

## ARTICLE

Received 2 Jun 2015 | Accepted 16 Feb 2016 | Published 11 Apr 2016

DOI: 10.1038/ncomms11067

OPEN

# Splicing misregulation of SCN5A contributes to cardiac-conduction delay and heart arrhythmia in myotonic dystrophy

Fernande Freyermuth<sup>1,\*†</sup>, Frédérique Rau<sup>2,\*</sup>, Yosuke Kokunai<sup>3</sup>, Thomas Linke<sup>4</sup>, Chantal Sellier<sup>1</sup>, Masayuki Nakamori<sup>3</sup>, Yoshihiro Kino<sup>5</sup>, Ludovic Arandel<sup>2</sup>, Arnaud Jollet<sup>2</sup>, Christelle Thibault<sup>1</sup>, Muriel Philipps<sup>1</sup>, Serge Vicaire<sup>1</sup>, Bernard Jost<sup>1</sup>, Bjarne Udd<sup>6,7,8</sup>, John W. Day<sup>9</sup>, Denis Duboc<sup>10</sup>, Karim Wahbi<sup>10</sup>, Tsuyoshi Matsumura<sup>11</sup>, Harutoshi Fujimura<sup>11</sup>, Hideki Mochizuki<sup>3</sup>, François Deryckere<sup>12</sup>, Takashi Kimura<sup>13</sup>, Nobuyuki Nukina<sup>14</sup>, Shoichi Ishiura<sup>15</sup>, Vincent Lacroix<sup>16</sup>, Amandine Campan-Fournier<sup>17</sup>, Vincent Navratil<sup>18</sup>, Emilie Chautard<sup>19</sup>, Didier Auboeuf<sup>19</sup>, Minoru Horie<sup>20</sup>, Keiji Imoto<sup>21</sup>, Kuang-Yung Lee<sup>22</sup>, Maurice S. Swanson<sup>23</sup>, Adolfo Lopez de Munain<sup>24</sup>, Shin Inada<sup>25</sup>, Hideki Itoh<sup>20</sup>, Kazuo Nakazawa<sup>25</sup>, Takashi Ashihara<sup>20</sup>, Eric Wang<sup>23</sup>, Thomas Zimmer<sup>4</sup>, Denis Furling<sup>2</sup>, Masanori P. Takahashi<sup>3</sup> & Nicolas Charlet-Berguerand<sup>1</sup>

Myotonic dystrophy (DM) is caused by the expression of mutant RNAs containing expanded CUG repeats that sequester muscleblind-like (MBNL) proteins, leading to alternative splicing changes. Cardiac alterations, characterized by conduction delays and arrhythmia, are the second most common cause of death in DM. Using RNA sequencing, here we identify novel splicing alterations in DM heart samples, including a switch from adult exon 6B towards fetal exon 6A in the cardiac sodium channel, SCN5A. We find that MBNL1 regulates alternative splicing of SCN5A mRNA and that the splicing variant of SCN5A produced in DM presents a reduced excitability compared with the control adult isoform. Importantly, reproducing splicing alteration of *Scn5a* in mice is sufficient to promote heart arrhythmia and cardiac-conduction delay, two predominant features of myotonic dystrophy. In conclusion, misregulation of the alternative splicing of SCN5A may contribute to a subset of the cardiac dysfunctions observed in myotonic dystrophy.

<sup>1</sup> Department of Translational medicine and neurogenetics, IGBMC, CNRS UMR7104, INSERM U964, Université de Strasbourg, Illkirch 67400, France. <sup>2</sup> Sorbonne Universités UPMC Univ Paris 06, Inserm, CNRS, Centre de Recherche en Myologie UMRS974/FRE3617, Institut de Myologie, GH Pitié-Salpêtrière, Paris 75013, France. <sup>3</sup> Department of Neurology, Osaka University Graduate School of Medicine, Osaka 565-0871, Japan. <sup>4</sup> Department of Physiology, Friedrich Schiller University Hospital, Jena 07743, Germany.

<sup>5</sup> Department of Bioinformatics and Molecular Neuropathology, Meiji Pharmaceutical University, Kiyose 205-8588, Japan. <sup>6</sup> Neuromuscular Research Center, Tampere University and University Hospital, Tampere 33520, Finland. <sup>7</sup> Department of Medical Genetics, Folkhälsan Institute of Genetics, Helsinki University, Helsinki 00250, Finland.

<sup>8</sup> Department of Neurology, Vaasa Central Hospital, Vaasa 65130, Finland. <sup>9</sup> Department of Neurology, Stanford University, Stanford, California 94304, USA. <sup>10</sup> Service de Cardiologie, Université Paris-Descartes, Hôpital Cochin, AP-HP, Paris 75014, France. <sup>11</sup> Department of Neurology, Toneyama National Hospital, Toyonaka 560-8552, Japan.

<sup>12</sup> CNRS UMR7175, Ecole Supérieure de Biotechnologies de Strasbourg, Illkirch 67400, France. <sup>13</sup> Division of Neurology, Hyogo Medical College, Nishinomiya 663-8501, Japan.

<sup>14</sup> Laboratory of Structural Neuropathology, Doshisha University Graduate School of Brain Science, Kyoto 610-0394, Japan. <sup>15</sup> Graduate School of Arts and Sciences, University of Tokyo, Tokyo 153-8902, Japan. <sup>16</sup> Université Lyon 1, CNRS, UMR5558 LBBE, Villeurbanne 69622, France. <sup>17</sup> Hôpices civils de Lyon, Laboratoire de cytogénétique constitutionnelle, Bron 69500, France. <sup>18</sup> Pôle Rhône Alpes de Bioinformatique, Université Lyon 1, Bâtiment Gregor Mendel, Villeurbanne 69100, France. <sup>19</sup> Centre de Recherche en Cancérologie de Lyon, Lyon 69373, France. <sup>20</sup> Department of Cardiovascular and Respiratory Medicine, Shiga Medical University, Otsu 520-2192, Japan. <sup>21</sup> Department of Information Physiology, National Institute for Physiological Sciences, Okazaki 444-8585, Japan. <sup>22</sup> Department of Neurology, Chang Gung Memorial Hospital, Keelung 20401, Taiwan. <sup>23</sup> Department of Molecular Genetics and Microbiology, Center for NeuroGenetics and the Genetics Institute, University of Florida, College of Medicine, Gainesville, Florida 32610, USA. <sup>24</sup> Department of Neurology, Hospital Universitario DONOSTIA, Neuroscience Area, Institute Biomedicina CIBERNED and University of Basque Country UPV-EHU, San Sebastián 20014, Spain. <sup>25</sup> Laboratory of Biomedical Sciences and Information Management, National Cerebral and Cardiovascular Center Research Institute, Osaka 565-8565, Japan. \* These authors contributed equally to the work. † Present address: Massachusetts General Hospital, MassGeneral Institute for Neurodegenerative Diseases, Charlestown, Massachusetts 02129, USA. Correspondence and requests for materials should be addressed to D.F. (email: denis.furling@upmc.fr) or to M.P.T. (email: mtakahas@neurol.med.osaka-u.ac.jp) or to N.C-B. (email: ncharlet@igbmc.fr).

**M**yotonic dystrophy (DM), the most common adult-onset muscular dystrophy, includes two genetically distinct forms. DM of type 1 (DM1) and its severe congenital form (CDM1) are caused by an expansion of CTG repeats in the 3'-untranslated region (UTR) of the *DMPK* gene<sup>1–3</sup>. In contrast, DM of type 2 (DM2) is caused by an expansion of CCTG repeats within the first intron of the *CNBP* (also known as *ZNF9*) gene<sup>4</sup>.

The pathogenesis of DM involves a RNA gain-of-function mechanism caused by expression of mutant RNAs containing hundred to thousands of CUG or CCUG repeats that interfere with the splicing of other pre-mRNAs through dysfunction of two classes of RNA-binding proteins. MBNL proteins (MBNL1, MBNL2 and MBNL3) are sequestered within nuclear RNA foci formed by expanded CUG and CCUG repeats<sup>5,6</sup>, whereas expression and phosphorylation of CUG-binding protein 1 (CUGBP1, encoded by the *CELF1* gene) are increased in DM1 heart samples<sup>7</sup>. MBNL and CUGBP1 proteins regulate alternative splicing, and alterations of their functional levels in myotonic dystrophic tissues results in reversion to fetal splicing patterns for several mRNAs, such as the insulin receptor (*INSR*) (ref. 8), the muscle chloride channel (*CLCN1*) (refs 9,10), dystrophin (*DMD*) (refs 11,12) and key components of the skeletal muscle excitation-contraction coupling process, including amphiphysin2 (*BIN1*) (ref. 13), ryanodine receptor 1 (*RYR1*) (ref. 14), sarcoplasmic/endoplasmic reticulum  $\text{Ca}^{2+}$ -ATPase SERCA1 (*ATP2A1*) (ref. 14) and the muscle calcium channel  $\text{Ca}_{\text{v}}1.1$  (*CACNA1S*) (ref. 15). Misregulation of the alternative splicing of the insulin receptor *INSR*, *CLCN1* and *DMD* mRNAs are associated with the insulin resistance<sup>8</sup>, myotonia<sup>9,10,16</sup> and dystrophic process<sup>12</sup>, respectively, while alterations of the alternative splicing of *BIN1*, *RYR1*, *ATP2A1* and *CACNA1S* may contribute to the skeletal muscle weakness observed in DM<sup>13–15</sup>.

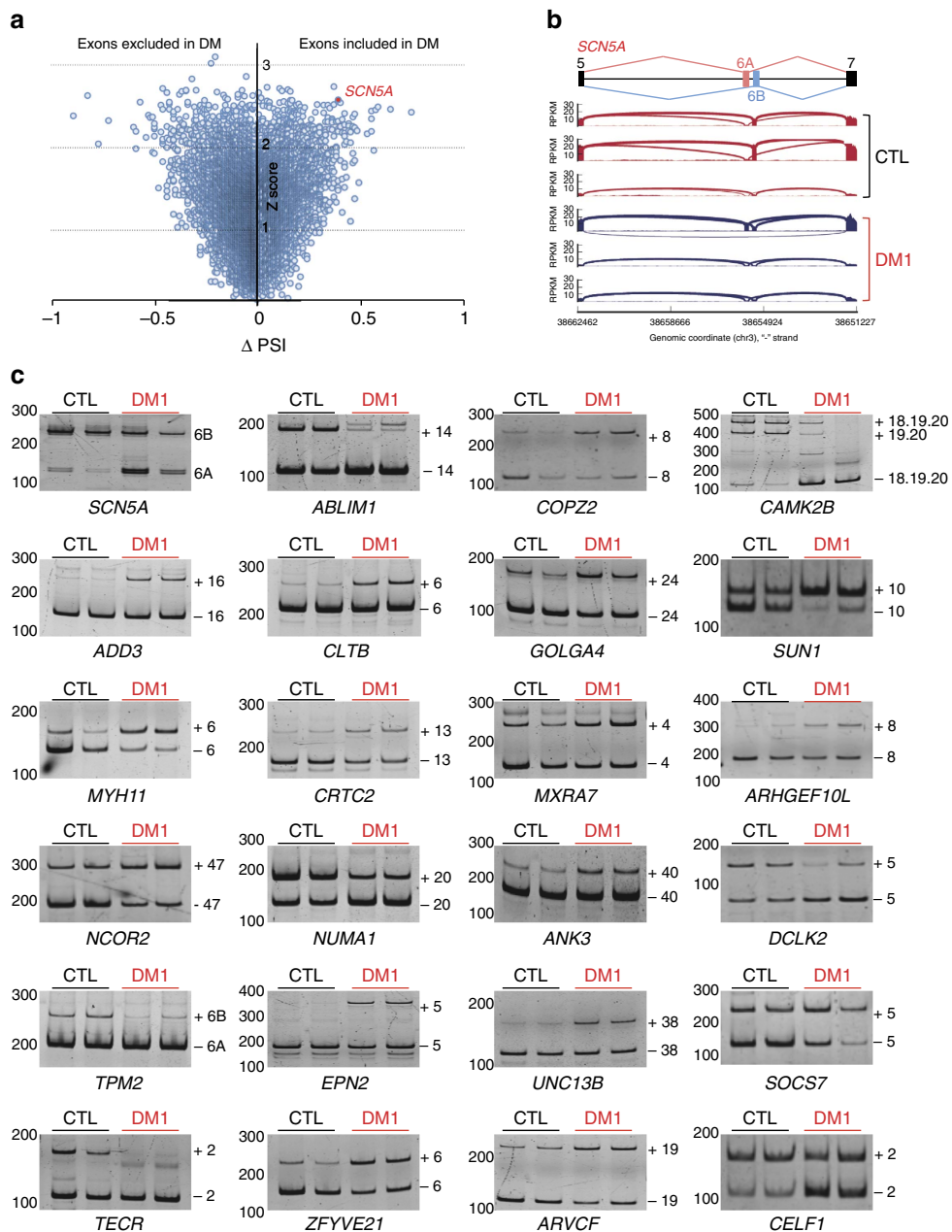
In contrast, the molecular mechanisms underlying the cardiac defects, which affect 80% of individuals with DM and represent the second most common cause of death in this disease<sup>17,18</sup>, are yet to be defined. Cardiac involvements in DM are characterized by cardiac-conduction delay that may result in fatal atrio-ventricular block, and by atrial or ventricular tachycardia<sup>17,18</sup>. Electrocardiography (ECG) analyses in DM patients indicate prolonged conduction time from the sinoatrial node to the ventricles (PR interval) and elongated ventricular depolarization (QRS duration). Interestingly, cardiac dysfunctions in DM are reminiscent in some aspect to an alteration of the cardiac sodium current. The  $\alpha$ -subunit of the cardiac voltage-gated  $\text{Na}^+$  channel,  $\text{Na}_v1.5$ , is encoded by the *SCN5A* gene and plays a key role in the excitability of cardiomyocytes and for rapid propagation of the impulse through the cardiac-conduction system. Mutations in *SCN5A* lead to a variety of arrhythmic disorders, including long QT3, progressive and non-progressive cardiac-conduction disease (also known as Lev-Lenègre disease), atrial fibrillation, sick sinus syndrome, Brugada syndrome and numerous overlapping syndromes<sup>19–21</sup>.

