Delayed conduction and its implications in murine Scn5a+/- hearts: independent and interacting effects of genotype, age, and sex

Kamalan Jeevaratnam · Sui Poh Tee · Yanmin Zhang · Rebecca Rewbury · Laila Guzadhur · Rudolf Duehmke · Andrew A. Grace · Ming Lei · Christopher L.-H. Huang

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Abstract We explored for relationships between SCN5A haploinsufficiency, implicated in clinical arrhythmogenicity, and right ventricular (RV) conduction disorders in Langendorff-perfused, male and female, and young (3 months) and old (>12 month old) Scn5a+/- and wild type (WT) hearts. The investigated conditions of genotype, age, and sex affected latencies but not repolarization time courses of RV monophasic action potentials. This prompted examination of the patterns of RV epicardial activation, its dispersion, and their interrelationships as possible arrhythmogenic mechanisms using a 64-channel, multi-electrode array. Mean ventricular activation times (T*MEAN), spatial dispersions (D*S) between recording channels/cardiac cycle, and maximum activation times (T*MAX) representing the slowest possible conduction in any given heart were all higher in old male Scn5a+/- compared with young male and old female Scn5a+/- and old male WT. Temporal dispersions (D*T) of recording channels were similarly higher in old male Scn5a+/- compared with old male WT. All groupings of D*T, D*S, and T*MAX nevertheless linearly correlated with T*MEAN, with indistinguishable slopes. The variates explored thus influence D*T, D*S, and T*MAX through actions on T*MEAN. These findings in turn correlated with increased levels of fibrosis in young male, young female, and old male Scn5a+/- compared with the corresponding WTs. We thus demonstrate for the first time independent and interacting effects of genotype, age, and sex on epicardial conduction and its dispersions at least partially attributable to fibrotic change, resulting in the greatest effects in old male Scn5a+/- in an absence of alterations in repolarization time courses. This directly implicates altered depolarization in the clinical arrhythmogenicity associated with Scn5a+/-.

Keywords Multi-array · Biophysics · Conduction · Dispersion · Depolarization · Repolarization

Introduction

There has been increasing interest in the possible physiological implications of haploinsufficiencies involving...
voltage-gated Na\(^+\) channels for clinical cardiac electrophysiological disorders. The latter include the Brugada syndrome (BrS) and progressive cardiac conduction disease (PCCD), both of which are associated with arrhythmic and conduction abnormalities [8]. Pathophysiological findings have implicated the right ventricle (RV) particularly the right ventricular outflow tract as the primary location for electrical disorders related to BrS [4]. BrS typically presents as syncope and cardiac arrest typically occurring in the third or fourth decade of life in males rather than females [14]. Thus, men with BrS have a higher prevalence of life-threatening arrhythmias than women corresponding to incidences of 9.5% and 3.8%, respectively [18]. BrS is also the leading cause of death of men under the age of 50 years in endemic regions [2]. Eighty percent of patients in Western populations and 90% in Asian countries showing clinical manifestations of BrS caused by SCN5A mutation are males [22]. The corresponding incidences have not been established for PCCD.

Genetically modified murine systems have been useful in exploring the consequences of SCN5A haploinsufficiencies. They potentially fully replicate the genetic changes associated with arrhythmogenic conditions without requiring pharmacological manipulation. Langendorff-perfused Scn5a\(^+/-\) hearts were more arrhythmic than their wild type (WT) counterparts [14, 23]. In addition, age and sex influenced murine Scn5a\(^+/-\) conduction phenotypes. Scn5a\(^+/-\) mice showed ~20% increases in the durations of their QRS complexes [17]. More recent detailed electrocardiographic (ECG) studies demonstrated independent and interacting effects of age and sex on both sino-atrial and atrioventricular function, consistent with marked conduction alterations particularly in old male Scn5a\(^+/-\) that paralleled clinical arrhythmogenic findings [7].

Electrophysiological changes in BrS have been replicated in arterially perfused canine right ventricular wedges through pharmacological manipulation. Such experiments described localized losses of the AP dome with potential arrhythmogenic effects [1, 5, 26]. RV conduction block has also been implicated in the development of arrhythmias in BrS [9, 12]. A recent human clinical trial on BrS patients attributed their type 1 ECG patterns to local depolarization as opposed to repolarization abnormalities [15]. Murine Scn5a\(^+/-\) mice similarly demonstrated prolonged ECG P wave and PR durations, right axis deviation, and second-degree AV block suggesting conduction deficiencies [14]. Their electrogram durations (EGDs) during bipolar electrogram recordings particularly following premature stimulation were also increased [23]. Both these abnormal patterns of myocardial activation and changes in EGD were attributed to re-entrant substrates resulting from altered (“fractionated”) conduction velocity distributions [19–21, 24].

The present study explores the hypothesis that Scn5a\(^+/-\) genotype, age, and sex alter cardiac conduction. This in turn would alter the dispersions in the resulting myocardial activation times at the tissue level, thereby influencing arrhythmic tendency. The electrophysiological experiments combined monophasic action potential and multi-electrode array recordings. They studied action potential properties, followed by detailed analyses of epicardial activation times and their dispersion for the first time, for subsequent comparison with morphological studies exploring for possible roles of tissue fibrosis in such changes. The observed variations in activation properties with genotype, age, and sex directly correlated with the corresponding fibrosis, suggesting a mechanism for such conduction changes. These in turn directly accounted for the previously reported clinical differences in cardiac arrhythmogenicity.

Methods

Experimental groups

Mice were housed in an animal facility at 21°C with 12-h light/dark cycles. Animals were fed sterile chow (RM3 Maintenance Diet, SDS, Witham, Essex, UK) and had free access to water. All procedures complied with the UK Home Office regulations (Animal (Scientific Procedures) Act 1986). In order to develop a comprehensive risk assessment model of the two types of mice studied, a total of 23 WT and 21 Scn5a\(^+/-\) knockout mice were used. All mice used for this study were derived from their respective 129/sv background strains to avoid any possible strain related variation in the study. The mice were divided into eight groups. Group 1 was WT male aged 3 months (\(n=7\)), group 2 was WT male aged >12 months (\(n=5\)), group 3 was WT female aged 3 months (\(n=6\)), group 4 was WT female aged >12 months (\(n=5\)), group 5 was Scn5a\(^+/-\) male aged 3 months (\(n=6\)), group 6 was Scn5a\(^+/-\) male aged >12 months (\(n=5\)), group 7 was Scn5a\(^+/-\) female aged 3 months (\(n=5\)), and group 8 was Scn5a\(^+/-\) female aged >12 months (\(n=5\)). Experiments used a Langendorff-perfused preparation adapted for the murine heart, as described previously [6, 23]. All assessments were performed fully blinded, and codes were broken only after all measurements had been performed.

Multi-electrode array recordings

A custom-made electrode array consisting of 64 separate electrodes (Teflon-coated silver wires; 0.125 mm diameter; Science Products) in an 8×8 configuration with an inter-electrode distance of 0.55 mm was used. Unipolar recordings were performed by placing the multi-electrode array on the right epicardium with the reference electrode, acting as
the indifferent pole at the base of the heart. The 64 recording electrodes were connected through shielded wires to two 32-channel amplifiers (SCXI-1102 C, National Instruments Corporation (U.K.) Ltd, Newbury, UK). Sampling frequency for each signal was set at 1 kHz. The signals were continuously stored on disk and displayed on screen using a custom-developed program, written in Labview 7.0 (National Instruments Corporation (U.K.) Ltd, Newbury, UK).

