for maturation.³⁰ In our experience, long-term culture of hiPSC-CMs was accompanied by some cell loss, the proliferation of non-cardiomyocyte cells and the risk of detachment of the beating monolayer from the dish. Importantly, cardiac MTs showed higher expression of the exon 6B fraction (five- to sixfold, MT vs. 20 days 2D hiPSC-CMs) compared with long-term monolayer hiPSC-CM culture (two- to three-fold, 60 days vs. 20 days).²⁹ Nevertheless, levels of exon 6B expression comparable to adult human heart were not reached, suggesting that splicing regulation may still differ between in vitro and in vivo conditions. It remains to be investigated whether the effects of exon 6B mutation are revealed in other 3D or monolayer maturation systems or media.

The possibility to derive all cardiac MT cell components from hiPSCs allowed us to include either mutant, corrected isogenic, or WT hiPSC-CMs while the other cell types derived from the same (WT) stock, avoiding possible confounding factors. Since hiPSC-CM maturation achieved in MTs is maintained after dissociation, ³⁰ we could analyse the electrical properties of single CMs. This revealed the contribution of p.R225W Nav1.5 mutation to the disease phenotype, uncovering the gene-dosage relationship and linking the presence of one or two mutated SCN5A alleles with a mild (\sim 45%) or severe (\sim 70%) $I_{\rm Na}$ reduction, respectively. Reduced $I_{\rm Na}$ density but no differences in the channel kinetics of activation or inactivation were measured in the p.R225W hiPSC-CMs compared with the corrected or WT hiPSC-CMs, much as has been observed for another mutation (p.I230T) in the same exon. ²⁹

A positive shift in the voltage dependence of $I_{\rm Na}$ activation was instead measured in the double mutant hiPSC-CMs, similar to the effects of the homozygous p.I230T Nav1.5 mutation.²⁹ This shift can be attributed to the p.R225W mutation carried on the maternal—and only functional—allele, since the p.W156X mutation likely generates a non-functional protein, due to the stop codon in the paternal allele. In hiPSC^{corr/R225W}, this change was instead masked by the corrected (and functional) allele. A shift in the voltage dependence of p.R225W Nav1.5 was also observed in transfected *X. laevis* oocytes³¹; however, this was more pronounced and involved both activation and inactivation, underscoring the fact that ion channel properties are influenced by species-specific cellular context (for example ancillary proteins/subunits).

The comparison between immature hiPSC-CMs and more mature CMs derived from MTs allowed us to investigate the mechanism underlying the switch between foetal and adult SCN5A isoforms. Mbnl1 was previously identified as necessary for exon 6B expression in mice, as the Mbnl1-knockout mice expressed only the foetal isoform of SCN5A in the heart. ^{7,10} Here, we showed that MBNL1 expression is low in 2D hiPSC-CM monotypic cultures, while its expression increases significantly when CMs are included in MTs. Interestingly, only MBNL1 was specifically up-regulated in the CM population, and not MBNL2, another member of the same family. MBNL1 is also characterized by different developmentally regulated splicing isoforms and it is known to regulate its own splicing. 44,51 In particular, exon 5 is included in the embryonic isoform and is gradually excluded postnatally. 51,52 In MTs, we observed reduced expression of MBNL1 exon 5 compared with monolayer hiPSC-CMs, suggesting a functional change in MBNL1. Overall this indicates that MBNL1 is up-regulated and alternatively spliced in mature hiPSC-CMs and it might be specifically required to induce the switch in SCN5A isoform. Indeed, transient overexpression of MBNL1 increased SCN5A exon 6B fraction in hiPSC-CMs and lack of MBNL1 prevented splicing isoform switch in MTs. This confirms a key role for this factor in regulating SCN5A splicing also in human cells. Since we performed transient expression of MBNL1 in hiPSC-CMs, it remains to be elucidated whether the sole expression of the adult MBNL1 is sufficient to promote functional maturation of Nav1.5. Moreover, further studies are needed to understand how MBNL1 overexpression and knockout influence the expression of other target genes beside SCN5A. Indeed, Mbnl1 was shown to regulate the post-natal isoform switch of many other genes during mouse heart development 7-9 and to repress human stem cell pluripotency while promoting differentiation. 53,54 The expression of adult MBNL1 in hiPSC-CMs within the MT opens the possibility to study other developmentally/postnatally regulated genes using hiPSC-derived cells. The alternative splicing of the region of the voltage-sensing S4 segment is conserved also in voltage-gated sodium channels expressed in the brain, generating different isoforms underlying functional differences between foetal and adult tissues.55 lt is tempting to speculate that promoting hiPSC maturation in 3D organspecific tissues might facilitate the study of adult disease phenotypes in vitro in other organs.

In conclusion, our data demonstrate that a post-natal maturation of hiPSC-CMs is required to express the adult *SCN5A* isoform through MBNL1 regulation and thus reveal mutation contributions, allowing dissection of ionic current changes that cause adult arrhythmic disease phenotypes in humans.

Supplementary material

Supplementary material is available at Cardiovascular Research online.

Authors' contributions

M.B., C.L.M., and G.C.: conceptualization; G.C., G.K, A.O.V, D.O., and H.M.: formal analysis; C.L.M., M.B., and G.C.: funding acquisition; G.C., G.K., D.W.-v.O., L.Y., and A.O.V.: investigation; D.W.-v.O., M.B., and G.C.: methodology; M.B.: project administration; A.A.M.W., C.R.B., and C.C.V.: resources; M.B., A.O.V., R.P.D., and V.V.O.: supervision; and G.C., M.B., and C.L.M.: writing—original draft.

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Conflict of interest: C.L.M. is co-founder of Ncardia bv.

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Data Availability

Data are incorporated into the article and its Supplementary material online. RNA-seq raw data are deposited in GEO under the accession number GSE180290. All raw data are available from the corresponding author upon request.

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