Using transcriptomic approaches, we identified various novel splicing changes in heart samples of DM1 individuals. Analysis of the RNA motifs enriched in the vicinity of these misregulated exons indicates that sequestration of the MBNL proteins is probably the main cause of splicing misregulation in heart of individuals with DM. Among these novel splicing alterations, we focused on misregulation of alternative splicing of the *SCN5A* pre-mRNA. This splicing alteration results in expression of a fetal isoform of *SCN5A* with altered electrophysiological properties. Of importance, we demonstrate that reproducing the splicing alteration of *Scn5a* in mouse is sufficient to cause heart arrhythmia and cardiac-conduction delay with elevated PR interval, which are key characteristics of the heart alterations

observed in DM. These results suggest that altered splicing of *SCN5A* mRNA may participate to the electrical cardiac abnormalities observed in DM.

## Results

**Identification of splicing changes in DM heart samples.** To determine novel splicing abnormalities in DM heart samples, we first used whole-genome microarrays (GeneChip Human Exon 1.0 ST array) on polyadenylated RNA extracted from left ventricle samples of three adult DM1 patients compared with three age-matched control individuals. Bioinformatic analyses predicted significant (Fold Change  $\geq 2$ , Student *t*-test, *P* value  $\leq 0.01$ ) changes in the splicing of 24 exons between control and DM1 samples (Supplementary Table 1), including a misregulation of the alternative splicing of the *SCN5A* pre-mRNA. To extend this analysis, we performed paired-end RNA sequencing (RNA-seq) on the same DM1 and control heart samples, yielding 1,611 million of mapped 100 bp reads. DESeq and Cuffdiff were then applied to estimate differential gene expression and over or under-expressed mRNAs were selected by using the Benjamini and Hochberg adjusted *P* values (false discovery rate (FDR)  $\leq 0.1$ ). A total of 9 and 19 upregulated genes were predicted differentially expressed with DESeq and Cuffdiff, respectively, but none were confirmed by quantitative real-time RT-qPCR analyses. This low number of differentially expressed mRNAs suggests that cardiac pathology in DM is not associated with drastic modifications of gene expression levels. In contrast, DEXSeq (ref. 22), which tests differential exon usage between two conditions, predicted 134 significant (Log2 Fold Change  $\geq 1.2$ , FDR  $\leq 0.1$ ) alternative splicing changes between control and DM1 heart samples (Supplementary Data 1). Similarly, MISO (ref. 23) analysis, which computes the fraction of mRNA that includes a given cassette alternative exon, predicted 259 significant ( $\Delta\text{PSI} \geq 0.3$ ; *Z*-score  $\geq 1.2$ ) alternative splicing changes between control and DM1 heart samples (Fig. 1a and Supplementary Data 2), including a robust misregulation of the alternative splicing of *SCN5A* (Fig. 1b). MISO and DEXSeq predictions overlapped, but with some exceptions, such as the skipping of the consecutive exons 18, 19 and 20 of *CAMK2B* predicted by DEXSeq but not by MISO; or the shift of *SCN5A* exon 6B towards exon 6A identified by MISO but not by DEXSeq. These differences are inherent to their computation models, since MISO does not detect alterations of successive exons and DEXSeq does not identify mutually exclusive exons, highlighting that MISO and DEXSeq are complementary bioinformatics approaches. Next, we tested by PCR with reverse transcription (RT-PCR) forty candidate mRNAs having the highest probability of misregulation in DEXSeq and/or MISO analyses. We validated splicing alterations for 32 of them, including some that have been identified in previous studies (*TNNT2*, *TNNT3*, *ABLM1*, *LDB3*, *MBNL1*, *CAMK2B*, *MAPT* and so on)<sup>24–26</sup>, and 20 others that represent, to the best of our knowledge, novel alterations of alternative splicing (*ADD3*, *GOLGA4*, *CRTC2*, *ARHGEF10L*, *ANK3*, *DCLK2*, *EPN2*, *UNC13B*, *TECR*, *ARVCF*, *SOCS7*, *CELF1* and so on) in DM1 heart samples (Fig. 1c). Of interest, some of these splicing alterations may be of pathological consequence in DM. For example, knockout of the *Socs7* gene in mouse results in insulin resistance<sup>27</sup>. Whether the splicing misregulation of *SOCS7* in DM contributes to insulin resistance remains to be tested. Also, RNA sequencing predicts an increased retention of the penultimate intron of *FCGRT*, which encodes the Fc fragment of the IgG receptor transporter  $\alpha$  (FCRN) protein, involved in IgG recycling<sup>28</sup>. Whether splicing alteration of *FCGRT* in DM is responsible to the decreased level of IgG in blood of these patients is an attractive hypothesis that remains to be tested.

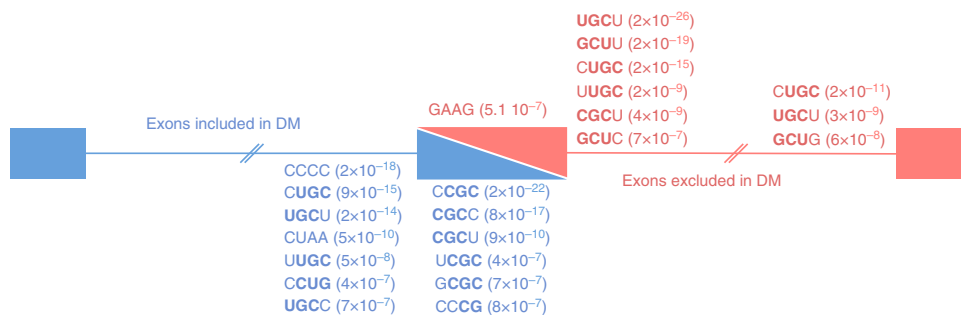


**Figure 1 | Identification of novel splicing misregulations in DM1 heart samples.** (a)  $\Delta$ -PSI versus Z-score plot of exon cassettes misregulations predicted by MISO analysis. (b) Exons structure and coverage of RNA-seq reads across *SCN5A* exons 5–7 show increased inclusion of exon 6A and decreased inclusion of exon 6B in heart samples of three DM1 patients (bottom, blue) versus three control samples (top, red). (c) Validation by RT-PCR of RNA-seq predictions in human heart samples of normal adult individuals (CTL, black) versus adult DM1 patients (DM1, red). Molecular size markers in bps are reported to the left of each RT-PCR gels. bp, base pair.

Finally, RNA sequencing predicts misregulation of the alternative splicing of a cardiac-specific exon located in the 5'-UTR of *CELF1*, which encodes CUGBP1. Whether this alternative splicing may contribute to the increase levels of CUGBP1 protein observed in DM1 hearts remains also to be evaluated.

**Splicing altered in DM are enriched for MBNL-binding sites.** Mutants RNAs containing expanded CUG or CCUG repeats interfere with the functional levels of CUGBP1 and MBNL proteins. Earlier studies determined that MBNL proteins bind to YGC RNA motifs (where Y is a pyrimidine)<sup>29–32</sup>, while CUGBP1 binds to UGU-enriched sequences<sup>33,34</sup>. To determine whether

these RNA motifs are indeed present in the vicinity of exons misregulated in DM, we determined all 4-mer RNA motifs enriched within, upstream or downstream of the exons predicted as misregulated by MISO in DM1 heart samples, compared with 2,000 control exons (Fig. 2). Most RNA motifs significantly enriched (binomial test,  $P$  value  $<1.0 \cdot 10^{-7}$ ) contained YGC sequences, while none were found to contain UGU sequences. Furthermore, YGC sequences were enriched upstream of exons abnormally included in DM1, while YGC motifs were enriched downstream of exons repressed in DM1. These results matched the MBNL splicing regulatory map determined by CLIP experiments, where binding of MBNL upstream of an exon tends to inhibit exon inclusion whereas binding of MBNL downstream



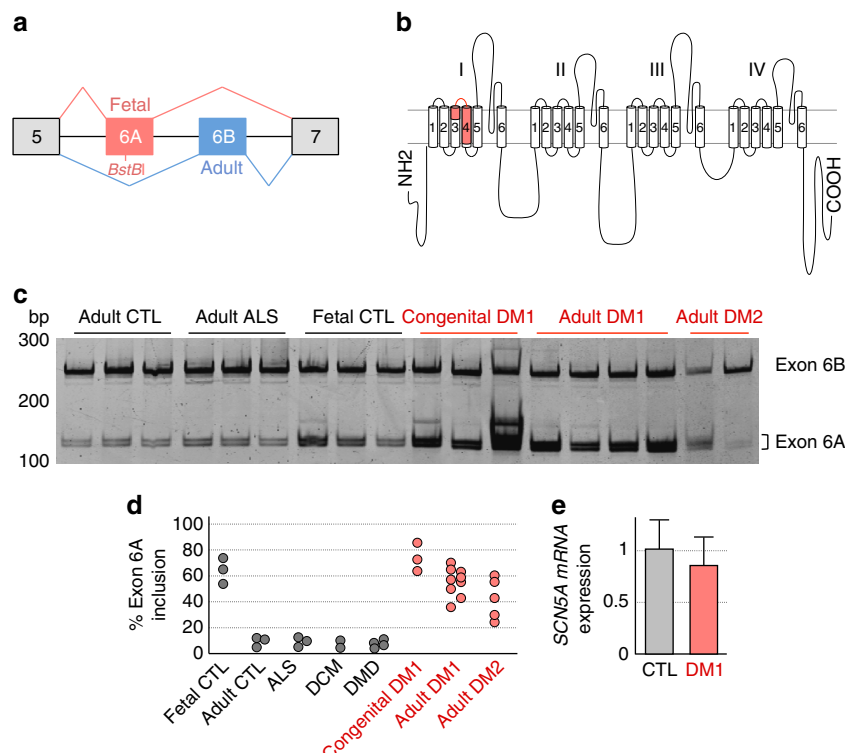
**Figure 2 | MBNL-binding motifs are enriched in vicinity of exons misregulated in DM1.** Sequence and binomial test *P* values of 4-mer RNA motifs enriched downstream, within and upstream of exons misregulated in DM1 heart samples. Sequences enriched in exons excluded in DM are indicated in red, while sequences enriched in exons included in DM are indicated in blue.

of an exon generally stimulates exon inclusion<sup>35,36</sup>. In contrast, we found no enriched motifs for other RNA-binding proteins, including CUGBP1, rbFOX1, hnRNP H or Staufen. These results, as well as previous data<sup>36–38</sup>, support a model in which titration of MBNL proteins is the main cause of splicing change in DM1 heart, while misregulation of other RNA-binding proteins may contribute to a subset of splicing alterations.

**Splicing of SCN5A is misregulated in DM heart samples.** Both microarray and RNA-seq predicted misregulation of alternative splicing of SCN5A pre-mRNA in DM1 heart samples. Splicing of SCN5A is developmentally regulated, such that exon 6A is included in fetal heart but rapidly replaced by exon 6B after birth<sup>39</sup>. Consequently, SCN5A exon 6A is named as embryonic or fetal, while exon 6B is known as adult. Exons 6A and 6B are mutually exclusive exons encoding part of the voltage sensor, segments 3 and 4 located in the domain I of the sodium channel (Fig. 3a,b). These are key segments for the electrical activity of the sodium channel, and inclusion of either fetal exon 6A or adult exon 6B results in channel isoforms, named, respectively, hNa<sub>v</sub>1.5e and hNa<sub>v</sub>1.5, with different electrophysiological properties<sup>39–41</sup>. We confirmed our microarray and RNA-seq predictions by RT-PCR and found that adult SCN5A exon 6B is partly replaced by its fetal exon 6A in heart samples of individuals with DM, including adult DM1 and adult DM2 cases (Fig. 3c). Note that to differentiate exon 6B from exon 6A that have the exact same length of 92 bp, we took advantage of a *Bst*BI restriction site present only in exon 6A, which thus appears as a *Bst*BI-digested doublet band in Fig. 3c. These results are consistent with the recent report of a splicing misregulation of SCN5A in one DM1 heart sample<sup>42</sup>. Although splicing of SCN5A is misregulated in DM1, we observed no correlation between the percentage of SCN5A exon 6A inclusion and the increased duration of the PR interval and only a very limited, if any, correlation between misregulation of SCN5A exon 6A splicing and alteration of the QRS duration in individuals with DM1 (*R*<sup>2</sup> of 0.2 with six DM1 samples; Supplementary Fig. 1). Misregulation of SCN5A splicing was specific to DM, as we did not observe inclusion of exon 6A in heart samples from individual affected with Duchenne muscular dystrophy (DMD), amyotrophic lateral sclerosis (ALS) or dilated cardiomyopathy (DCM) (Fig. 3d). Moreover, misregulation of splicing in DM1 was specific and not global, as we observed no splicing changes of SCN5A alternative exon 18, of CACNA1C mutually exclusive exons 8A and 8B, of KCNAB1 alternative exons 2 and 11, or of KCNQ1 alternative exons 2 and 5 (Supplementary Fig. 2). Finally, we observed no significant alteration of the expression level of SCN5A mRNA by quantitative real-time RT-qPCR (Fig. 3e). Overall, these results indicate a specific misregulation of

alternative splicing of SCN5A resulting in expression of a fetal form of this channel in adult DM heart. These results are consistent with previous studies where alternative splicing changes in DM resume a MBNL-dependent fetal splicing pattern that persist in adult tissues<sup>26,37</sup>.