For the off-line analysis, signals were displayed on screen in sets of eight to 16 electrograms. The activation time was determined as the point of maximal negative slope and marked with a cursor. After marking all significant waveforms in all leads, the activation times were then displayed in a grid representing the layout of the original recording array. All activation times, in milliseconds, were related to the timing of the first detected waveform. Activation maps were drawn using Microsoft Office Excel 2007 (Microsoft Corporation, Silicon Valley, USA).

Monophasic action potential recordings

A bipolar platinum stimulating electrode (1-mm interpolar spacing) was placed on the interventricular septum, and a monophasic action potential (MAP) recording electrode (Hugo Sachs, Harvard Apparatus) was placed on the corresponding right epicardium. During this time, MAPs showed stable baselines, rapid upstroke phases that reached consistent amplitudes, and smooth repolarization phases, all of which are known to be characteristic of high-fidelity MAP recordings. MAPs were amplified, band-pass filtered (0.5 Hz–1 kHz: Neurolog NL104 Amplifer, NL125/6 Filter; Digitimer, Welwyn Garden City, Herts, UK) and digitized at a sampling frequency of 5 kHz (micro1401, Cambridge Electronic Design, Cambridge, UK). Hearts were paced at a cycle length of 125 ms. Analysis of MAPs was performed using Spike II software (Cambridge Electronic Design).

Quantification of cardiac fibrosis

Isolated hearts were flushed with Krebs buffer and perfused with 4% buffered formalin for 5 min before being immersed in formalin overnight. After fixation, gross transverse sections were cut from base to apex, and the hearts were subjected to routine tissue processing and paraffin embedding. Paraffin sections of 7 μm thickness were then cut and stained with Sirius red stain (Sigma-Aldrich, Dorset, UK). All sections were subsequently viewed, magnified, and digitally acquired using the Nano Zoomer 2.0 Digital Pathology system (Hamamatsu, Hertfordshire, UK). For the quantification of fibrosis, two randomly selected photomicrographs with a minimum distance of 14 μm from each other were taken for every heart.

Following a × 2 magnification, a custom-made 17 cm × 23 cm morphometric grid, consisting of square boxes of dimension 1 cm × 1 cm, corresponding to a 0.26 mm × 0.26 mm area of tissue, was then superimposed on each photomicrograph. Only squares occupied either completely or partially by cardiac tissues were counted. Similarly, squares containing fibrotic changes were then counted, and this was expressed as a percentage of the total cardiac tissue area. An average percentage of fibrotic changes for every heart taken from the two photomicrographs were obtained and then accumulated for every group to provide the mean percentage of fibrotic tissue.

Statistical analysis

Comparisons where appropriate were performed by analysis of variance or t test with SPSS software (SPSS UK, Woking, Surrey, UK). Regression coefficients were calculated by

\[ t = r\sqrt{(n - 2)/\sqrt{(1 - r^2)}} \]

with \( n - 2 \) degrees of freedom [16]. Statistical significance was assumed at \( P < 0.05, P < 0.01, \) or \( P < 0.001 \) where applicable.

Results

MAP assessments of action potential repolarization and latencies

The initial MAP studies observed alterations in action potential latencies that prompted the more detailed studies of such latencies that followed. Thus, Fig. 1a–b shows typical action potential waveforms including their latencies from each of the experimental groups. Figure 1i, j quantifies the resulting measures of action potential duration (APD), obtained at 30%–90% full repolarization, in the eight experimental groups. APD\(_{30}\), APD\(_{50}\), APD\(_{70}\), and APD\(_{90}\) values (Fig. 1i, j) showed no statistical differences between experimental groups in tests for either independent or interacting stratifications by genotype, age, or sex.

In contrast, significant differences were observed concerning the MAP latencies (Table 1). Thus, the initial stratifications simply by genotype demonstrated that Scn5a\(^{+/-}\) hearts had significantly longer (\( P < 0.001 \)) latencies compared with WT hearts. Stratifications by genotype and age then demonstrated that there were no latency differences between young and old WT but longer latencies (\( P = 0.002 \)) in old Scn5a\(^{+/-}\) when compared with young Scn5a\(^{+/-}\). Both young and old Scn5a\(^{+/-}\) showed longer (\( P = 0.029; P = 0.001 \)) latencies compared with young and old WT, respectively. A third stratification assessed for independent effects of genotype and sex. This confirmed an absence of latency differences between male and female WT. However, latency
was longer ($P=0.032$) in male Scn5a$^{+/−}$ compared with female Scn5a$^{+/−}$. Male Scn5a$^{+/−}$ showed longer ($P<0.001$) latencies compared with male WT; differences were absent in the corresponding females.

The final stratification examined for the interacting effects of genotype, age, and sex. Both old male Scn5a$^{+/−}$ and old female Scn5a$^{+/−}$ hearts showed longer ($P=0.044$; $P=0.046$) latencies compared with the young male Scn5a$^{+/−}$ and young female Scn5a$^{+/−}$ hearts, respectively. Such differences were absent in the WT population. Furthermore, young male Scn5a$^{+/−}$ and old male Scn5a$^{+/−}$ had longer ($P=0.006$; $P=0.004$) latencies compared with young male WT and old male WT, respectively. All other comparisons were not significantly different. Together, these findings suggest that MAP latencies were influenced by independent and interacting effects of genotype, age, and sex. These findings prompted the more detailed multi-array studies that follow.

Multi-array analysis for conduction and its dispersion

Figure 2 shows representative waveforms obtained using a 64-channel multi-electrode array placed on the right ventricles of Langendorff-perfused, intrinsically beating hearts. The records were obtained from two rows, marked 

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**Fig. 1** Mean repolarization durations represented by values of APD$_{30}$, APD$_{50}$, APD$_{70}$, and APD$_{90}$ for WT and Scn5a$^{−/−}$ hearts sorted by age and sex. APDs were obtained from Langendorff-perfused hearts isolated from healthy mice. Representative MAPs waveform recordings are shown for all the eight experimental groups (a–h). Durations of APDs for the stratification by genotype only, genotype and age, and genotype and sex are not shown. The full stratification of both WT and Scn5a$^{−/−}$ mice by both age and sex is represented by i and j, respectively. All APD values were not significantly different irrespective of genotype, age, or sex through all stratification levels.
Table 1  Latency (ms) in monophasic action potential activation

| Animal       | Samples | Average ± SE |
|--------------|---------|--------------|
| WT (total)   | 27      | 15.66±0.77 a |
| WT young     | 14      | 14.79±0.54 b |
| WT old       | 13      | 16.60±1.49 c |
| WT male      | 14      | 16.36±1.09 d |
| WT female    | 13      | 14.91±1.01   |
| WT young male| 6       | 15.71±0.64 e |
| WT young female| 8     | 14.10±0.76   |
| WT old male  | 8       | 16.84±1.88 f |
| WT old female| 5       | 16.22±2.67   |
| Scn5a +/- (total) | 21    | 21.20±1.11 a |
| Scn5a +/- young | 11    | 17.85±1.36 b, g |
| Scn5a +/- old  | 10     | 24.25±1.10 c, g |
| Scn5a +/- male | 11     | 23.41±1.17 d, h |
| Scn5a +/- female | 10    | 18.77±1.67 h |
| Scn5a +/- young male | 4   | 20.10±1.23 e, i |
| Scn5a +/- young female | 6   | 16.15±1.88 j |
| Scn5a +/- old male | 7 | 25.14±1.35 f, i |
| Scn5a +/- old female | 4 | 22.70±1.88 j |

