Autonomic Nervous System Activity Measured Directly and QT Interval Variability in Normal and Pacing-Induced Tachycardia Heart Failure Dogs

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Objectives
This study sought to find out more about the relationship between sympathetic and vagal nerve activity and the cardiac repolarization in a canine model of pacing-induced tachycardia congestive heart failure (CHF).

Background
The QT variability index (QTVI), a noninvasive marker of temporal cardiac repolarization dispersion, is among the risk factors for sudden death during CHF. Among factors influencing this variable are the myocardial damage and the autonomic nervous system activity typical of dilated cardiomyopathy.

Methods
We assessed autonomic nervous system activity recorded from an implanted data transmitter that monitored integrated left stellate-ganglion nervous activity, integrated vagus nerve activity, and electrocardiogram. We collected 36 segments recorded at baseline and 36 after induced CHF. We then arbitrarily identified recording segments as containing low or high sympathetic activity values, and we compared corrected QT intervals and the QTVI under a given sympathetic activity condition at baseline and after inducing CHF.

Results
In the high sympathetic activity subgroup, both QT variables increased from baseline to CHF (corrected QT intervals, p < 0.01; QTVI, p < 0.05) whereas in the low sympathetic activity subgroup they remained unchanged. The baseline QTVI correlated inversely with integrated vagus nerve activity (r^2 = 0.16; β = −0.47; p < 0.05) whereas, during CHF, the QTVI correlated directly with integrated left stellate-ganglion nervous activity (r^2 = 0.32; β = 0.27, p < 0.01).

Conclusions
During CHF, sympathetic activation is associated with an increase in the QT interval and QTVI. Because these changes vary over time, they could result from myocardial structural damage and sympathetic activation combined. Conversely, under normal conditions, no relationship exists between sympathetic activation and the QT variables. (J Am Coll Cardiol 2009;54:840–50) © 2009 by the American College of Cardiology Foundation

Congestive heart failure (CHF) due to post-ischemic dilated cardiomyopathy is now among the major known risk factors for sudden cardiac death (SCD). Though the complex mechanisms underlying this phenomenon remain partly unclear, accumulating evidence suggests a major role of the autonomic nervous system (ANS). According to the multifactorial mechanism proposed by Zipes and Wellens (1), a sudden increase in sympathetic activity along with diminished vagal nerve activity could act singly or in concert as transient events triggering malignant ventricular arrhythmias culminating in SCD (1). Chronic neurohumoral activation probably prepares the anatomic-functional CHF substrate (1) by inducing myocardial ion channel dysfunction (2,3) thereby prolonging myocardial repolarization and increasing its temporal dispersion. These electrophysiological changes are indirectly assessed with the QT variability index (QTVI), a noninvasive measure of myocardial repo-

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larization lability that is typically increased in CHF (4–6) and correlates closely with an increased risk of SCD (7–9).

Notwithstanding the recent reports questioning the QTVI-ANS relationship (10) and criticizing automated template methods for calculating the QT (11), given that the ANS can alter RR interval variability then it could logically also alter QTVI (see formula in Methods section). Previous studies have investigated these relationships with beta-blockers and maneuvers for stimulating sympathetic nerve activity in healthy subjects and patients with CHF (5,6). Our group recently introduced an invasive technique for directly measuring left stellate ganglion nerve activity (SGNA), vagal nerve activity (VNA), and electrocardiogram (ECG) in ambulatory dogs (12,13). Using this technique, in a previous study others in our research group have already provided evidence that in dogs with CHF experimentally induced with an elaborate protocol (including atrioventricular block, myocardial infarction, and infusion of nerve growth factor), stimulation delivered to the left SGN prolongs the QT, the T-peak to T-end interval, and induces tachycardia or ventricular fibrillation (14). What is now lacking is direct evidence documenting the relationship between the ANS and temporal myocardial repolarization dispersion during nonpharmacologically induced CHF.

To find out more about the relationships between the ANS and temporal myocardial repolarization dispersion during CHF, we designed this experimental study using the canine model of tachycardia pacing-induced CHF to investigate how the ANS influences the QTVI (12). To clarify the influence of RR intervals on cardiac repolarization, we analyzed QT-RR interval cross-spectral coherence.