**Alternative splicing of SCN5A is regulated by MBNL1.** To determine the mechanisms underlying misregulation of SCN5A splicing, we first determined its splicing pattern in cell models of DM. Since SCN5A is expressed at low level in culture of immature skeletal muscle cells, we investigated its splicing in primary cultures of differentiated skeletal muscle cells originating from muscle biopsies of control and DM1 individuals. RT-PCR experiments determined a switch of exon 6B towards exon 6A in DM1 muscle cells compared with control, reproducing the splicing alteration observed in cardiac tissue (Fig. 4a). Of technical interest, the basal level of exon 6A inclusion was higher in muscle cell cultures than in adult heart samples (compare Fig. 4a to Fig. 3c), which probably reflect the immature aspect of cell cultures. Since mutant RNAs containing expanded CUG or CCUG repeats interfere with alternative splicing through titration of MBNL proteins, we tested whether MBNL1 regulates SCN5A splicing. Reduction of *MBNL1* expression through a siRNA-mediated approach in human control primary muscle cells mimicked the effect of CUG repeats and promoted a switch from adult exon 6B towards fetal exon 6A (Fig. 4b). Western blotting analysis confirmed the successful depletion of MBNL1 expression (Supplementary Fig. 3A). Next, we assessed alternative splicing of *Scn5A* in heart samples of *Mbnl* knockout mice<sup>43</sup>. RT-PCR analysis shows that inclusion of the exon 6A of *Scn5A* is increased in heart samples of mice with no *Mbnl1* and reduced level of *Mbnl2* (*Mbnl1*<sup>-/-</sup>, *Mbnl2*<sup>+/-</sup>) (Fig. 4c). The increased inclusion of *Scn5A* exon 6A in *Mbnl* knockout mice is significant (Student *t*-test, *P* value  $\leq 0.01$ ) but rather mild, probably reflecting difference in regulation of alternative splicing between human and mouse or the compensatory effect of residual *Mbnl2* expression<sup>43</sup>. This hypothesis is consistent with the mild splicing alteration of *Scn5A* observed in the sole *Mbnl1* knockout mice<sup>44</sup>. Overall, these results suggest that MBNL proteins regulate the alternative splicing of SCN5A exons 6A and 6B. To determine whether this regulation is direct or indirect, we constructed a minigene containing exons 6A and 6B of SCN5A bordered by their intronic regions. Expression of this construct in mouse C2C12 myoblasts reproduced a fetal pattern with mainly inclusion of exon 6A (Fig. 4d). Since inclusion of exon 6B was repressed, reduction of *Mbnl1* activity through siRNA or expression of expanded CUG repeats had no further repressive effect on exon 6B. In contrast, expression of MBNL1 promoted a switch from fetal exon 6A towards adult exon 6B, while

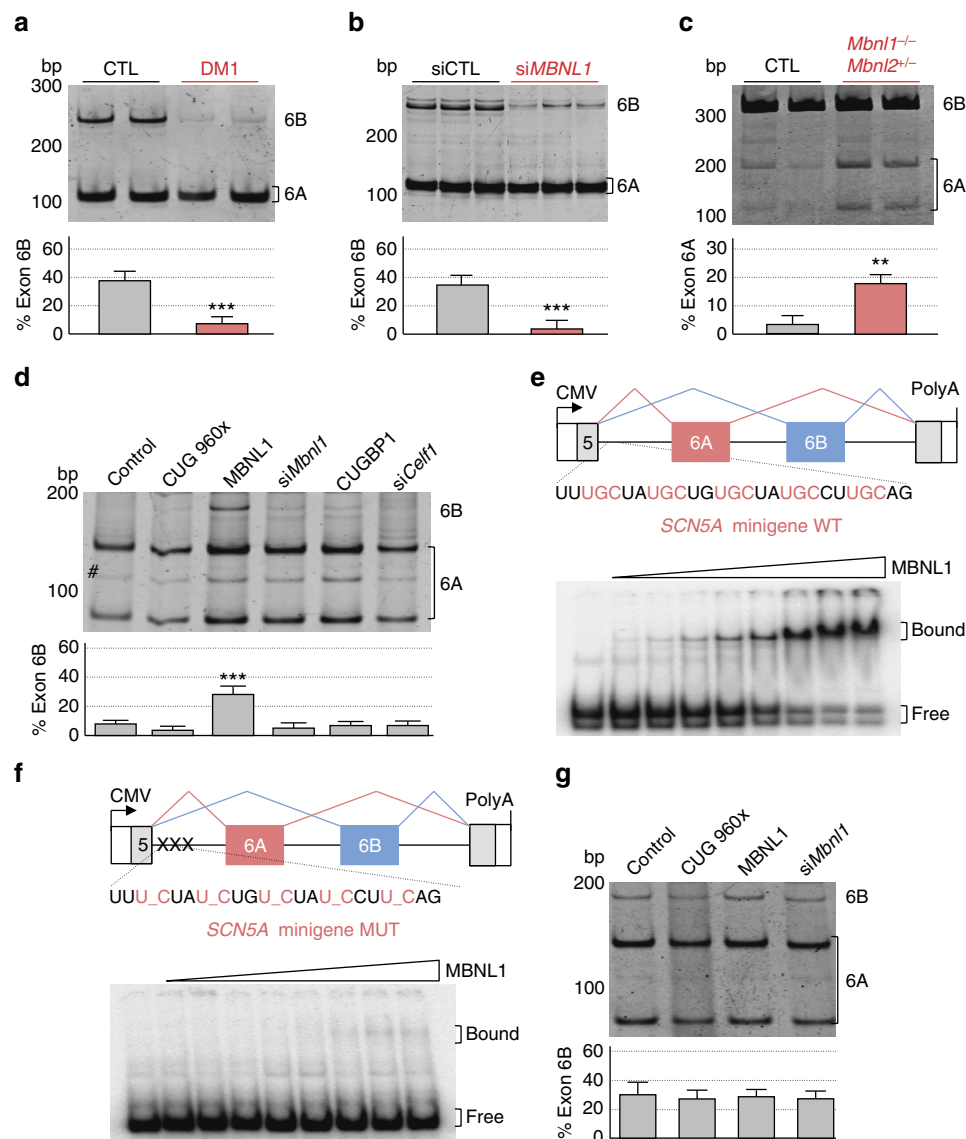


**Figure 3 | Splicing of SCN5A exon 6A is altered in DM heart samples.** (a) Schematic representation of mutually exclusive exons 6A and 6B of SCN5A. SCN5A mRNA includes exon 6A (red) in fetal heart, while SCN5A mRNA expresses exon 6B (blue) in adult heart. (b) Schematic representation of SCN5A topology expressing exon 6A (red). Exons 6A or 6B encodes part of segment 3, connecting loop between S3 and S4 and most part of the voltage-sensitive segment 4 of domain 1 of the sodium channel SCN5A. (c) Representative *Bst*BI-digested RT-PCR analysis of endogenous SCN5A mRNA from human heart samples of normal adult (CTL), adult ALS, non-DM fetuses (20, 24 and 35 weeks), congenital DM1 fetuses (CDM1 of 22, 25 and 28 weeks), adults DM1 and DM2 individuals. Molecular size marker is indicated in bp. (d) Graphical representation of RT-PCR analysis depicting the percentage of SCN5A mRNA including exon 6A in left ventricular heart samples from fetal and adult control, ALS, DCM, DMD, CDM1 and adult DM1 and DM2 individuals. (e) Graphical representation of quantitative real-time RT-qPCR depicting the mRNA expression of SCN5A relative to *RPLP0* in control normal adults ( $n=5$ ) versus adult DM1 ( $n=5$ ) heart samples. Bars indicate s.e.m. bp, base pairs.

expression or siRNA-mediated depletion of CUGBP1 had no effect (Fig. 4d). Western blotting analysis confirmed that siRNA transfection efficiently reduced endogenous Mbnl1 or Cugbp1 expression (Supplementary Fig. 3B and C). Next, gel-shift assays determined that recombinant purified GST-tagged MBNL1 bound to UGC RNA motifs located upstream of exon 6A (Fig. 4e). Of interest, this UGC sequence is absent from the mouse genome, which may explain the mild splicing alteration of *Scn5A* observed in mice knockout for Mbnl proteins. Mutation of these UGC motifs abolished MBNL1 binding (Fig. 4f), as well as the regulatory effect of MBNL1 on a mutant SCN5A minigene (Fig. 4g). Overall, these results establish that MBNL1 regulates directly alternative splicing of SCN5A exons 6A/6B.

**SCN5A splicing forms present different electrical properties.** SCN5A encodes  $\text{Na}_v1.5$ , the main cardiac voltage-gated sodium channel, and loss-of-function mutations in SCN5A lead to a variety of arrhythmic disorders, which share some common pathological features with DM. Furthermore, exons 6A and 6B differ at seven amino acid positions, resulting in channel variants with different electrophysiological properties<sup>39–41</sup>. To investigate the consequences of the switch from SCN5A exon 6B towards exon 6A observed in DM, we first examined in *Xenopus* oocytes the sodium currents generated by either h $\text{Na}_v1.5\text{e}$ , the splice variant of SCN5A containing the fetal exon 6A, or h $\text{Na}_v1.5$ , encoded by SCN5A containing the adult control exon 6B

(Fig. 5a and Supplementary Table 2). Injection of RNA encoding h $\text{Na}_v1.5\text{e}$ , which is the splicing isoform of SCN5A found in DM, indicated a significant reduction of the sodium current amplitude of 45%, compared with h $\text{Na}_v1.5$ , the normal adult SCN5A exon 6B form (Fig. 5b,c). Since, the extent of splicing misregulation varies among DM individuals, which typically express a mix of SCN5A splicing forms containing either exon 6A or exon 6B (cf. Fig. 3c), we analysed sodium currents generated by a mix of both SCN5A isoforms (Fig. 5a). Injecting *Xenopus* oocytes with an equimolar mix of RNA encoding each channel, namely 50% of h $\text{Na}_v1.5\text{e}$  (SCN5A containing fetal exon 6A) and 50% of h $\text{Na}_v1.5$  (SCN5A expressing adult exon 6B), resulted in a reduction of 30% of the current amplitude compared with the control h $\text{Na}_v1.5$  (Fig. 5b,c). Next, two-electrode voltage clamp recording experiments revealed that the steady-state activation of the fetal h $\text{Na}_v1.5\text{e}$  was shifted by 7 mV towards depolarized potential compared with the control adult h $\text{Na}_v1.5$  form (Fig. 5d and Supplementary Table 2). This shift is consistent with the shift observed previously in transfected mammalian cells<sup>39–41</sup>, thus validating our approach in *Xenopus* oocytes. To better reproduce the situation observed in DM, we injected in *Xenopus* oocytes an equimolar mix of DM (h $\text{Na}_v1.5\text{e}$ , fetal exon 6A) and control (h $\text{Na}_v1.5$ , adult exon 6B) RNA isoforms of SCN5A. Importantly, this mix of splicing forms also presented a significant shift of steady-state activation towards depolarized potentials by 3.8 mV, compared with the control h $\text{Na}_v1.5$  form (Fig. 5d, Supplementary Table 2). Correspondingly, a similar shift