Means with similar lowercase letters differ significantly from each other at P<0.05

1–8 and 9–16, respectively, in the array from (a) old male WT, (b) old male Scn5a +/-, (c) old female WT, and (d) old female Scn5a +/- hearts. Most of the recording elements indicated a presence of electrical activity reflecting effective contact with the apposed tissue. This took the form of similar biphasic waveforms with different latencies. It was therefore possible to characterize each trace by a single activation time during offline analysis for all 64 channels. Such latencies were measured by locating a cursor to the point of maximal negative slope in the trace. This was repeated in five consecutive beats in all hearts that were studied. The resulting activation times obtained were then displayed in a matrix representing the layout of the original recording array through the use of a false color code in which the time of earliest to latest activation were represented by red and a gradual color change to green.

Figure 3 exemplifies the resulting activation maps in five successive cardiac cycles from typical young (a, b, e, and f) and old (c, d, g, and h), male (a, e, c, and g), female (b, f, d, and h), WT (a, b, c, and d), and Scn5a +/- hearts (e, f, g, and h). Most of the activation maps obtained from the WT hearts showed a general pattern of sequential activation in which one region of the matrix appeared to be activated first, and this was then followed by a successive spread of activation through the remainder of the area from which recordings were made. This would be compatible with either a spread of excitation within the plane of the epicardium or an orderly arrival of activation from the endocardium to the epicardium. However the Scn5a +/- hearts often did not show this orderly pattern and often exhibited a more random excitation of different channels. Rather than describing our results solely in terms of a sequential activation within the plane of the epicardium, we therefore adopted a more empirical description quantifying the activation times, their spatial and temporal dispersions, and their maximum values.

Figure 3e–h thus indicates that the Scn5a +/- hearts showed a greater frequency of areas reflecting late activation times in comparison with WT. This trend was more prominent in males compared with females and was particularly marked in the old male Scn5a +/- (Fig. 3g) hearts whether compared with remaining groups of either Scn5a +/- or WT hearts. In contrast, among the WT hearts...
The old female WT (Fig. 3d) appeared to have the most delayed activation time compared with the other WT groups, although this distinction was not as clear as for the Scn5a\(^{+/−}\). These findings suggest a greater propensity to delayed activation times in Scn5a\(^{+/−}\) in comparison with WT. They further suggest that the interacting influences of age, sex, and genotype, with older male Scn5a\(^{+/−}\) showing particularly longer delays. These observations were subsequently quantified through determinations of the following parameters. All these involved a collection of multi-array data over five consecutive cardiac cycles, i.e., over a time span of 1 s.

Mean activation times

Firstly, a mean activation time, \(T_{\text{MEAN}}\), was used to describe the overall activation time within any given heart. This was obtained from the activation times in all channels averaged over five beats. These values were then used to obtain average values, \(T^*_{\text{MEAN}}\), for each group of hearts investigated. Table 2 summarizes the resulting \(T^*_{\text{MEAN}}\) values, expressed as means ± standard errors of the mean from the different levels of stratification of the experimental population. Such successive stratifications demonstrated significant independent and interacting effects of genotype, age, and sex.

Thus, initial comparison of the population as a whole by genotype prior to the subsequent stratifications involving age and sex demonstrated that \(T^*_{\text{MEAN}}\) was significantly higher (\(P=0.046\)) in the Scn5a\(^{+/−}\) hearts when compared with WT hearts. Stratification of the dataset by both genotype and age yielded four groups: young WT, old WT, young Scn5a\(^{+/−}\) and old Scn5a\(^{+/−}\). The resulting comparisons showed that there were no significant differences in \(T^*_{\text{MEAN}}\), between any of these groups. Stratifications by both genotype and sex gave contrasting groups consisting of male WT, female WT, male Scn5a\(^{+/−}\) and female Scn5a\(^{+/−}\). Values of \(T^*_{\text{MEAN}}\) between male and female WT were then indistinguishable. However, they were significantly higher (\(P=0.042\)) for male Scn5a\(^{+/−}\) compared with females Scn5a\(^{+/−}\). They were also significantly higher (\(P=0.042\)) for male Scn5a\(^{+/−}\) compared
Secondly, a temporal dispersion of activation times, $D_T$, was derived to describe the variation in activation time at a given channel. This was determined from five successive cardiac cycles. Readings from each of the 64 individual channels in the array were then averaged. This yielded the temporal dispersion $D_T$ for the heart under study. These values were then used to obtain average values, $D^*_T$, for each group of hearts investigated.

Table 3 summarizes the resulting $D^*_T$ values from the different levels of stratification. Stratification by genotype alone showed that $D^*_T$ was higher in the Scn5a$^{+/−}$ hearts compared with the WT hearts ($P<0.048$). However, stratifications of the study population by both genotype and age demonstrated that $D^*_T$ was indistinguishable whether between young and old animals within a given genotype or between WT and Scn5a$^{+/−}$ within a given age group. Stratifications by genotype and sex similarly revealed that values of $D^*_T$ showed no significant differences between male WT and female WT and between male Scn5a$^{+/−}$ and female Scn5a$^{+/−}$. However, values of $D^*_T$ were significantly higher ($P=0.028$) in the male Scn5a$^{+/−}$ mice compared with the male WT mice. The final level of stratification separating interacting effects between genotype, age, and sex showed that there were no statistical differences in $D^*_T$ between the four groups of WT hearts. These similarities were also observed within the groups of Scn5a$^{+/−}$ hearts. However, $D^*_T$ was significantly higher ($P=0.046$) in old male Scn5a$^{+/−}$ compared with old male WT.

| Table 2 | $T^*_\text{MEAN}$ (ms) values in right ventricular murine hearts |
|---------|-------------------------------------------------------------|
| Animal  | Samples | Average ± SE |
| WT (total) | 23 | 2.45±0.25 a |
| WT young | 13 | 2.29±0.20 |
| WT old | 10 | 2.65±0.52 |
| WT male | 12 | 2.22±0.37 b |
| WT female | 11 | 2.71±0.33 |
| WT young male | 7 | 2.30±0.28 |
| WT young female | 6 | 2.29±0.32 |
| WT old male | 5 | 2.09±0.85 c |
| WT old female | 5 | 3.21±0.57 |
| Scn5a$^{+/−}$ (total) | 21 | 3.37±0.38 a |
| Scn5a$^{+/−}$ young | 11 | 2.92±0.45 |
| Scn5a$^{+/−}$ old | 10 | 3.87±0.62 |
| Scn5a$^{+/−}$ male | 11 | 4.08±0.63 b, d |
| Scn5a$^{+/−}$ female | 10 | 2.59±0.24 d |
| Scn5a$^{+/−}$ young male | 6 | 2.84±0.81 e |
| Scn5a$^{+/−}$ young female | 5 | 3.02±0.31 |
| Scn5a$^{+/−}$ old male | 5 | 5.58±0.42 c, c, f |
| Scn5a$^{+/−}$ old female | 5 | 2.17±0.27 f |

Means with similar lowercase letters differ significantly from each other at $P<0.05$

With the male WT hearts. However, such a distinction was not observed between female Scn5a$^{+/−}$ and female WT.