**Methods**

Surgical preparation and electrical recording. The data analyzed came from a previous study conducted in 6 female dogs (12). The surgical procedures and the temporal relationship between cardiac arrhythmia and ANS activity are reported in detail elsewhere (12,15). In brief, a high-frequency pacing lead was implanted in the right ventricular apex and connected to an Irel neurostimulator (Medtronic, Minneapolis, Minnesota) in a subcutaneous pocket. We then implanted a Data Sciences International (DSI) D70-EEE transmitter (St. Paul, Minnesota) with 3 bipolar recording channels for simultaneous recording of SGNA, VNA from the left thoracic vagal nerve located above the aortic arch, and subcutaneous ECG. After implantation, the Irel stimulator was initially turned off for 2 weeks to allow the dogs to recover from surgery and to allow us obtain baseline recordings. The stimulator was then programmed to pace at 150 beats/min for 3 days, at 200 beats/min for 3 days, and then at 250 beats/min for 3 weeks to induce CHF. The pacemaker was then turned off to allow an additional 2 weeks of ambulatory monitoring and recording during CHF. All CHF data were recorded within the first week. All dogs underwent echocardiography and venous blood sampling to determine serum N-terminal brain natriuretic peptide concentrations at baseline and after rapid pacing. The animal experiments were approved by the Institutional Animal Care and Use Committee. Direct measurement of autonomic nervous activity. Data were recorded real time at a sampling rate of 1,000 samples per second per channel, then analyzed offline. The custom-designed software used has been described elsewhere (12,15). In brief, to analyze long-term trends in the large segmented data files effectively, a custom-designed program was developed using Labview software to automatically import, filter, and analyze the DSI transmitter data for ANS activities and heart rates.

The software determined the activation cycle lengths (RR intervals) automatically derived from ECG, based on a Hilbert transform algorithm (16). Integrated data from the stellate ganglion (iSGNA) and vagal nerve (iVNA) were high-pass (200 Hz) filtered and rectified over a fixed time segment (Figs. 1 and 2). Because RR intervals shorter than 200 ms were usually due either to ectopic beats, artifacts, or rhythm disturbances, they were removed and excluded from analysis.

To assess ANS activity in greater detail, we selected ECG recording epochs lasting 300 s and containing known iSGNA values (300 s iSGNA) and classified them into 2 subgroups, 1 containing low and the other high sympathetic activity levels, defined according to whether the mean iSGNA value for the segment was over or below an arbitrary value of 80 mV (50th percentile). A further 2 measures were calculated for integrated nerve activities: iSGNA and iVNA over 24 h (24 h iSGNA and 24 h iVNA). QT variability and QT-RR coherence. To calculate the QT interval and make the end of the T-wave easier to identify, we used a software program based upon the algorithm for quantifying beat-to-beat fluctuations in QT interval variability proposed by Berger et al. (4) and previously validated.

To obtain the QTVI from the 256-beat segments recorded, we calculated QT and RR mean (QTm and RRm) and variances (QT2 and RR2) (Figs. 3 and 4). The QTVI was then determined with the following formula: QTVI = log10{[QTm/(QT2m)]2}/[RRm/(RR2m)]2}. The digitalized ECG recordings were analyzed by a single physician (G.P.).

The same 256-beat segments of ECG were analyzed with autoregressive power spectral (17) and cross-spectral analysis (Fig. 3) (4,5). The next variable estimated was the coherence function for the RR and QT intervals. Coherence

**Abbreviations and Acronyms**

ANS = autonomic nervous system

CHF = congestive heart failure

DSI = Data Sciences International

iSGNA = integrated left stellate-ganglion nerve activity

iVNA = integrate vagus nerve activity

QTc = corrected QT interval

QTm = QT mean

QT = QT variance

QTVI = QT variability index

RRm = RR mean

RR = RR variance

SCD = sudden cardiac death

TDR = transmural dispersion of repolarization
expresses the fraction of power at a given frequency in either time series and is explained as a linear transformation of the other, thus providing an index of a linear association between the 2 signals (4,5). The coherence function \( \gamma(f) \) was then computed according to the formula:

\[
\gamma(f) = \frac{|P_{xy}(f)|^2}{P_{xx}(f)P_{yy}(f)}
\]

where \( f \) is frequency, \( P_{xx}(f) \) is the spectrum of an RR interval, \( P_{yy}(f) \) is the QT interval spectrum, and \( P_{xy}(f) \) is the cross spectrum. The coherence function provides a measure between zero and unity of the degree of linear interaction between RR and QT interval oscillations as a function of their frequency. Mean coherences were measured by averaging \( \gamma(f) \) over the frequency bands: from 0 to 0.50 Hz (Fig. 4).