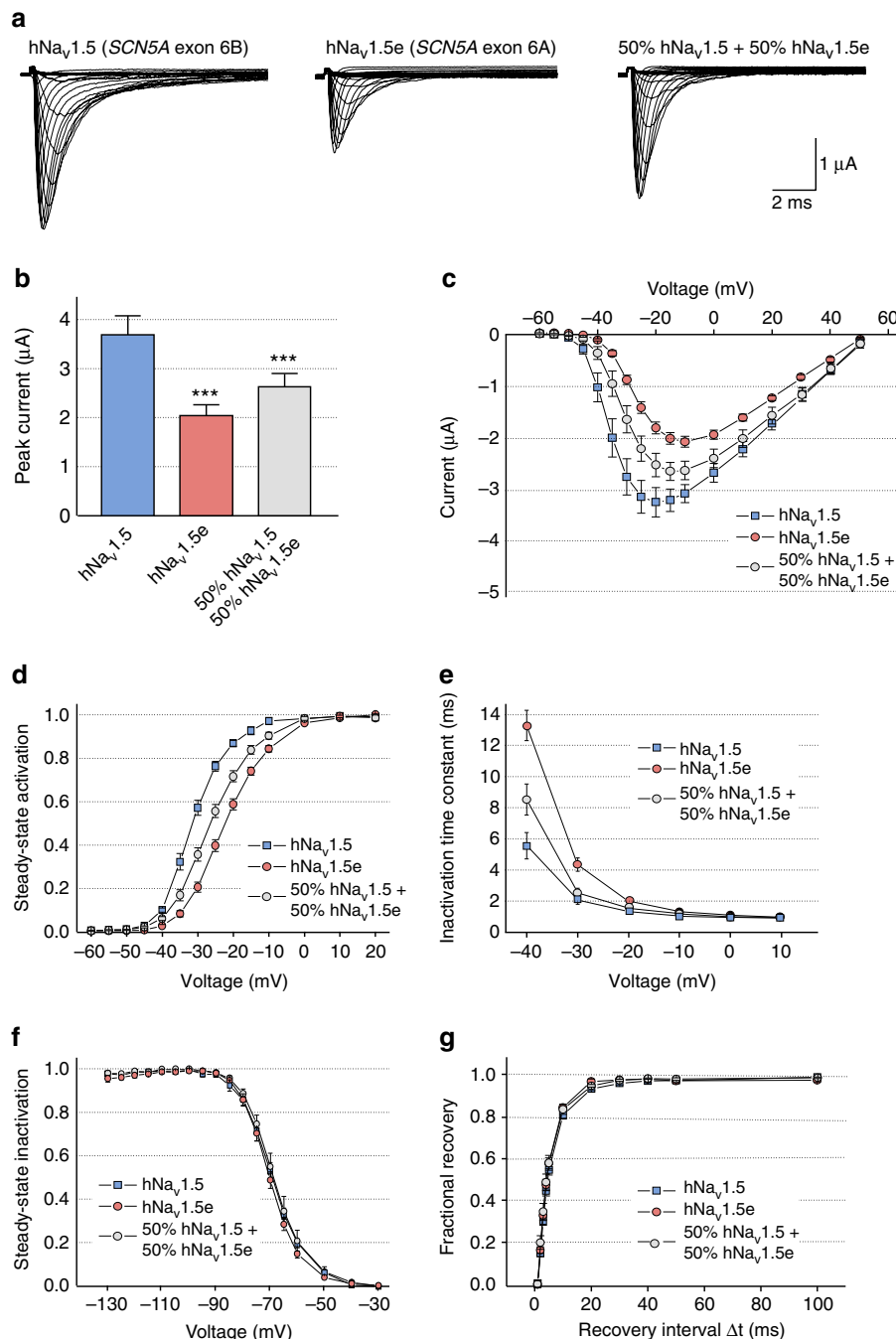


**Figure 4 | MBNL1 regulates alternative splicing of SCN5A.** (a) Upper panel, RT-PCR analysis of endogenous SCN5A mRNA from differentiated primary muscle cell cultures derived from biopsies of control or DM1 individuals. (lower) Quantification of the percentage of SCN5A mRNA including exon 6B. (b, upper) RT-PCR analysis of endogenous SCN5A mRNA from human differentiated cultures of control primary muscle cells transfected with a scrambled siRNA (siCTL) or a siRNA targeting MBNL1 mRNA (siMBNL1). (lower) Percentage of SCN5A mRNA including exon 6B. (c, upper) RT-PCR analysis of endogenous Scn5a mRNA in heart samples of wild-type and compound *Mbnl1*<sup>-/-</sup>, *Mbnl2*<sup>+/-</sup> double knockout mice. (lower) Percentage of *Scn5a* mRNA including exon 6A. (d, upper) RT-PCR analysis of exogenous SCN5A mRNA from differentiated C2C12 muscle cells co-transfected with a SCN5A minigene containing exons 6A and 6B bordered by their introns and with either a plasmid expressing 960 CTG repeats, MBNL1, CUGBP1 or with a siRNA directed against *Mbnl1* (siMbnl1) or *Celf1* (encoding Cugbp1; siCelf1). # Indicates usage of a cryptic splice site inherent to the minigene. (lower) Percentage of SCN5A mRNA including exon 6B. (e, upper) Schematic representation of SCN5A minigene, including the UGC-rich sequence used for binding assays. (lower) Gel-shift assays were performed using 5–1,000 nM of purified bacterial recombinant GST-MBNL1Δ<sup>101</sup> and a uniformly <sup>32</sup>P-CTP labelled RNA. (f, upper) Schematic representation of mutant SCN5A minigene, including the mutant sequence, used for binding assays. (lower) Gel-shift assay performed as in e. (g, upper) RT-PCR analysis of exogenous SCN5A mRNA from differentiated C2C12 muscle cells co-transfected with mutant SCN5A minigene and with a plasmid expressing 960 CTG repeats or MBNL1 or with a siRNA directed against *Mbnl1* (siMbnl1). (lower) Percentage of SCN5A mRNA including exon 6B. All transfection and gel-shift experiments were repeated three to five times. Molecular size markers are indicated in bp. Bars indicate s.e.m. Student test, \*\* indicates  $P < 0.01$ , \*\*\* indicates  $P < 0.001$ . bp, base pairs.

was observed for the time constant of inactivation (Fig. 5e). Consistent with previous electrophysiological studies<sup>39–41</sup>, no significant differences were observed between hNa<sub>v</sub>1.5 and hNa<sub>v</sub>1.5e regarding steady-state inactivation and recovery from inactivation (Fig. 5f,g). Overall, our results are consistent with previous studies<sup>39–41</sup>, and demonstrate that hNa<sub>v</sub>1.5e, the splicing form of SCN5A expressed in DM and containing the fetal exon

6A, presents a reduced excitability compared with hNa<sub>v</sub>1.5, which is the adult control SCN5A isoform containing exon 6B.

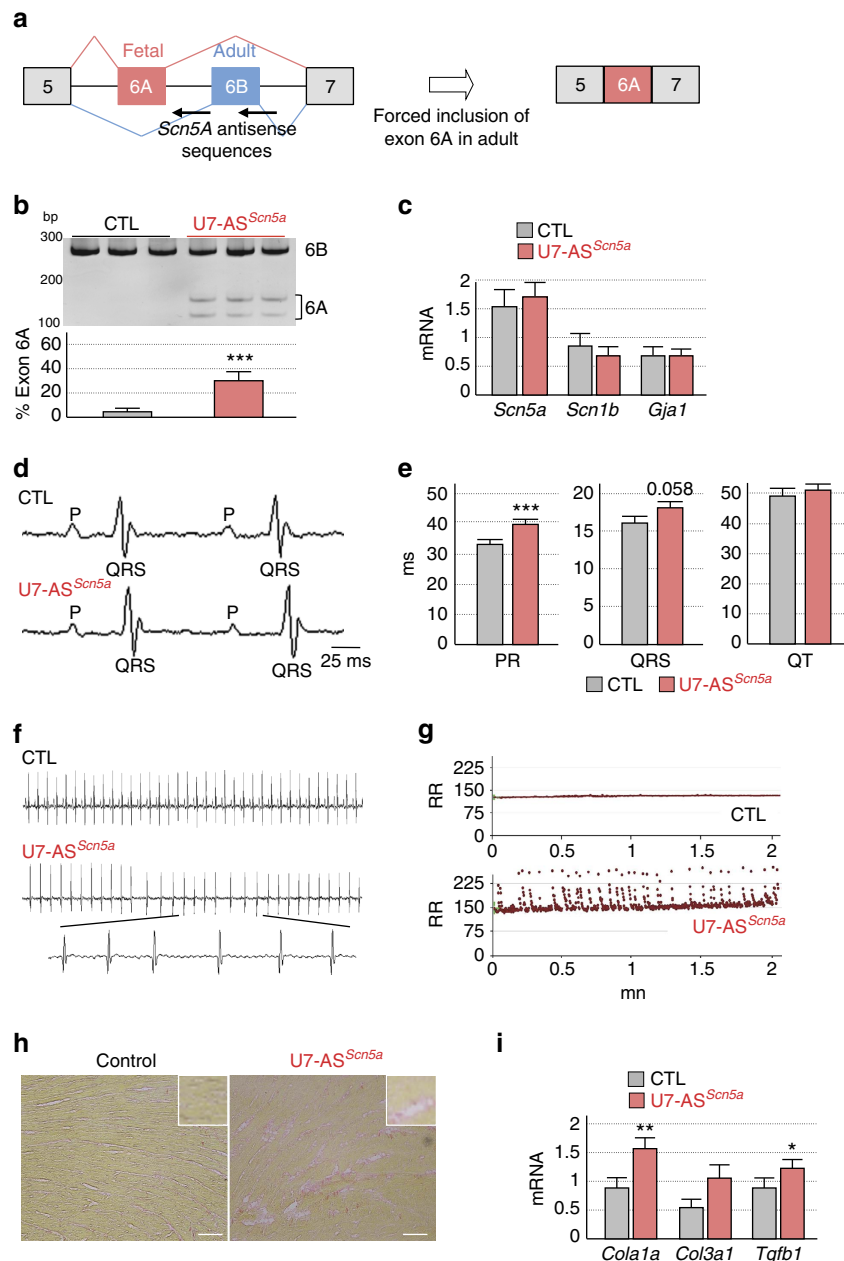
**Alteration of SCN5A splicing leads to heart conduction defects.** Misregulation of the alternative splicing of SCN5A in DM is one alteration identified among many others, thus questioning the



**Figure 5 | Electrophysiological properties of hNa<sub>v</sub>1.5 and hNa<sub>v</sub>1.5e channels.** (a) Representative Na<sup>+</sup> currents generated in *Xenopus* oocytes by hNa<sub>v</sub>1.5 (encoded by SCN5A containing the adult exon 6B), hNa<sub>v</sub>1.5e (encoded by SCN5A including the fetal exon 6A), and simultaneously expressed Na<sub>v</sub>1.5 and Na<sub>v</sub>1.5e channels at a 1:1 ratio. (b) Peak current amplitudes at the test potential of -10 mV in *Xenopus* oocytes injected with equimolar amount of cRNA encoding hNa<sub>v</sub>1.5, hNa<sub>v</sub>1.5e or 1:1 combination of Na<sub>v</sub>1.5 and Na<sub>v</sub>1.5e channels. (c) Current-voltage relationships. (d) Steady-state activation curves. (e) Inactivation time constants  $\tau_{\text{h}}$  (ms) at different test pulses. (f) Steady-state inactivation curves. (g) Fractional recovery curves. Data were obtained from 11 different batches of oocytes. To illustrate steady-state activation, steady-state inactivation and recovery from inactivation, we used 3–5 representative measurements. For total number of measurements ( $n=25$ –27) and for statistical data evaluation ( $V_m$ , s) see the Supplementary Table 2. Bars indicate s.e.m. Student test, \*\*\* indicates  $P < 0.001$ .

contribution of SCN5A misregulation to the cardiac symptoms observed in DM. To test the physiological importance of SCN5A splicing misregulation, we artificially forced the switch from adult exon 6B towards fetal exon 6A into wild-type adult mouse heart using an exon-skipping strategy (Fig. 6a). To insure efficient transduction of the cardiac muscle and continuous expression of nuclear antisense oligonucleotides, we engineered and

produced adeno-associated virus (AAV2/9) expressing optimized U7-snRNA fused to *Scn5a* antisense sequences (U7-AS<sup>Scn5a</sup>). Splicing analysis revealed that combination of two U7-AS constructs, spanning intron 6/exon 6B junction and exon 6B of *Scn5a*, promoted a switch from inclusion of adult exon 6B towards inclusion of the fetal exon 6A (Supplementary Fig. 4A). Thus, AAV2/9 expressing both U7-AS constructs



**Figure 6 | Alteration of *Scn5a* splicing causes heart conduction defects and arrhythmias.** (a) Schematic representation of mutually exclusive exons 6A and 6B of *Scn5a* and of antisense sequences driven by optimized U7-snRNAs (U7-AS<sup>Scn5a</sup>) to force fetal exon 6A inclusion in adult wild-type mouse heart. (b, upper) RT-PCR analysis of the alternative splicing of endogenous *Scn5a* mRNA from heart samples of mice injected with AAV2/9 expressing U7-AS<sup>Scn5a</sup> compared with control injected mice. Molecular size marker is indicated in bp. (lower) Percentage of *Scn5a* mRNA including exon 6A. (c) Real-time RT-qPCR quantification of the expression of *Scn5a*, *Scn1b* and *Gja1* (connexin 43) mRNAs in heart samples of mice expressing U7-AS<sup>Scn5a</sup> ( $n=6$ ) compared with control injected mice ( $n=6$ ). (d) Representative ECG traces show prolongation of the PR interval in U7-AS<sup>Scn5a</sup>-injected mice compared with control mice. (e) ECG measures of PR interval, QRS and QT intervals in 4-month-old mice injected with AAV2/9 expressing U7-AS<sup>Scn5a</sup> ( $n=25$ ) compared with age-matched control mice ( $n=17$ ). (f) Representative ECG traces reveal atrial fibrillation in U7-AS<sup>Scn5a</sup>-injected mice compared with control mice. (g) Variation of the RR interval indicates evidences of heart arrhythmias in U7-AS<sup>Scn5a</sup>-injected mice ( $n=25$ ) compared with control mice ( $n=17$ ). (h) Representative image of six analysed heart samples showing mild fibrosis revealed by Red Sirius histology staining in AAV-U7-AS<sup>Scn5a</sup>-injected mice. Scale bar, 100  $\mu$ m. (i) Real-time RT-qPCR quantification of the expression of *Cola1a*, *Col3a1* and *Tgfb1* mRNAs in heart of control ( $n=6$ ) or AAV-U7-AS<sup>Scn5a</sup>-injected mice ( $n=6$ ). Bars indicate s.e.m. Student test, \* indicates  $P<0.5$ , \*\* indicates  $P<0.01$ , \*\*\* indicates  $P<0.001$ . bp, base pair.