The final level of stratification additionally resolved interacting effects between the variables of genotype, age, and sex. This gave the groups: young male WT, old male WT, young female WT, old female WT, young male Scn5a$^{+/−}$, old male Scn5a$^{+/−}$, young female Scn5a$^{+/−}$, and old female Scn5a$^{+/−}$. This showed that regardless of age and sex, there were no statistical differences observed in $T^*_\text{MEAN}$ among the four groups of WT hearts. However, $T^*_\text{MEAN}$ showed a more diverse pattern of statistical significances within the Scn5a$^{+/−}$ group. $T^*_\text{MEAN}$ in the old male Scn5a$^{+/−}$ was significantly higher whether compared with the old female Scn5a$^{+/−}$ ($P=0.009$), the young male Scn5a$^{+/−}$ ($P=0.021$), as well as the old male WT ($P=0.028$). Thus $T^*_\text{MEAN}$ was greater in the old male Scn5a$^{+/−}$ hearts than in any of the remaining groups. In contrast, it was statistically identical between young male Scn5a$^{+/−}$ and young female Scn5a$^{+/−}$, old female Scn5a$^{+/−}$, and young female Scn5a$^{+/−}$ as well as between old female Scn5a$^{+/−}$ and old female WT.

Temporal dispersions of activation times

Table 3

| Animal | Samples | Average ± SE |
|--------|---------|--------------|
| WT (total) | 23 | 0.88±0.11 a |
| WT young | 13 | 0.79±0.10 |
| WT old | 10 | 1.01±0.21 |
| WT male | 12 | 0.78±0.12 b |
| WT female | 11 | 1.00±0.19 |
| WT young male | 7 | 0.82±0.11 |
| WT young female | 6 | 0.75±0.20 |
| WT old male | 5 | 0.72±0.24 c |
| WT old female | 5 | 1.29±0.31 |
| Scn5a$^{+/−}$ (total) | 21 | 1.24±0.14 a |
| Scn5a$^{+/−}$ young | 11 | 1.08±0.15 |
| Scn5a$^{+/−}$ old | 10 | 1.43±0.25 |
| Scn5a$^{+/−}$ male | 11 | 1.41±0.25 b |
| Scn5a$^{+/−}$ female | 10 | 1.07±0.12 |
| Scn5a$^{+/−}$ young male | 6 | 1.07±0.27 |
| Scn5a$^{+/−}$ young female | 5 | 1.09±0.15 |
| Scn5a$^{+/−}$ old male | 5 | 1.82±0.40 c |
| Scn5a$^{+/−}$ old female | 5 | 1.03±0.20 |

Means with similar lowercase letters differ significantly from each other at $P<0.05$
Spatial dispersions of activation times

Thirdly a spatial dispersion, $D_s$, was obtained to describe the dispersion of activation times between recording channels in a given cardiac cycle averaged through five successive cycles. Activation times were obtained for each of the 64 channels in any given cycle. The standard deviation of these was then calculated. Averaged values of this standard deviation were then obtained from five consecutive cardiac cycles to derive $D_s$. These values were then used to obtain average values, $D^*_s$, for each group of hearts investigated.

Table 4 summarizes such $D^*_s$ values following stratification into WT and Scn5a$^{+/−}$ hearts and then demonstrated no differences in $D^*_s$. Further stratification of the dataset by genotype and age demonstrated no significant differences in $D^*_s$ regardless of genotype or age. Similarly, stratifications by genotype and sex showed no significant differences in $D^*_s$ between any pair of experimental groups. Nevertheless, the final level of stratification resolved interacting effects between genotype, age, and sex. There were then no statistical differences observed in $D^*_s$ between the four groups of WT hearts. However, $D^*_s$ appeared to show a more diverse pattern of statistical significances within the Scn5a$^{+/−}$ group. Thus, it was significantly higher in the old male Scn5a$^{+/−}$ compared with the young male Scn5a$^{+/−}$ ($P=0.009$), as well as between old female Scn5a$^{+/−}$. Thus, $D^*_s$ was greater in the old male Scn5a$^{+/−}$ hearts than in any of the remaining groups. In contrast, $D^*_s$ was indistinguishable between young male Scn5a$^{+/−}$ and young female Scn5a$^{+/−}$, old female Scn5a$^{+/−}$ and young female Scn5a$^{+/−}$, as well as between old female Scn5a$^{+/−}$ and old female WT.

Maximum activation times

Finally, a maximum activation time, $T_{MAX}$, provided a simple measure of the slowest possible conduction that could take place in any given heart. It was determined by measuring the activation time from the channel showing the greatest delay in electrical activation through all five cardiac cycles from which data were collected. Such data from all the hearts in any given group were used to obtain an average, $T^*_{MAX}$.

Table 5 summarizes these $T^*_{MAX}$ values, obtained from the different levels of stratification of the experimental groups. Hearts were also sorted by individual $T_{MAX}$ values falling in the respective ranges of 0–3, 4–6, 7–10, and >10 ms, in order to emphasize groups showing either the greatest or smallest extremes in such values. These successive stratifications demonstrated significant independent and interacting effects of genotype, age, and sex. Thus, even the simple initial, genotypic, comparison of the population as a whole demonstrated significantly higher ($P=0.002$) $T^*_{MAX}$ values in the Scn5a$^{+/−}$ compared with the WT. Scn5a$^{+/−}$ and WT correspondingly showed distributions biased toward greater and smaller $T^*_{MAX}$ values, respectively.

Further stratification of the dataset by both genotype and age demonstrated different effects of age within each genotype. Thus, there were statistically indistinguishable and similarly distributed $T^*_{MAX}$ values in old WT and young WT hearts. However, such $T^*_{MAX}$ values were significantly greater ($P=0.031$) in old Scn5a$^{+/−}$ than young Scn5a$^{+/−}$, reflecting distributions skewed in old Scn5a$^{+/−}$ towards higher values, >10 ms, of $T_{MAX}$. In contrast, analysis of the effects of genotype within each age group demonstrated significantly higher ($P=0.046$) $T^*_{MAX}$ values in the young Scn5a$^{+/−}$ hearts compared with the young WT. Similarly, $T^*_{MAX}$ values were significantly higher ($P=0.016$) in the old Scn5a$^{+/−}$ compared with old WT.