(AAV-U7-AS<sup>Scn5a</sup>) were injected systemically into newborn wild-type mice and cardiac functions were investigated 4 and 6 months post injection. Control animals injected either with saline or empty AAV2/9 presented no splicing alterations of *Scn5a* and normal cardiac functions. In contrast, mice injected with AAV-U7-AS<sup>Scn5a</sup> presented a decreased inclusion of adult exon 6B with

a concomitant 30–40% increase of the inclusion of exon 6A, thus reproducing the situation observed in DM (Fig. 6b). Quantitative RT-PCR demonstrated no changes in the expression of *Scn5a* mRNA or of its associated subunit *Scn1b* between control- and AAV-U7-AS<sup>Scn5a</sup>-injected mice (Fig. 6c). Importantly, AAV-U7-AS<sup>Scn5a</sup>-injected mice reproduce some of the key

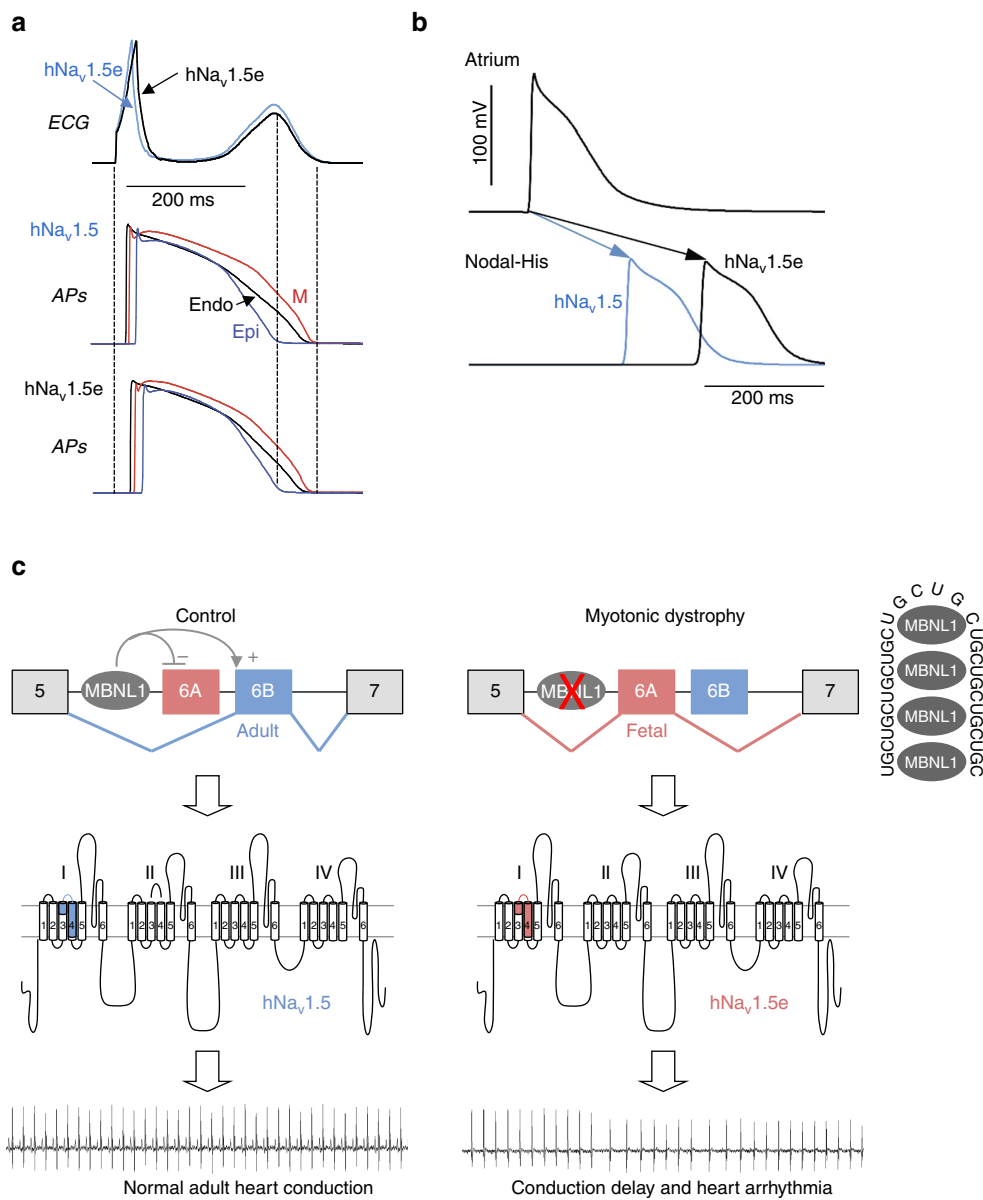
pathological features of DM, including conduction defects and heart arrhythmias. Indeed, ECG performed 4 months post injection revealed a significant prolongation of the PR intervals (Student *t*-test, *P* value  $\leq 0.001$ ) in AAV-U7-AS<sup>Scn5a</sup>-injected mice compared with control injected mice (Fig. 6d,e). In contrast, QT interval was not significantly altered, and we identified only a trend towards increased QRS duration (Student *t*-test, *P* value of 0.058 with 8 AAV-U7-AS<sup>Scn5a</sup>-injected mice on 25 presenting a QRS higher than 19 ms versus 16.5 ms in control mice) (Fig. 6e and Supplementary Fig. 4B). Similarly, analysis of heart functions in 6-month-old animals showed that AAV-U7-AS<sup>Scn5a</sup>-injected mice present a consistent increase of the PR interval compared with control injected mice (40.5 ms versus 34.8 ms respectively; Student *t*-test, *P* value  $\leq 0.05$ ), without significant changes of the QRS and QT intervals (Supplementary Fig. 4B). Of interest, a similar elongation of the PR interval was observed in *Scn5a*<sup>+/−</sup> mice, which are hemizygote for *Scn5a* expression and represent an established model for cardiac-conduction disease<sup>45–47</sup>. Furthermore, ECG analyses also revealed that 44% of AAV-U7-AS<sup>Scn5a</sup>-injected mice develop significant (Student *t*-test, *P*<0.001) heart arrhythmia at 4 months post injection with an average of five arrhythmic events, defined as variation of the RR interval, per minute whereas control injected animals showed no alterations (Fig. 6f,g). We did not detect ventricular fibrillations or second and third-degree heart blocks in any injected animals. In contrast, we observed supraventricular premature contractions and atrial fibrillation in AAV-U7-AS<sup>Scn5a</sup>-injected mice (Fig. 6f), and five of these injected mice died suddenly between 4 and 6 months post injections (none of the control mice died). These electrical alterations were specific and not caused by global cardiac remodelling since we observed neither systolic nor diastolic alterations by doppler echocardiography (Supplementary Table 3) and no change in heart/body weight ratio ( $4.4 \pm 0.1 \text{ mg g}^{-1}$  in control, *n*=9, versus  $4.7 \pm 0.2 \text{ mg g}^{-1}$  in AAV-U7-AS<sup>Scn5a</sup>-injected animals, *n*=14). As further control, H&E-staining revealed normal heart structures with no evident cardiomyopathy or dilation at 6 months post AAV injections (Supplementary Fig. 4C). Similarly, quantitative RT-PCR experiments show no alteration in the expression levels of *Nppa*, *Nppb* (encoding *Anp* and *Bnp*, respectively) and *Myh7* mRNAs (Supplementary Fig. 4D), suggesting no overt cardiac remodelling in antisense AAV-U7-AS<sup>Scn5a</sup>-injected mice. Moreover, Sirius Red staining confirmed normal heart structures but also revealed some mild fibrosis (Fig. 6h), which was confirmed by increased expression of collagen *Cola1a* and *Tgfb1* mRNAs (Fig. 6i). Interestingly, mild fibrosis is also observed in DM cardiac samples<sup>17,18</sup>, as well as in individuals and mice models with loss-of-function mutations of the *SCN5A* gene<sup>20,21,46,47</sup>. Overall, heart arrhythmias and prolonged PR interval in AAV-U7-AS<sup>Scn5a</sup>-injected animals demonstrate that inclusion of the fetal exon 6A of *Scn5a* is inappropriate to adult mouse heart physiology. However, while we found a clear elongation of the PR interval, we did not detect a significant alteration of the QRS duration as only a third of AAV-U7-AS<sup>Scn5a</sup>-injected mice present increased QRS duration ( $> 19 \text{ ms}$ ). Interestingly, similar findings have been described in *Scn5a*<sup>+/−</sup> mice, which all show elongation of the PR interval, while only a subset of *Scn5a*<sup>+/−</sup> animals present a prolongation of the QRS interval<sup>45</sup>. Hence, elongation of the PR interval is not systematically associated with increased duration of the QRS in mouse model of *Scn5a* dysfunction. Thus, to strengthen our data, we mathematically tested whether human cardiac parameters would be altered by the electrophysiological differences caused by the switch from adult exon 6B towards fetal exon 6A of *SCN5A*. Simulation based on a modified O'Hara-Rudy model<sup>48,49</sup>

predicted a change of the QRS duration from 72 ms with control adult hNa<sub>v</sub>1.5 to 88 ms with fetal hNa<sub>v</sub>1.5e, hence a 22% increase (Fig. 7a and Supplementary Fig. 5). Furthermore, we also tested extent of atrio-ventricular change<sup>50</sup>. Mathematical simulation predicted a change of the atrium-His interval from 81 ms with control hNa<sub>v</sub>1.5 to 143 ms with fetal hNa<sub>v</sub>1.5e (Fig. 7b). Overall, these results support our mouse results and provide additional evidences that misregulation of *SCN5A* alternative splicing causes cardiac-conduction abnormalities, which is a key pathological feature of DM (Fig. 7c).

## Discussion

Cardiac defects affect 80% of individuals with DM and represent the second most common cause of death in this disease<sup>17,18</sup>. However, the molecular mechanisms responsible for cardiac-conduction delay and ventricular tachycardia in DM are unclear. Using RNA sequencing we identified various novel splicing misregulation events in DM1 heart samples. Among these changes, the splicing switch from adult exon 6B to fetal exon 6A in *SCN5A* mRNA is of particular interest. Previous studies<sup>39–41</sup> as well as ours indicate that hNa<sub>v</sub>1.5e, the splicing variant of *SCN5A* found in DM and that contains the fetal exon 6A, possesses a reduced excitability compared with the normal adult splicing form of *SCN5A* containing the exon 6B. Consequently, the switch from the hNa<sub>v</sub>1.5 to the hNa<sub>v</sub>1.5e channel in DM may cause a slower upstroke velocity of the cardiac action potential, leading to conduction slowing. Importantly, this hypothesis is supported by mathematical simulation as well by animal model, since imposing a switch from inclusion of the control adult exon 6B towards using the fetal exon 6A of *Scn5a* in adult mouse heart led to cardiac-conduction delay and heart arrhythmias, two key features of DM. Moreover, clinical evidence also supports an alteration of the sodium current in DM. Indeed, the electrophysiological features<sup>39–41</sup> of the fetal isoform of *SCN5A* expressed in DM are similar to the electrophysiological characteristics observed with loss-of-function mutations of *SCN5A* causing cardiac-conduction disease<sup>51–53</sup>. Also, there are some similarities of ECG recording, including prolongation of the PR interval and of the QRS duration, between individuals with DM and individuals affected by cardiac-conduction disease caused by loss-of-function mutations in *SCN5A*<sup>42,54</sup>. Finally, the induction of abnormal ECG pattern in DM patients treated with ajmaline<sup>55,56</sup>, a class Ia antiarrhythmic agent acting on the cardiac sodium channel and the abnormal sodium current observed in a mouse model of DM<sup>57</sup>, are also evocative of a dysfunction of the sodium channel in DM. Overall, our results suggest that misregulation of the splicing of *SCN5A* participates in a subset of electrical cardiac alterations observed in DM, namely the cardiac-conduction delay and the heart arrhythmias. However, it is likely that other alternative splicing alterations and/or mechanisms<sup>58–61</sup> are participating to the full pattern of cardiac alterations in DM since knockout of *Mbnl1* and *Mbnl2* in mice leads to only mild alteration of *Scn5a* splicing, while these mice show severe conduction disease and cardiac dilatation<sup>43,44</sup>.