Stratifications by both genotype and sex revealed that there were no differences attributable to sex within each genotype. Thus, $T^*_{MAX}$ values were similar both between male and female WT and between male and female Scn5a$^{+/−}$. However, comparisons of the effects of genotype within each sex group showed significantly higher ($P=0.007$) $T^*_{MAX}$ values in the male Scn5a$^{+/−}$ than the male WT. In contrast, $T^*_{MAX}$ values were statistically indistinguishable between female WT and female Scn5a$^{+/−}$.

Table 4 $D^*_s$ (ms) values in right ventricular murine hearts

| Animal                     | Samples | Average ± SE |
|----------------------------|---------|--------------|
| WT (total)                 | 23      | 1.61±0.18    |
| WT young                   | 13      | 1.51±0.21    |
| WT old                     | 10      | 1.74±0.32    |
| WT male                    | 12      | 1.55±0.29    |
| WT female                  | 11      | 1.68±0.21    |
| WT young male              | 7       | 1.65±0.31    |
| WT young female            | 6       | 1.35±0.28    |
| WT old male                | 5       | 1.41±0.59 a  |
| WT old female              | 5       | 2.07±0.22    |
| Scn5a$^{+/−}$ (total)      | 21      | 1.99±0.24    |
| Scn5a$^{+/−}$ young        | 11      | 1.57±0.25    |
| Scn5a$^{+/−}$ old          | 10      | 2.46±0.36    |
| Scn5a$^{+/−}$ male         | 11      | 2.40±0.40    |
| Scn5a$^{+/−}$ female       | 10      | 1.55±0.14    |
| Scn5a$^{+/−}$ young male   | 6       | 1.52±0.47 b  |
| Scn5a$^{+/−}$ young female | 5       | 1.62±0.13    |
| Scn5a$^{+/−}$ old male     | 5       | 3.44±0.24 a, b, c |
| Scn5a$^{+/−}$ old female   | 5       | 1.48±0.27 c  |

Means with similar lowercase letters differ significantly from each other at $P<0.05$
marked effects upon all four WT groups. In contrast, age and sex exerted ranges of frequencies of hearts with different compared with old female Scn5a old male groups of animals. The most striking finding was that the means with similar lowercase letters differ significantly from each other at P<0.05.

Means with similar lowercase letters differ significantly from each other at P<0.05.

The final level of stratification additionally resolved interacting effects between the variables of genotype, age and sex. Firstly, it considered the effects of age and sex within any given genotypic group. This importantly demonstrated that age and sex exerted no effects upon *T*<sub>MAX</sub> within all four WT groups of animals. Thus, *T*<sub>MAX</sub> values were statistically indistinguishable between all four WT groups. In contrast, age and sex exerted marked effects upon *T*<sub>MAX</sub> among the four *Scn5a<sup>−/−</sup>* groups of animals. The most striking finding was that the old male *Scn5a<sup>−/−</sup>* showed the highest *T*<sub>MAX</sub> compared with all the other groups. An analysis of differences attributable to age in sex-matched *Scn5a<sup>−/−</sup>* showed that *T*<sub>MAX</sub> values were significantly higher (P=0.006) in the old male *Scn5a<sup>−/−</sup>* than in the young male *Scn5a<sup>−/−</sup>*. However, such values were similar in the old female *Scn5a<sup>−/−</sup>* compared with the young female *Scn5a<sup>−/−</sup>*. Analysis of differences attributable to sex in age-matched *Scn5a<sup>−/−</sup>* showed that *T*<sub>MAX</sub> values were significantly higher (P=0.008) in the old male *Scn5a<sup>−/−</sup>* with old female *Scn5a<sup>−/−</sup>*. This was in contrast to indistinguishable *T*<sub>MAX</sub> values in the young male *Scn5a<sup>−/−</sup>* and young female *Scn5a<sup>−/−</sup>*.

Secondly, it considered the effects of genotype on animals matched by both age and sex. This demonstrated that *T*<sub>MAX</sub> values were significantly higher (P=0.034) in old male *Scn5a<sup>−/−</sup>* than old male WT. This finding was also reflected in the distributions. Thus, all the *Scn5a<sup>−/−</sup>* old male hearts had a *T*<sub>MAX</sub> value of >10 ms in comparison with the rest of the group. In contrast, there were no other significant differences in *T*<sub>MAX</sub> values among the remaining groups.

Correlations between activation times and measures of their dispersion

The above results demonstrate significant interacting effects of age, sex, and genotype upon measures of activation times, *T*<sub>MEAN</sub>, its dispersions, as represented by *D*<sub>T</sub>, *D*<sub>S</sub>, and *T*<sub>MAX</sub>. Thus, in the group as a whole, genotype exerted significant effects upon *T*<sub>MEAN</sub>, *D*<sub>T</sub>, and *T*<sub>MAX</sub>. Stratification by genotype and age influenced only the *T*<sub>MAX</sub> values and that by genotype and sex influenced *T*<sub>MEAN</sub>, *D*<sub>T</sub>, and *T*<sub>MAX</sub>. The final stratifications revealed effects on all four variables. All significant changes tended to occur in the same direction, consistent with causal relationships between them. Furthermore, significant changes in *D*<sub>T</sub> and *D*<sub>S</sub> were always accompanied by corresponding changes in *T*<sub>MEAN</sub>, but the reverse was not always the case. Together, these findings suggest a trend toward increased dispersions of activation with increasing activation time, establishing the latter variant as a major determinant of such dispersion. These features prompted us to explore for quantitative relationships between the measures of dispersion with their corresponding *T*<sub>MEAN</sub>.

Firstly, we accordingly compared *D*<sub>T</sub>, *D*<sub>S</sub>, and *T*<sub>MAX</sub> values from each individual heart against their corresponding *T*<sub>MEAN</sub>. Figure 4 exemplifies the resulting

| Animal               | Samples | Average ± SE | Intervals |
|----------------------|---------|--------------|-----------|
|                      |         |              | 0–3 ms    | 4–6 ms  | 7–10 ms | >10 ms |
| WT (total)           | 23      | 5.87±0.65 a  | 8         | 6       | 8       | 1      |
| WT young             | 13      | 5.31±0.73 b  | 5         | 4       | 4       | 0      |
| WT old               | 10      | 6.60±1.17 c  | 3         | 2       | 4       | 1      |
| WT male              | 12      | 5.33±1.03 d  | 5         | 4       | 2       | 1      |
| WT female            | 11      | 6.46±0.79    | 3         | 2       | 6       | 0      |
| WT young male        | 7       | 5.29±0.99    | 2         | 3       | 2       | 0      |
| WT young female      | 6       | 5.33±1.17    | 3         | 1       | 2       | 0      |
| WT old male          | 5       | 5.40±2.21    | 3         | 1       | 0       | 1      |
| WT old female        | 5       | 7.80±0.74    | 0         | 1       | 4       | 0      |
| *Scn5a<sup>−/−</sup>* (total) | 21  | 9.29±0.86 a  | 0         | 6       | 9       | 6      |
| *Scn5a<sup>−/−</sup>* young | 11        | 7.46±0.62 b, f | 0        | 4       | 6       | 1      |
| *Scn5a<sup>−/−</sup>* old | 10       | 11.30±1.44 c, f | 0     | 2       | 3       | 5      |
| *Scn5a<sup>−/−</sup>* male | 11      | 10.46±1.48 d | 0         | 3       | 2       | 6      |
| *Scn5a<sup>−/−</sup>* female | 10      | 8.00±0.62    | 0         | 3       | 7       | 0      |
| *Scn5a<sup>−/−</sup>* young male | 6         | 6.83±1.01 g  | 0         | 3       | 2       | 1      |
| *Scn5a<sup>−/−</sup>* young female | 5        | 8.20±0.58    | 0         | 1       | 4       | 0      |
| *Scn5a<sup>−/−</sup>* old male | 5       | 14.80±1.36 e, g, h | 0    | 0       | 0       | 5      |
| *Scn5a<sup>−/−</sup>* old female | 5       | 7.80±1.16 h  | 0         | 2       | 3       | 0      |
plots of $D^*_T$ (a), $D^*_S$ (b), and $T^*_MAX$ (c) against $T^*_MEAN$ when these were sorted by the WT (i) and Scn5a$^{+/-}$ (ii) genotypes. The analysis then went on to explore such correlations when the experimental groups were instead sorted by age and sex. In all six resulting cases, these demonstrated significant correlations between the measures of dispersion and the mean activation time. Thus, Table 6 summarizes the results of correlations of the $n$ available values of $D^*_T$, $D^*_S$, and $T^*_MAX$ with $T^*_MEAN$ following sorting of the data by genotype (WT, Scn5a$^{+/-}$), age (young and old), and sex (male and female). All the regression coefficients, $r^2$, demonstrated significant correlations with $T^*_MEAN$.

Secondly, having established that $T^*_MEAN$ significantly influences $D^*_T$, $D^*_S$, and $T^*_MAX$ in all groups analyzed, we examined if such influences may vary with genotype, age,
and sex. Values of the slope for each regression line, $m$, were obtained and analyzed. As demonstrated by Table 6, $m$ values were statistically indistinguishable between WT and Scn5a+−, between young and old and between male and female. Thus, in contrast to activation time, each of these factors (i.e., genotype, age, and sex) is unlikely to make major determinants by themselves to the degree of dispersion. Thus, in the present study, the major determinant for dispersion appears to be activation time.

Quantitative evaluation of cardiac fibrosis

The present study quantitatively demonstrated major differences in electrophysiological conduction characteristics as a result of the interacting effects of age, sex, and genotype for the first time. These findings agreed with recent qualitative histological evidence that the Scn5a+− mutation interacts with age in producing microscopic myocardial fibrotic changes [10, 17]. The present experiments therefore proceeded to make quantitative histological explorations for effects of genotype, age, and sex upon such fibrotic change. It was then possible to assess the extent to which these might correlate with the corresponding conduction properties.

Figure 5 exemplifies typical histological sections from young (a, b, e, and f) and old (c, d, g, and h), male and (a, e, c, and g) female (b, f, d, and h), and WT (a, b, c, and d) and Scn5a−− hearts (e, f, g, and h). Table 7 summarizes the results of morphometric assessments of the percentage of fibrotic tissue using a stratification scheme for the experimental groups identical to that adopted for the electrophysiological results. The initial stratification by genotype demonstrated a significantly ($P<0.001$) greater proportion of fibrotic tissue in the Scn5a+− than the WT hearts. The second level of stratification by genotype and age demonstrated more fibrosis in old WT and old Scn5a+− hearts ($P<0.001$ and $P=0.013$, respectively) than their young counterparts within the same genotypic groups. Comparisons across genotype similarly demonstrated more fibrosis in the young and old Scn5a+− compared with the young and old WT ($P<0.001$ and $P=0.002$ respectively).

The third level of stratification by genotype and sex additionally demonstrated effects of sex on levels of fibrosis but only in the Scn5a+− hearts. There was more ($P=0.013$) fibrosis in male compared with female Scn5a+−, but no such differences between male and female WT. Comparisons across genotype revealed greater fibrosis ($P<0.001$) in the male Scn5a+− compared with male WT. Such differences were absent between female Scn5a+− and WT. The final level of stratification by genotype, age, and sex similarly corresponded closely to the electrophysiological results. Thus, within genotypic groups, there was more fibrosis in the old male Scn5a−− than either the young male Scn5a+− or the old female Scn5a+− ($P<0.001$ and $P<0.001$ respectively). There was more fibrosis in the old male WT than the young male WT ($P=0.006$) but not the old female WT ($P>0.05$). Comparisons across genotypic group demonstrated more fibrosis in young male Scn5a+−, young female Scn5a+− and old male Scn5a+− compared with the young male WT, young female WT, and old male WT ($P=0.001$, $P=0.002$, and $P<0.001$, respectively). Such differences were not observed between the old female Scn5a+− hearts and old female WT hearts.

Table 6 Results of linear correlation analysis of $D^*_{t}, D^*_{s},$ and $T^*_{\text{MAX}}$ against $T^*_{\text{MEAN}}$ showing values of correlation coefficient, $r^2$, $t$ values of its significance and values of slope, $m$

| Samples | df | $D^*_{t}$ | | | $D^*_{s}$ | | | $T^*_{\text{MAX}}$ | | |
|---------|----|----------|---|---|----------|---|---|----------|---|---|
|         | $r^2$ | $t$ | $m$ | | | $r^2$ | $t$ | $m$ | | |
| WT      | 23  | 21     | 0.36 | 3.45$^a$ | 0.26±0.08 | 0.83 | 10.28$^b$ | 0.70±0.12 | 0.85 | 10.96$^b$ | 1.97±0.46 |
| Scn5a+− | 21  | 19     | 0.48 | 4.16$^b$ | 0.28±0.08 | 0.87 | 11.24$^b$ | 0.45±0.11 | 0.59 | 5.21$^b$ | 2.71±0.30 |
| Young   | 24  | 22     | 0.49 | 4.64$^b$ | 0.21±0.09 | 0.74 | 7.95$^b$ | 0.63±0.16 | 0.54 | 5.11$^b$ | 0.92±0.45 |
| Old     | 20  | 18     | 0.43 | 3.65$^a$ | 0.34±0.05 | 0.89 | 12.24$^b$ | 0.52±0.10 | 0.81 | 8.67$^b$ | 2.76±0.26 |
| Male    | 23  | 21     | 0.57 | 5.32$^b$ | 0.24±0.07 | 0.89 | 13.1$^b$ | 0.63±0.12 | 0.87 | 11.85$^b$ | 2.42±0.14 |
| Female  | 21  | 19     | 0.33 | 3.07$^a$ | 0.53±0.15 | 0.62 | 5.54$^b$ | 0.31±0.09 | 0.21 | 2.24$^c$ | 2.26±0.61 |

Populations were sorted by genotype, age, and sex.

df degrees of freedom

$^a P<0.01$, significant correlations deduced from values of $t$

$^b P<0.001$, significant correlations deduced from values of $t$

$^c P<0.05$, significant correlations deduced from values of $t$
in genetically modified murine systems. These reported increased arrhythmogenic tendencies [14] that were accompanied by increases in electrogram durations during extracellular recording studies in murine $Scn5a^{+/−}$ hearts [23]. These were followed by in vivo studies that followed cohorts of $Scn5a^{+/−}$ mice stratified by the severity of their conduction changes defined using 18–22- and >22-ms cutoffs in their QRS durations, in parallel with similar clinical stratifications shown.