In conclusion, this work may also have some clinical importance such as considering with caution the treatments of DM patients with pharmaceutical agents that reduce the activity of the cardiac sodium channel, including mexiletine, flecainide and other antiarrhythmic drugs of class I. In that aspect, this study may provide a molecular explanation to the adverse cardiac reaction of some patients with myotonic dystrophic to treatment with drugs reducing activity of *SCN5A* (refs 62,63). Involvement of the cardiac sodium channel in DM might also highlight the importance of considering polymorphism in the *SCN5A* gene, as



**Figure 7 | Computer simulation predicts cardiac conduction abnormalities in DM.** (a) Simulations of ECG (upper) and action potential (AP) alterations (middle and lower) caused by the switch from adult *SCN5A* exon 6B towards fetal exon 6A, using a modified human ventricular ORd model. (b) Simulations of the atrio-ventricular changes caused by inclusion of *SCN5A* fetal exon 6A instead of adult exon 6B, employing an atrio-ventricular nodal model. APs of atrial and AV node cells were simulated and atrium-His interval was measured as the difference in the latency of APs between atrial and nodal-His cells. (c) Model of splicing alteration of the cardiac sodium channel, *SCN5A*, in DM. MBNL proteins regulate the switch from *SCN5A* exon 6A in fetal heart to exon 6B in adult. In DM, titration of MBNL proteins by mutant RNA containing expanded CUG repeats leads to expression of a fetal splicing form of *SCN5A*, inappropriate to adult heart physiology, ultimately resulting in cardiac-conduction delay and heart arrhythmias, which are two key features of DM.

well as in other genes such as *SCN10A*, as a possible cause of the clustering of cardiac alterations in some families of DM<sup>64</sup>, an hypothesis supported by the recent report that DM exacerbates Brugada syndrome caused by a mutation in *SCN5A* (ref. 65). Inversely, it remains to be tested whether mutations in intronic regions regulating alternative splicing of *SCN5A* exon 6A/6B might be considered as a cause of cardiac-conduction abnormalities in patients in whom no mutations were identified in the coding sequence of *SCN5A*. Also, if dysfunction of the cardiac sodium channel in DM is comparable in some aspects to cardiac-conduction disease caused by mutations in *SCN5A*, an attractive supposition would be that therapeutic approaches developed for these diseases could be considered for DM. Finally, if cardiac-conduction defects in DM are associated with an

alteration of the alternative splicing of *SCN5A*, one may hope that correction of splicing misregulations through approaches releasing MBNL1 from CUG expanded repeats or through approaches reducing the expression of pathogenic CUG RNA<sup>66,67</sup> may alleviate the cardiac symptoms of DM.

## Methods

**Human samples.** All samples were heart left ventricles that were sampled with the informed consent of individuals and approved by the Institutional Review Board of the Pitié-Salpêtrière hospital, of the Neuromuscular Research Center of the Tampere University Hospital, of the Hospital Donostia and of the Toyeyama National Hospital. Non-affected heart samples (CTL #1 to #3) were purchased at Ambion and Stratagene, respectively. DCM #1 and #2 were patients suffering from DCM of uncharacterized genetic origin. DMD #1 to #4 were patients affected with DMD. ALS #1 to #3 were patients with ALS described previously<sup>68</sup>. DM1 samples #1 and

2 were from cardiac transplantation for end stage heart failure with conduction system disease in a 45-year-old male and a 48-year-old female suffering from DM1. DM1 patient #3 is a 58-year-old female with 500 CTG repetitions in blood. DM1 patients #4 to #6 have been described previously<sup>68</sup> with expansion of 4,300 (female, 58 years); 4,800 (male, 63 years); 5,800 (female, 56 years); and ~6,000 CTG repeats in heart. DM1 samples #7 to #9 were patients with BAV1 and sadly unknown number of CTG repeats. DM2 patients #1 to #4 were described previously<sup>69,70</sup>. DM2 patients #5 heart sample was obtained at immediate autopsy after sudden bilateral renal thrombosis at the age of 71 years. He had ~4,800 CCTG repeats in leucocytes.

**mRNA sequencing.** RNA sequencing was performed on DM1 samples #4 to #6 described previously<sup>68</sup>. After isolation of total cellular RNA from human heart samples, libraries of template molecules suitable for high throughput DNA sequencing were created using 'TruSeq RNA Sample Preparation v2 Kit' (Illumina). Briefly, mRNA was purified from 1 µg total RNA using poly-T oligo-attached magnetic beads and fragmented using divalent cations at 94 °C for 8 min. The cleaved mRNA fragments were reverse transcribed to cDNA using random primers then the second strand of the cDNA was synthesized using DNA Polymerase I and RNase H. The double-stranded cDNA fragments were blunted using T4 DNA polymerase, Klenow DNA polymerase and T4 PNK. A single 'A' nucleotide was added to the 3' ends of the blunt DNA fragments using a Klenow fragment (3'- to 5'-exo minus) enzyme. The cDNA fragments were ligated to double-stranded adapters using T4 DNA Ligase. The ligated products were enriched by PCR amplification (30 s at 98 °C; (10 s at 98 °C, 30 s at 60 °C, 30 s at 72 °C) × 12 cycles; 5 min at 72 °C). Then surplus PCR primers were removed by purification using AMPure XP beads (Agencourt Biosciences Corporation). DNA libraries were checked for quality and quantified using 2100 Bioanalyzer (Agilent). The libraries were loaded in the flow cell at 7 pM concentration and clusters were generated in the Cbot and sequenced in the Illumina Hiseq 2000 as paired-end 2 × 100 base reads. FASTQ files were generated with CASAVA v1.8.2.

**Bioinformatic analysis.** A total of 1,857,442,522 short reads were mapped onto the human genome (hg19 assembly) by using TopHat2 software<sup>71</sup> (release v2.0.5) with a mapping rate ranging from 83 to 90%. Htsq-count (release 0.5.3p3). The Cufflinks software<sup>72</sup> (release 2.0.2) was used to call the gene expression level according to the ensembl gene annotation (release 66). DEXseq software<sup>22</sup> (release v1.4.0) was then used to identify differential exons usage and significant over and under-expressed genes were selected by using Benjamini & Hochberg adjusted P value (FDR ≤ 0.1).

**Computer simulation of action potential durations and ECG.** Two mathematical models, O'Hara-Rudy (ORD) dynamic model<sup>48,49</sup> of ventricular myocytes and rabbit atrio-ventricular node model<sup>50</sup>, were employed to simulate action potential of cardiac cells and ECG. For the simulation, parameters of the activation gates for control hNav1.5, an equimolar mix of hNav1.5 and hNav1.5e, and hNav1.5e were estimated to match the experimental data obtained with *Xenopus* oocytes. Na currents and curves of voltage dependence of control hNav1.5, an equimolar mix of hNav1.5 and hNav1.5e, and hNav1.5e were reconstructed using these estimated parameters (Supplementary Fig. 5A–D). The parameters for the inactivation gate were unchanged. To get ECG similar to the left precordial ECG, a unipolar recording electrode was located 2.6 cm far from the epicardial border of the ventricular wedge preparation model (Supplementary Fig. 5E). The model consists of endocardial (Endo), midmyocardial (M) and epicardial (Epi) layers of thicknesses 0.12, 0.65 and 0.24 cm, respectively. The membrane kinetics and the transmural heterogeneity were based on the ORD model<sup>48</sup> with twice sodium channel conductance to reproduce physiological ranges of both ventricular conduction velocity (~60 cm s<sup>-1</sup> along the fibre) and tissue excitability. The pacing stimuli of 2 ms and twice diastolic threshold were applied transmembranously to the part of endocardial border at 1 Hz to achieve steady state. The action potentials of Endo, M and Epi are recorded at the centre of each myocardial layer. Temperature employed in the simulation was 37 °C. Ionic concentrations employed in the simulation were 96 mM NaCl, 2 mM KCl, 1.8 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, 10 mM HEPES/KOH and pH 7.4. Other model parameters and the numerical approach have been described elsewhere<sup>49</sup>. One-dimensional mathematical model<sup>50</sup> of atrio-ventricular node was employed to simulate the conduction between atrium to ventricle. Although this model was based on the experimental data in rabbit heart, this is, in our knowledge, the only model for AV conduction. The simulation parameters for the equimolar mix of hNav1.5 and hNav1.5e and hNav1.5e were set to reproduce the shift inactivation and the decrease of peak Na current observed in oocytes. The multicellular model consisted of atrium and AV node, and action potentials of atrium and AV node cells were simulated with the stimulus intervals of 350 ms.

**Construction and analysis of the SCN5A minigenes.** Exons 5, 6A, 6B and 7 were PCR amplified from human DNA (Clontech) with FWD primer: 5'-aaaaggctagcgatccatcaccattacacc-3' and REV primer: 5'-aaaaggccggatactgggttaaggctca gg-3', and inserted between the NheI and NotI restriction sites of pCDNA3.1+ (Invitrogen). Mutations in SCN5A intron 5 were introduced by primer-directed PCR

mutagenesis. C2C12 cells plated in 6-well plates were co-transfected with Lipofectamine 2000 (Invitrogen) according to the manufacturer's instructions with SCN5A minigene and DMPKS DT960, V5-CUGBP1, V5-MBNL1 40 kDa (NM\_207292) and siRNA directed against MBNL1 (5'-CACGGAAUGUAAAUA UGCAdTdT-3'; Eurogentec) or against CUGBP1 (NM\_198700; NM\_006560; NM\_001025596; NM\_001172640; NM\_001172639; Thermo Scientific) in DMEM medium containing 1 g per l glucose and 2% horse serum (37 °C, 5% CO<sub>2</sub>) during 24 h. Total RNAs were extracted using Tri Reagent (MRC) and subjected to reverse transcription using Transcripter Reverse Transcriptase kit (Roche). PCR were performed using Taq polymerase (Roche), one denaturation step at 94 °C for 2 min, 26 cycles of amplification 94 °C for 1 min, 58 °C for 30 s, 72 °C for 45 s and a final step at 72 °C for 7 min using the forward primer (5'-cttctgcctgcacgcgttcac-3') and the reverse (5'-acggggcccttagctcg-3') specific to pCDNA3.1- SCN5A minigene. The PCR products were digested by *Bst*BI enzyme. PCR products of mRNAs including exon 6A and 6B are of 116 + 71 bp and 187 bp, respectively. PCR products were precipitated, analysed by electrophoresis on 6.5% polyacrylamide gel, BET labelled and quantified using Typhoon scanner.

**Electrophysiological measurements.** Ovarian lobes were obtained from *Xenopus laevis* under anaesthesia (0.4% 3-aminobenzoic acid ethyl ester) and transferred to Ca<sup>2+</sup>-free Barth medium (84 mM NaCl, 1 mM, 2.4 mM NaHCO<sub>3</sub>, 0.82 mM MgSO<sub>4</sub> and 7.5 mM Tris/HCl pH 7.4). Oocytes in stages V and VI were incubated in Ca<sup>2+</sup>-free Barth medium containing 1.2 mg ml<sup>-1</sup> collagenase for 60 min. After washing repeatedly with Barth medium, the oocytes were isolated and defolliculated mechanically. Plasmids containing cDNAs of hNav1.5 or hNav1.5e were linearized using the *Not*I restriction enzyme and *in vitro* transcribed using the T7 RNA polymerase. Concentrations of the cRNA variants were adjusted to ~0.002 µg µl<sup>-1</sup> using the gel documentation system from Herolab (Wiesloch, Germany) and a reference RNA sample (0.24–9.5 kb RNA ladder, Invitrogen Life Technologies), before injecting ~50–80 nl cRNA per oocyte. After 3 days incubation at 18 °C in Barth medium, the peak current amplitude of the whole-cell Na<sup>+</sup> current was between 0.5 and 7.5 µA, depending on the quality of the oocyte batch. Cells producing currents larger than 6 µA were not selected for the evaluation of the channel kinetics. Measurements were performed in totally 11 different batches of oocytes. Whole-cell Na<sup>+</sup> currents were recorded with the two-microelectrode voltage-clamp technique using the TEC-05-S amplifier (npi electronic GmbH, Tamm, Germany) and the following bath solution (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, 10 mM HEPES/KOH and pH 7.4). Currents were elicited by test potentials from –80 to 20 mV in 5 or 10 mV increments at a pulsing frequency of 1.0 Hz (holding potential –120 mV). Steady-state activation ( $m_\infty$ ) was evaluated by fitting the Boltzmann equation  $m_\infty = \{1 + \exp[-(V - V_m) / s]\} - 1$  to the normalized conductance as function of voltage, where V is the test potential, V<sub>m</sub> the mid-activation potential, and s the slope factor in mV. Recording and analysis of the data was performed on a personal computer with the ISO3 software (MFK, Niedernhausen, Germany). The sampling rate was 50 kHz. Student's t-test was used to test for statistical significance. Statistical significance was assumed for  $P < 0.05$ .

**In vivo gene transfer.** All mouse procedures were done according to protocol approved by the Committee on Animal Resources at the Centre d'Exploration Fonctionnelle of UPMC animal facility and under appropriate biological containment. AAV2/9 vectors expressing *Scn5a* antisense sequences driven by an optimized U7-snRNA (U7-AS) construct described previously<sup>73</sup> were produced by co-transfection in HEK293 cells of pAAV2(U7-AS), pAD-DELTA-F16 encoding adenovirus helper functions and p5E18-VD2/9. Vector particles were purified on iodixanol gradients from cell lysates obtained 72 h after transfection, and titres were measured by RT-qPCR. The vector preparation used in this study had a titer of  $1 \times 10^{10}$  EXP13 vector genomes (vg) ml<sup>-1</sup>. U7-AS constructs that contain antisense sequences to target intron 6/exon 6B border or exon 6B of *Scn5a* were cloned in the same AAV backbone. 2 days old C57BL/6 mice were injected in the temporal vein for systemic delivery with 50 µl of either with saline, empty AAV2/9 or AAV2/9-U7-AS ( $1 \times 10^{12}$  vector genomes ml<sup>-1</sup>). Six months after injections, mice were killed, hearts were collected and snap-frozen in liquid nitrogen-cooled isopentane and stored at –80 °C.

**ECG recordings and echocardiographic examination.** Mice were anaesthetized with isoflurane and surface ECG was recorded using 23-gauge, subcutaneous needle electrodes (Gould Instrument Systems), affixed to each limb. ECG was continuously monitored for 3 min (software IOX 2, EMKA Technologies, Paris France). Heart rate, PR and QRS duration were measured from recording traces using ECG auto V.1.5.7 software (Emka Technologies). Non-invasive measurements of left ventricular dimensions were evaluated under isoflurane anaesthesia, using echocardiography-Doppler (General Electric Medical systems Co, Vivid 7 Dimension/Vivid 7 PRO) with a probe emitting ultrasounds with 9–14 MHz frequency. The two-dimensionally guided Time Motion mode recording of the left ventricle provided the following measurements: diastolic and systolic septal and posterior wall thicknesses, internal end-diastolic and end-systolic diameters, ejection fraction, fractional shortening of left ventricular diameter. Each set of

measurements was obtained from a same cardiac cycle. At least three sets of measures were obtained from three different cardiac cycles.

**Recombinant protein production and purification.** *Escherichia coli* BL21(RIL) pRARE competent cells (Invitrogen) were transformed with peGEX-MBNL1 $\Delta^{101}$ , which is a C-terminal truncated form of MBNL1 with higher solubility but identical affinity than MBNL1 full-length, (gift from Prof. C. Branstetter), grown at 37 °C in 400 ml of LB medium supplemented with Kanamycin until OD600 = 0.5, 0.5 mM IPTG was added and the culture was further incubated 4 h at 30 °C. Harvested cells were sonicated in 50 mM Tris-Cl pH 7.5, 300 mM NaCl, 5% glycerol, 1 mM DTT, 5 mM EDTA, centrifuged 20 min at 20,000g and recombinant GST-MBNL1 $\Delta^{101}$  protein was purified using GST-Kits (Novagen) and stored in 20% glycerol.

**RNA transcription and gel-shift assays.** Templates for transcription were obtained by PCR with pCDNA3.1-SCN5A or pCDNA3.1-SCN5A mutant minigenes and the primers (FWD that include a T7 promoter sequence: 5'-TAA TACGACTCACTATAGGGCCATGGAACTGGCTGGACTTT-3' and REV: 5'-AGAAAGAGAGGGGTGGTCG-3'). Transcription reaction was performed using T7 transcription kit (Ambion) in presence of 1  $\mu$ l of  $\alpha$ P32-CTP (10  $\mu$ Ci, 800 Ci per mmole, Perkin Elmer), analysed on 8% denaturing polyacrylamide and quantified with LS-6500 counter (Beckman). After transcription, 1 unit of DNase I (Invitrogen, Carlsbad, CA) was added, and the sample was incubated for additional 30 min at 37 °C. Transcribed RNAs were then purified by micro Bio-Spin 6 chromatography columns (Bio-rad) according to the manufacturer's instructions. The sizes of RNAs were checked by gel electrophoresis on a denaturing 6% polyacrylamide gel. For gel shift, 1  $\mu$ l of labelled RNA was annealed in binding buffer (BB, 50 mM Tris-HCl (pH 7.0), 75.0 mM NaCl, 37.5 mM KCl, 5.25 mM DTT, 0.1 mg ml $^{-1}$  BSA, 0.1 mg ml $^{-1}$  *E. Coli* tRNA) at 90 °C for 5 min. and allowed to cool to room temperature by placing on a bench top. After cooling, MgCl $_2$  and RNasin were added to a final concentration of 0.75 mM and 0.4 U ml $^{-1}$  respectively. GST-MBNL1- $\Delta^{101}$  protein was then added to the above solution and the mixture was incubated at 25 °C for 20 min. The solution mixture was loaded onto a non-denaturing 6.0% (w/v) polyacrylamide gel (acrylamide/bisacrylamide, 40:1, w/w) containing 0.5  $\times$  TBE (1X TBE is 90 mM Tris-base, 89 mM Boric acid and 2 mM EDTA (pH 8.0)), which had been pre-electrophoresed at 110 V for 20 min. at 4 °C. The gel was electrophoresed at 110 V at 4 °C for 3 h. The gel was then dried and exposed to a phosphorimager screen and imaged using a Typhoon 9410 variable mode imager. The data were fit to the following equation:  $y = \min + ((\max - \min) / (1 + (x/IC50) \cdot \text{HillSlope}))$ , where  $y$  is the percentage of RNA bound to MBNL1,  $x$  is the concentration of MBNL1, min and max are the minimum and maximum percentage of RNA bound to MBNL1 (0–100%) and IC50 is the concentration where 50% of maximum binding is achieved.

**RT-PCR analyzes.** Total RNA was isolated by TriReagent (Molecular Research Center). A total of 1  $\mu$ g of total RNA isolated from heart samples of control or DM individual was subjected to reverse transcription using random hexanucleotide and the Transcripter High Fidelity cDNA synthesis kit (Roche Diagnostics). PCR was performed with Taq polymerase (Roche), using the following protocol: one denaturation step at 94 °C for 2 min, 30 cycles of amplification 94 °C for 1 min, 60 °C for 1 min, 72 °C for 2 min and a final step at 72 °C for 5 min using primers described in the Supplementary Table 4. The PCR products were precipitated, analysed by electrophoresis on a 6.5% polyacrylamide gel, stained with ethidium bromide and quantified using a PhosphorImager Typhoon scanner (GE Healthcare) and the ImageQuaNT software. Concerning analysis of alternative splicing of human SCN5A, the forward primer was located within SCN5A exon 5 (FWD: 5'-CTTCT GCCTGCACCGCTTAC-3') and the reverse primer within SCN5A exon 7 (REV: 5'-CAGAAGACTGTGAGGACCATC-3'). Since SCN5A exons 6A or 6B are of similar size (92 nts), PCR products were digested by *Bst*BI enzyme before loading on 6.5% polyacrylamide gel, resulting in PCR products of 240 bp (exon 6B) and a doublet of 115 bp and 125 bp (exon 6A digested by *Bst*BI).

**Quantitative real-time PCR analyzes.** Total RNA was isolated by TriReagent (Molecular Research Center). cDNAs were generated using the Transcripter High Fidelity cDNA synthesis kit (Roche Diagnostics). Real-time qPCR were realized using the LightCycler 480 SYBR Green I Master (Roche) in a Lightcycler 480 (Roche) with 15 min at 94 °C followed by 50 cycles of 15 s at 94 °C, 20 s at 58 °C and 20 s at 72 °C using primers described in the Supplementary Table 5. *Rplpo* mRNA was used as standard and data were analysed using the Lightcycler 480 analysis software (2Δ $\Delta$ Ct method).

## References

- Brook, J. D. *et al.* Molecular basis of myotonic dystrophy: expansion of a trinucleotide (CTG) repeat at the 3' end of a transcript encoding a protein kinase family member. *Cell* **68**, 799–808 (1992).
- Fu, Y. H. *et al.* An unstable triplet repeat in a gene related to myotonic muscular dystrophy. *Science* **255**, 1256–1258 (1992).
- Mahadevan, M. *et al.* Myotonic dystrophy mutation: an unstable CTG repeat in the 3' untranslated region of the gene. *Science* **255**, 1253–1255 (1992).
- Liquori, C. L. *et al.* Myotonic dystrophy type 2 caused by a CCTG expansion in intron 1 of ZNF9. *Science* **293**, 864–867 (2001).
- Miller, J. W. *et al.* Recruitment of human muscleblind proteins to (CUG)n expansions associated with myotonic dystrophy. *EMBO J.* **19**, 4439–4448 (2000).
- Kanadia, R. N. *et al.* A muscleblind knockout model for myotonic dystrophy. *Science* **302**, 1978–1980 (2003).
- Kuyumcu-Martinez, N. M., Wang, G. S. & Cooper, T. A. Increased steady-state levels of CUGBP1 in myotonic dystrophy 1 are due to PKC-mediated hyperphosphorylation. *Mol. Cell* **28**, 68–78 (2007).
- Savkur, R. S., Philips, A. V. & Cooper, T. A. Aberrant regulation of insulin receptor alternative splicing is associated with insulin resistance in myotonic dystrophy. *Nat. Genet.* **29**, 40–47 (2001).
- Mankodi, A. *et al.* Expanded CUG repeats trigger aberrant splicing of ClC-1 chloride channel pre-mRNA and hyperexcitability of skeletal muscle in myotonic dystrophy. *Mol. Cell* **10**, 35–44 (2002).
- Charlet-B, N. *et al.* Loss of the muscle-specific chloride channel in type 1 myotonic dystrophy due to misregulated alternative splicing. *Mol. Cell* **10**, 45–53 (2002).
- Nakamori, M., Kimura, T., Fujimura, H., Takahashi, M. P. & Sakoda, S. Altered mRNA splicing of dystrophin in type 1 myotonic dystrophy. *Muscle Nerve* **36**, 251–257 (2007).
- Rau, F. *et al.* Abnormal splicing switch of DMD's penultimate exon compromises muscle fibre maintenance in myotonic dystrophy. *Nat. Commun.* **6**, 7205 (2015).
- Fugier, C. *et al.* Misregulated alternative splicing of BIN1 is associated with T tubule alterations and muscle weakness in myotonic dystrophy. *Nat. Med.* **17**, 720–725 (2011).
- Kimura, T. *et al.* Altered mRNA splicing of the skeletal muscle ryanodine receptor and sarcoplasmic/endoplasmic reticulum Ca $^{2+}$ -ATPase in myotonic dystrophy type 1. *Hum. Mol. Genet.* **14**, 2189–2200 (2005).
- Tang, Z. Z. *et al.* Muscle weakness in myotonic dystrophy associated with misregulated splicing and altered gating of Ca(V)1.1 calcium channel. *Hum. Mol. Genet.* **21**, 1312–1324 (2012).
- Wheeler, T. M., Lueck, J. D., Swanson, M. S., Dirksen, R. T. & Thornton, C. A. Correction of ClC-1 splicing eliminates chloride channelopathy and myotonia in mouse models of myotonic dystrophy. *J. Clin. Invest.* **117**, 3952–3957 (2007).
- Groh, W. J. *et al.* Electrocardiographic abnormalities and sudden death in myotonic dystrophy type 1. *N. Engl. J. Med.* **358**, 2688–2697 (2008).
- Lazarus, A. *et al.* Long-term follow-up of arrhythmias in patients with myotonic dystrophy treated by pacing: a multicenter diagnostic pacemaker study. *J. Am. Coll. Cardiol.* **40**, 1645–1652 (2002).
- Chen, Q. *et al.* Genetic basis and molecular mechanism for idiopathic ventricular fibrillation. *Nature* **392**, 293–296 (1998).
- Tan, H. L. *et al.* A sodium-channel mutation causes isolated cardiac conduction disease. *Nature* **409**, 1043–1047 (2001).
- Schott, J. J. *et al.* Cardiac conduction defects associate with mutations in SCN5A. *Nat. Genet.* **23**, 20–21 (1999).
- Anders, S., Reyes, A. & Huber, W. Detecting differential usage of exons from RNA-seq data. *Genome Res.* **22**, 2008–2017 (2012).
- Katz, Y., Wang, E. T., Aioldi, E. M. & Burge, C. B. Analysis and design of RNA sequencing experiments for identifying isoform regulation. *Nat. Methods* **7**, 1009–1015 (2010).
- Philips, A. V., Timchenko, L. T. & Cooper, T. A. Disruption of splicing regulated by a CUG-binding protein in myotonic dystrophy. *Science* **280**, 737–741 (1998).
- Nakamori, M. *et al.* Splicing biomarkers of disease severity in myotonic dystrophy. *Ann. Neurol.* **74**, 862–872 (2013).
- Kalsotra, A. *et al.* A postnatal switch of CELF and MBNL proteins reprograms alternative splicing in the developing heart. *Proc. Natl Acad. Sci. USA* **105**, 20333–20338 (2008).
- Banks, A. S. *et al.* Deletion of SOCS7 leads to enhanced insulin action and enlarged islets of Langerhans. *J. Clin. Invest.* **115**, 2462–2471 (2005).
- Wani, M. A. *et al.* Familial hypercatabolic hypoproteinemia caused by deficiency of the neonatal Fc receptor, FeRn, due to a mutant beta2-microglobulin gene. *Proc. Natl Acad. Sci. USA* **103**, 5084–5089 (2006).
- Yuan, Y. *et al.* Muscleblind-like 1 interacts with RNA hairpins in splicing target and pathogenic RNAs. *Nucleic Acids Res.* **35**, 5474–5486 (2007).
- Kino, Y. *et al.* Muscleblind protein, MBNL1/EXP, binds specifically to CHHG repeats. *Hum. Mol. Genet.* **13**, 495–507 (2004).
- Goers, E. S., Purcell, J., Voelker, R. B., Gates, D. P. & Berglund, J. A. MBNL1 binds GC motifs embedded in pyrimidines to regulate alternative splicing. *Nucleic Acids Res.* **38**, 2467–2484 (2010).
- Teplova, M. & Patel, D. J. Structural insights into RNA recognition by the alternative-splicing regulator muscleblind-like MBNL1. *Nat. Struct. Mol. Biol.* **15**, 1343–1351 (2008).