Fig. 5 Representative slide for cardiac fibrosis staining in WT and $Scn5a^{+/−}$ hearts grouped by age and sex. The study population was stratified into: WT young male (a), WT young female (b), WT old male (c), WT old female (d), $Scn5a^{+/−}$ young male (e), $Scn5a^{+/−}$ young female (f), $Scn5a^{+/−}$ old male (g), and $Scn5a^{+/−}$ old female (h). Hearts were routinely stained with Sirius red, and morphometric analysis for percentage of fibrosis was performed for all eight groups. Areas of increased red uptake signify presence of fibrotic changes. Horizontal bar below sections in each panel denotes a 3 mm distance.
Table 7 Percentage of fibrotic tissue in murine hearts

| Animal           | Samples | Average ± SE |
|------------------|---------|--------------|
| WT               | 23      | 4.73±0.80 a  |
| WT young         | 13      | 2.18±0.26 b, c|
| WT old           | 10      | 7.27±1.08 b, d|
| WT male          | 12      | 4.50±1.02 e  |
| WT female        | 11      | 4.95±1.27    |
| WT young male    | 7       | 2.22±0.41 f, g|
| WT young female  | 6       | 2.12±0.33 h  |
| WT old male      | 5       | 7.93±0.95 f, i|
| WT old female    | 5       | 6.84±1.75    |
| Scn5a+/− young   | 11      | 9.26±0.93 c, j|
| Scn5a+/− old     | 10      | 18.92±3.14 d, j|
| Scn5a+/− male    | 11      | 18.42±2.82 e, k|
| Scn5a+/− female  | 10      | 8.60±1.00 k  |
| Scn5a+/− young male | 6    | 8.78±1.40 g, l|
| Scn5a+/− young female | 5   | 9.74±1.37 h  |
| Scn5a+/− old male | 5     | 24.84±1.53 i, l, m|
| Scn5a+/− old female | 5   | 7.08±1.08 m  |

Means with similar lowercase letters differ significantly from each other at P<0.05

by BrS patients containing SCN5A mutations [10]. These demonstrated parallel changes in conduction and in semi-quantitative measures of fibrosis that varied with age, confirming previous correlations between increased PR intervals and age specifically in the Scn5a+/− [17]. Our subsequent study using closely similar sample numbers then described conduction velocity alterations associated with increased cardiac fibrosis and down-regulation of connexin expression in old Scn5a+/− mice. We thus demonstrated that such alterations in conduction velocity were not solely dependent on reduced Scn5a expression. We then however did not stratify for the interacting effects of age and sex [25]. Finally, recent ECG measurements particularly of PR and RR intervals then went on to fully clarify both the independent and the interacting effects of genotype, age, and sex upon conduction characteristics in fully stratified experimental groups. These showed in particular that old male Scn5a+/− showed marked increases in such conduction durations [7].

We proceeded to examine mean conduction and its dispersion at the tissue, rather than the ECG level, using multi-array recording methods, comparing these with MAP assessments of action potential waveform durations and latencies. Our multi-electrode array studies were less extensive (550 vs 300 μm; 64 vs 247 channels; 1- vs 2–4-kHz sampling) than those used previously [25]. These findings were then correlated with differences in levels of fibrosis, though they did not explore for changes in connexin and Na+ channel expression. Nevertheless, they complemented our earlier studies in exploring for interacting effects of genotype, age, and sex and permitted a novel statistical analysis of activation times and their dispersions both over the regions of interest studied and through successive intrinsic beats. Stratification of these results into experimental groups paralleled and therefore permitted direct comparisons with the earlier, similarly stratified, ECG results [7], thus adding to these a full clarification of alterations in electrogram latencies at the tissue level. They went further to achieve direct correlations of such altered conduction with quantitative assessments of the level of fibrosis. The latter suggested possible mechanisms for such conduction changes, thereby extending and complementing earlier findings that had also assessed for possible existence of fibrosis [10, 17].

The present experiments accordingly provided a more detailed analysis of these conduction disorders on a more microscopic scale as reflected in the activation times shown by epicardial excitation during multi-electrode array recording. They systematically examined the effects of genotype, age, and sex, corresponding to eight experimental groups, on cardiac conduction, determined how these effects alter the subsequent dispersions, and related these findings to morphometric analysis of fibrosis to test this as a possible mechanism underlying such electrophysiological changes.

Initially, the time course of monophasic action potential recovery was assessed at 30%, 50%, 70%, and 90% recovery. This yielded similar values through the stratifications indicated above, suggesting a similarity in all their recovery characteristics. A recent report [11] had described right epicardial APDs in Scn5a+/− that differed from WT. However, this only considered a single age of mouse between 4 and 8 months. It did not perform further stratifications by sex, in contrast to the present study. This difference in findings between 3 and ~6 months suggests that APD can vary with age. Such a notion would parallel recent descriptions of atrial remodeling phenomena in the presence of Scn5a modification [6] and merits further investigation. Nevertheless, the present findings do demonstrate differing action potential latencies corroborating earlier ECG findings [7] under circumstances in which action potential characteristics remained similar.

Our initial qualitative assessment of activation maps derived from activation times at each channel demonstrated that the Scn5a+/− group generally showed higher activation times compared with WT, particularly the older males. They also showed a more heterogeneous pattern of activation, at times with multiple random firing points. Such random activation suggests that the spread of electrical activity in the right ventricle is fragmented, conceivably reflecting patterns of spread suggested on recent occasions [3]. In this respect, it would resemble recent reports describing fragmented QRS complexes characteristics in
BrS patients [13]. It further suggests the presence of areas within the right ventricle that depolarize later than the rest particularly in the Scn5a<sup,+−</sup> mice. Such a finding is compatible with the late potentials, which have previously been observed in signal-averaged ECG thought to represent regions undergoing delayed depolarization than the remaining ventricular tissue [27]. Other clinical studies have previously identified the presence of similarly heterogeneous patterns in BrS patients [9].

In this study, rather than suggesting a sequential spread of conduction within the plane of the epicardium, the Scn5a<sup,+−</sup> mice suggested an irregular pattern of activation in which each epicardial site was separately activated by a radial propagation of excitation through the myocardium. These patterns would be compatible with a greater likelihood of re-entrant events between neighboring epicardial sites and thereby result in arrhythmic substrate. This would be particularly the case if neighboring sites showed strongly different activation times. These might be detectable through the maximum activation times observed and might even then serve as an arrhythmic trigger in parallel with the arrhythmogenicity observed in the corresponding clinical conditions. Quantitative assessments compared such patterns in terms of overall activation times and their dispersions, for the first time. These in turn demonstrated differences that were attributable to both independent and interacting effects of genotype, age, and sex.

Firstly, a mean activation time (T<sub>MEAN</sub>) provided an overall representation of epicardial activation properties resulting from myocardial conduction. This was significantly higher in the Scn5a<sup,+−</sup> compared with WT hearts. However, such differences were less clear with further stratification by both genotype and age. Nevertheless, stratification by both genotype and sex resulted in higher T<sub>MEAN</sub> values in Scn5a<sup,+−</sup> male compared with Scn5a<sup,+−</sup> female hearts. This established an interacting effect of genotype and sex on conduction disorders in mutation carriers. Thus, sex affects T<sub>MEAN</sub> values only in the mutant and not in the WT. The final stratification involving genotype, age, and sex further established interacting effects on T<sub>MEAN</sub> values particularly within the Scn5a<sup,+−</sup> group. Taken together, this analysis demonstrated that alterations in T<sub>MEAN</sub> values, likely reflecting altered conduction, are a major feature of Scn5a<sup,+−</sup> hearts particularly in old male hearts. They also demonstrated that the earlier electrocardiographic findings [7], suggestive of alterations in overall conduction, further extended to the pattern of epicardium activation. Such a finding in the murine Scn5a<sup,+−</sup> model reflects recent clinical findings implicating conduction disorder as feature in arrhythmogenic BrS patients [15].

Secondly, a temporal dispersion D<sub>T</sub> described the temporal variation in activation time at epicardial sites between cardiac cycles. D<sub>T</sub> values were significantly higher in the Scn5a<sup,+−</sup> hearts. Although stratifications by genotype and age did not yield significant differences, those by genotype and sex revealed difference between male Scn5a<sup,+−</sup> and male WT. Finally, stratification by all three variates revealed differences between old male Scn5a<sup,+−</sup> and old male WT. Taken together, such findings suggest that where an influence occurs, temporal dispersions of the ventricular activation time are primarily affected by genotype and not by age and sex.

Thirdly, a spatial dispersion (D<sub>S</sub>) described the spatial variation in activation time of the different epicardial sites in a single cardiac cycle. Unlike both T<sub>MEAN</sub> and D<sub>T</sub>, values for D<sub>S</sub> were indistinguishable when stratified by genotype, genotype and age, and genotype and sex. However, the final stratification by genotype, age, and sex yielded distinct significances. Thus, it demonstrated that the old male Scn5a<sup,+−</sup> hearts had higher dispersions than not only the old male WT but also the other Scn5a<sup,+−</sup> groups.

Such a finding suggests that differences in dispersions of activation times arise only following interacting influences involving all three variates.

Fourthly, the maximum activation time (T<sub>MAX</sub>) described the features of those epicardial sites showing the greatest activation delays that might potentially actually trigger arrhythmia as suggested above. T<sub>MAX</sub> values were significantly higher in the Scn5a<sup,+−</sup> compared with WT hearts, thus establishing an independent effect of genotype on this value. Stratification by genotype and age showed marked effects. They revealed higher T<sub>MAX</sub> values in young Scn5a<sup,+−</sup> than young WT, in old Scn5a<sup,+−</sup> than old WT, and in old Scn5a<sup,+−</sup> than young Scn5a<sup,+−</sup>. Stratification by genotype and sex revealed higher T<sub>MAX</sub> values in male Scn5a<sup,+−</sup> than male WT. Finally, stratification by all three variates revealed strongly interacting effects specifically in the Scn5a<sup,+−</sup> and not the WT. This resulted in higher T<sub>MAX</sub> values in old male Scn5a<sup,+−</sup> compared with old male WT, old female Scn5a<sup,+−</sup>, and young male Scn5a<sup,+−</sup>.

The D<sub>T</sub>, D<sub>S</sub>, and T<sub>MAX</sub> findings thus suggested significant differences in dispersions of epicardial activation times between particular experimental groups. The final analysis assessed the extent to which these differences could be attributed to variations in conduction as represented by T<sub>MEAN</sub>. This would extend the electrocardiographic observations in our previous paper [7] with the physiological findings at the tissue level described here. Such a notion would be compatible with our statistical observations in which significant changes in D<sub>T</sub> and D<sub>S</sub> were always accompanied by corresponding changes in T<sub>MEAN</sub>, but with the reverse not being always the case. Accordingly, we explored for correlations between the above measures of dispersion with such activation times. This demonstrated linear correlations between each of the variables, D<sub>T</sub>, D<sub>S</sub>, and T<sub>MAX</sub> upon T<sub>MEAN</sub>. Each gave
statistically significant linear correlations whether the experimental populations were sorted by genotype, age, or sex. Furthermore, this yielded statistically indistinguishable slopes, suggesting that these correlations were similar through all these conditions. This analysis is thus compatible with a situation in which conduction properties form the major determinant for these dispersions, regardless of genotype, age, and sex.

A final series of investigations tested a hypothesis that implicated the degree of fibrosis in the conduction properties found in each experimental group. They complemented previous qualitative reports that had not fully assessed effects of both age and sex in connection with fibrotic changes associated with the Scn5a+/−/ΔKPQ mutation [10, 17]. The Scn5a+/− hearts then showed generally greater fibrosis than WT hearts. Furthermore, even young Scn5a+/− mice showed increased fibrosis relative to the corresponding young WT. However, this predisposition appeared to apply only to the male and not the female Scn5a+/−/ΔKPQ mice. Consequently, old male Scn5a+/− showed greater fibrosis than the young male Scn5a+/−, the old female Scn5a+/−/ΔKPQ, and the old male WT hearts. The quantitative morphometric analysis of collagen staining used here thus demonstrated patterns of fibrosis that closely matched the electrophysiological findings among the experimental groups sorted by genotype, age, and sex.

In summary, we have investigated for independent and interacting effects of the Scn5a+/− genotype, age, and sex on epicardial activation and its dispersion in murine hearts, which have been previously used to model the corresponding human clinical conditions. Our findings show that genotype, age, and sex all exert independent and interacting effects on cardiac conduction with the greatest effects on ventricular activation times in the old male Scn5a+/−. Such changes directly correlated with the corresponding levels of fibrosis, thereby directly implicating fibrotic changes as important participants in these conduction differences. These effects took place despite an absence of corresponding differences in action potential recovery times as reflected in action potential durations measured at 30%, 50%, 70%, and 90% full recovery. These effects on conduction correlated with greater dispersions of activation times in the Scn5a+/− hearts as a whole compared with WT. The involved variations in activation time are both between cardiac cycles at any given recording site and between recording sites within a given cardiac cycle, as well as the maximum activation times that were obtained during such recordings. These dispersions have been implicated in arrhythmic substrate, as assessed by electrogram analysis in clinical studies [19]. In murine Scn5a+/−, the increased EGD that had been reported previously [23] are directly comparable with the increased dispersion in epicardial activation reported here. This finding again was particularly marked in hearts from old male Scn5a+/− mice. The subsequent regression analysis gave regression coefficients that attributed each of these dispersions directly to differences in conduction. In each case, there was dependence between variates, m, that was similar through all the stratifications that the experiments explored. Such findings implicate similar actions of genotype, age, or sex upon each dispersion, which are mediated through actions on the conduction process. Together, these findings also demonstrate that in contrast to their similar epicardial repolarization features, conduction and dispersion, which are features of depolarization, are prominent in the murine Scn5a+/− model, particularly in older males. They may thus contribute to the reported arrhythmogenic properties of this system.

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