33. Marquis, J. *et al.* CUG-BP1/CELF1 requires UGU-rich sequences for high-affinity binding. *Biochem. J.* **400**, 291–301 (2006).
34. Teplova, M., Song, J., Gaw, H. Y., Teplov, A. & Patel, D. J. Structural insights into RNA recognition by the alternate-splicing regulator CUG-binding protein 1. *Structure* **18**, 1364–1377 (2010).
35. Wang, E. T. *et al.* Transcriptome-wide regulation of pre-mRNA splicing and mRNA localization by muscleblind proteins. *Cell* **150**, 710–724 (2012).
36. Goodwin, M. *et al.* MBNL sequestration by toxic RNAs and RNA misprocessing in the myotonic dystrophy brain. *Cell Rep.* **12**, 1159–1168 (2015).
37. Lin, X. *et al.* Failure of MBNL1-dependent post-natal splicing transitions in myotonic dystrophy. *Hum. Mol. Genet.* **15**, 2087–2097 (2006).
38. Kanadia, R. *et al.* Reversal of RNA missplicing and myotonia after muscleblind overexpression in a mouse poly(CUG) model for myotonic dystrophy. *Proc. Natl Acad. Sci. USA* **103**, 11748–11753 (2006).
39. Murphy, L. L. *et al.* Developmentally regulated SCN5A splice variant potentiates dysfunction of a novel mutation associated with severe fetal arrhythmia. *Heart Rhythm* **9**, 590–597 (2012).
40. Walzik, S., Schroeter, A., Benndorf, K. & Zimmer, T. Alternative splicing of the cardiac sodium channel creates multiple variants of mutant T1620K channels. *PLoS ONE* **6**, e19188 (2011).
41. Onkal, R. *et al.* Alternative splicing of Na<sub>v</sub>1.5: an electrophysiological comparison of 'neonatal' and 'adult' isoforms and critical involvement of a lysine residue. *J. Cell. Physiol.* **216**, 716–726 (2008).
42. Wahbi, K. *et al.* Brugada syndrome and abnormal splicing of SCN5A in myotonic dystrophy type 1. *Arch. Cardiovasc. Dis.* **106**, 635–643 (2013).
43. Lee, K. Y. *et al.* Compound loss of muscleblind-like function in myotonic dystrophy. *EMBO Mol. Med.* **5**, 1887–1900 (2013).
44. Dixon, D. M. *et al.* Loss of muscleblind-like 1 results in cardiac pathology and persistence of fetal splice isoforms. *Sci. Rep.* **5**, 9042 (2015).
45. Leoni, A. L. *et al.* Variable Na(v)1.5 protein expression from the wild-type allele correlates with the penetrance of cardiac conduction disease in the Scn5a(+/−) mouse model. *PLoS ONE* **5**, e9298 (2010).
46. Royer, A. *et al.* Mouse model of SCN5A-linked hereditary Lenègre's disease: age-related conduction slowing and myocardial fibrosis. *Circulation* **111**, 1738–1746 (2005).
47. van Veen, T. A. *et al.* Impaired impulse propagation in Scn5a-knockout mice: combined contribution of excitability, connexin expression, and tissue architecture in relation to aging. *Circulation* **112**, 1927–1935 (2005).
48. O'Hara, T., Virág, L., Varró, A. & Rudy, Y. Simulation of the undiseased human cardiac ventricular action potential: model formulation and experimental validation. *PLoS Comput. Biol.* **7**, e1002061 (2011).
49. Tsuji-Wakisaka, K. *et al.* Identification and functional characterization of KCNQ1 mutations around the exon 7-intron 7 junction affecting the splicing process. *Biochim. Biophys. Acta* **1812**, 1452–1459 (2011).
50. Inada, S., Hancox, J. C., Zhang, H. & Boyett, M. R. One-dimensional mathematical model of the atrioventricular node including atrio-nodal, nodal, and nodal-his cells. *Biophys. J.* **97**, 2117–2127 (2009).
51. Probst, V. *et al.* Progressive cardiac conduction defect is the prevailing phenotype in carriers of a Brugada syndrome SCN5A mutation. *J. Cardiovasc. Electrophysiol.* **17**, 270–275 (2006).
52. Zhang, Z. S., Tranquillo, J., Neplioueva, V., Bursac, N. & Grant, A. O. Sodium channel kinetic changes that produce Brugada syndrome or progressive cardiac conduction system disease. *Am. J. Physiol. Heart Circ. Physiol.* **292**, H399–H407 (2007).
53. Wang, D. W., Viswanathan, P. C., Balsler, J. R., George, Jr A. L. & Benson, D. W. Clinical, genetic, and biophysical characterization of SCN5A mutations associated with atrioventricular conduction block. *Circulation* **105**, 341–346 (2002).
54. Rudnik-Schöneborn, S. *et al.* Brugada-like cardiac disease in myotonic dystrophy type 2: report of two unrelated patients. *Eur. J. Neurol.* **18**, 191–194 (2011).
55. Pambrun, T. *et al.* Unmasked Brugada pattern by Ajmaline challenge in patients with myotonic dystrophy type 1. *Ann. Noninvasive Electrocardiol.* **20**, 28–36 (2015).
56. Maury, P. *et al.* Prevalence of type 1 Brugada-ECG pattern following administration of Class 1C drugs in patients with type 1 myotonic dystrophy: myotonic dystrophy as a part of the Brugada syndrome. *Heart Rhythm* **11**, 1721–1727 (2014).
57. Algalarondo, V. *et al.* Abnormal sodium current properties contribute to cardiac electrical and contractile dysfunction in a mouse model of myotonic dystrophy type 1. *Neuromuscul. Disord.* **25**, 308–320 (2014).
58. Yadava, R. S. *et al.* RNA toxicity in myotonic muscular dystrophy induces NKX2-5 expression. *Nat. Genet.* **40**, 61–68 (2008).
59. Koshelev, M., Sarma, S., Price, R. E., Wehrens, X. H. & Cooper, T. A. Heart-specific overexpression of CUGBP1 reproduces functional and molecular abnormalities of myotonic dystrophy type 1. *Hum. Mol. Genet.* **19**, 1066–1075 (2010).
60. Kalsotra, A. *et al.* The Mef2 transcription network is disrupted in myotonic dystrophy heart tissue, dramatically altering miRNA and mRNA expression. *Cell Rep.* **6**, 336–345 (2014).
61. Rau, F. *et al.* Misregulation of miR-1 processing is associated with heart defects in myotonic dystrophy. *Nat. Struct. Mol. Biol.* **18**, 840–845 (2011).
62. Gorog, D. A., Russell, G., Casian, A. & Peters, N. S. A cautionary tale: the risks of flecainide treatment for myotonic dystrophy. *J. Clin. Neuromuscul. Dis.* **7**, 25–28 (2005).
63. Otten, R. F. *et al.* Arrhythmia exacerbation after sodium channel blockade in myotonic dystrophy type 1. *Muscle Nerve* **40**, 901–902 (2009).
64. Groh, W. J., Lowe, M. R., Simmons, Z., Bhakta, D. & Pascuzzi, R. M. Familial clustering of muscular and cardiac involvement in myotonic dystrophy type 1. *Muscle Nerve* **31**, 719–724 (2005).
65. Pambrun, T. *et al.* Myotonic dystrophy type 1 mimics and exacerbates Brugada phenotype induced by Nav1.5 sodium channel loss-of-function mutation. *Heart Rhythm* **11**, 1393–1400 (2014).
66. Childs-DiNey, J. L. *et al.* Induction and reversal of myotonic dystrophy type 1 pre-mRNA splicing defects by small molecules. *Nat. Commun.* **4**, 2044 (2013).
67. Wheeler, T. M. *et al.* Targeting nuclear RNA for *in vivo* correction of myotonic dystrophy. *Nature* **488**, 111–115 (2012).
68. Nakamori, M. *et al.* Aberrantly spliced alpha-dystrobrevin alters alpha-syntrophin binding in myotonic dystrophy type 1. *Neurology* **70**, 677–685 (2008).
69. Schoser, B. G. *et al.* Sudden cardiac death in myotonic dystrophy type 2. *Neurology* **63**, 2402–2404 (2004).
70. Udd, B., Krahe, R., Wallgren-Pettersson, C., Falck, B. & Kalimo, H. Proximal myotonic dystrophy—a family with autosomal dominant muscular dystrophy, cataracts, hearing loss and hypogonadism: heterogeneity of proximal myotonic syndromes? *Neuromuscul. Disord.* **7**, 217–228 (1997).
71. Kim, D. *et al.* TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions. *Genome Biol.* **14**, R36 (2013).
72. Trapnell, C. *et al.* Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. *Nat. Biotechnol.* **28**, 511–515 (2010).
73. Goyenvalle, A. *et al.* Rescue of dystrophic muscle through U7 snRNA-mediated exon skipping. *Science* **306**, 1796–1799 (2004).

## Acknowledgements

We thank Tom Cooper (Baylor College of Medicine, Houston, USA) for the gift of the DT960 and tgCUGBP1 plasmids, Christiane Brantl (CNRS, Nancy, France) for the gift of the pGEX-MBNL1Δ<sup>101</sup> vector, the Research Resource Network Japan and the Myobank of the Institute of Myology for providing human heart tissue samples, Feriel Azibani (UMRS974, Paris, France) and all members of the French DM Network for fruitful discussion. We also deeply thank the IGBMC Microarray and Sequencing platform, the informatics resources of the PRABI, Nathalie Mougenot from the UPMC PECMV-CEF facility, the Fédération de Recherche en Imagerie Multimodalité ([www.bichat.inserm.fr](http://www.bichat.inserm.fr)), the AAV platform of the Centre de Recherche en Myologie, the Penn Vector Core, Gene Therapy Program, University of Pennsylvania (Philadelphia) for providing the pAAV2/9 plasmid (p5E18-VD29), members of the France Génomique program for RNA sequencing and all IGBMC common facilities for assistance. This work was supported by UPMC/INSERM/CNRS/AIM (D.F.); ANR-12-RARE-0005-01 E-RARE 'HEART DM' (N.C.-B. and D.F.); AFM #18833, ERC-2012-Stg #310659 'RNA DISEASES' and ANR-10-LABX-0030-INRT and ANR-10-IDEX-0002-02 (N.C.-B.); Research Grants for Practical Research Project for Rare / Intractable Diseases and for Comprehensive Research on Persons with Disabilities from AMED (H.M., M.P.T.); Grants-in-Aids from MEXT (25136705) (H.I.); Grants-in-Aids from JSPS (25713034 (M.N.) and 15K09314 (M.P.T.)); and Intramural Research Grant of NCNP 26-8 (M.P.T.).

## Author contributions

Experiments were performed by F.F., F.R., Y.K., T.L., E.C., F.D., M.N., H.M., Y.K., N.N., C.S., L.A., A.J., C.T., M.P., S.V. and B.J. Bioinformatic analyses were performed by S.I., T.A., K.N., H.I., M.H., K.I., V.L., A.F., V.N., E.W., C.B. and D.A. Clinical samples and patient data were obtained from A.L.d.M., B.U., J.W.D., D.D., K.W., T.M., H.F. and D.F. Data were collected and analysed by F.F., F.R., T.K., S.I., M.S., D.F., T.Z., M.P.T and N.C.B. The study was designed, coordinated and written by E.W., T.Z., D.F., M.P.T and N.C.B.

## Additional information

**Accession codes:** ExonArray and RNA-Seq data have been deposited in Gene Expression Omnibus (GEO) data base under accession codes GSE67067 and GSE67812, respectively.

**Supplementary Information** accompanies this paper at <http://www.nature.com/naturecommunications>

**Competing financial interests:** The authors declare no competing financial interests.

Reprints and permission information is available online at <http://npg.nature.com/reprintsandpermissions/>

**How to cite this article:** Freyermuth, F. *et al.* Splicing misregulation of SCN5A contributes to cardiac-conduction delay and heart arrhythmia in myotonic dystrophy. *Nat. Commun.* 7:11067 doi: 10.1038/ncomms11067 (2016).



This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>