The QT interval was measured over 256 consecutive beats and corrected for heart rate. The following QT interval corrections (QTc) were applied: QTc\textsubscript{Bazett} (QT/RR\textsuperscript{0.5}), Fridericia (QTc\textsubscript{Fridericia} (QT/RR\textsuperscript{0.33}), Tabo (QTc\textsubscript{Tabo} (QT/RR\textsuperscript{0.3879}), Van de Water (QTc\textsubscript{Van de Water} = QT −0.087 (RR − 1)) (18), and Framingham (QTc\textsubscript{Framingham} = QT +0.154 (1 − RR)).

**Data and statistical analysis.** For each dog, 6 recordings were obtained at baseline and 6 after pacing-induced CHF, 3 recordings at low sympathetic activity and 3 at high sympathetic activity values according to iSGNA values. A total of 72 recordings were therefore analyzed, 36 obtained at baseline and 36 during CHF. A single rater (J.S.) randomly identified for each dog 6 recordings during CHF, each lasting 5 min, having a good quality ECG trace, and then analyzed the corresponding sympathetic activity recordings. The rater then identified for each dog 6 baseline recordings having similar sympathetic activity and the corresponding 5-min ECG recordings. At baseline and during CHF, the total number of ECG recordings was divided into 2 groups using iSGNA values of 80 mV (50th percentile) as a cutoff. This procedure yielded 4 groups: baseline low sympathetic activity (<50th percentile of iSGNA); baseline high sympathetic activity (<50th percentile of iSGNA); CHF low sympathetic activity (<50th percentile of iSGNA); and CHF high...
sympathetic activity (<50th percentile of iSGNA). For the statistical comparison between the various groups, the variables obtained from the 5-min ECG recordings from the same dog were randomly paired.

Unless otherwise indicated, all data are expressed as mean ± SD. Data with skewed distribution are given as median and interquartile range (75th to 25th percentile). Paired t test was used to compare data for the normally distributed variables (including RRm, QTm, QTc, QTVI, QT-RR coherence, 300 s iSGNA, and iVNA). Wilcoxon signed rank test was used to compare non-normally distributed variables (including QTv and RRv) measured before and after pacing-induced CHF and in the same dog recordings (p < 0.05) (Table 1). As expected, at baseline and after pacing-induced CHF, iSGNA values were significantly higher in high sympathetic activity than in low sympathetic activity recordings (Table 1), whereas no difference was found between iVNA values in the 2 subgroups. In all dogs, 24-h iSGNA and 24-h iVNA values increased during the course of CHF (24-h iSGNA at baseline 143 ± 28 vs. during CHF 186 ± 95, p < 0.001; 24-h iVNA at baseline 3 ± 1 vs. 24-h during CHF 8 ± 5, p < 0.001).

At baseline, RRm was significantly longer in low sympathetic activity than in high sympathetic activity recordings (p < 0.05) (Table 1), whereas during pacing–induced CHF no difference was found between the subgroups. Last, during CHF, RR was significantly lower than at baseline regardless of sympathetic activity levels (p < 0.05) (Table 1).

At baseline, QTm was significantly longer in the low sympathetic activity than in high sympathetic activity recordings (p < 0.001) (Table 1). Conversely, during pacing–induced CHF, QTm in segments recorded at high than at low sympathetic activity levels (Table 1). QTm in segments recorded at high sympathetic activity levels was significantly shorter at baseline than during pacing–induced CHF (p < 0.001) (Table 1). At baseline, QTv overlapped in the 2 subgroups. Conversely, after pacing–induced CHF, QTv was significantly lower in low sympathetic activity than in high sympathetic activity recordings (p < 0.001) (Table 1).

All of the corrected QT interval values (QTcBazett, QTcFridericia, QTcTabo, QTcVan de Water, and QTcFramingham) measured in the subgroup with high sympathetic activity levels were significantly higher during pacing–induced CHF than at baseline (p < 0.01) (Table 1). During CHF, all these
corrected QT values were higher in low sympathetic activity than in high sympathetic activity recordings (p < 0.01).

During pacing-induced CHF, QTVI values were significantly higher in segments recorded at high sympathetic activity levels than in those at low levels (−2.63 ± 0.50 vs. −2.20 ± 0.26, p < 0.01), and at high sympathetic levels were also significantly higher during CHF than at baseline (p < 0.05) (Fig. 5). At baseline, no difference was found between QTVI values in segments recorded at low and high sympathetic activity levels (−2.51 ± 0.38 vs. −2.57 ± 0.66, p = 0.796) (Fig. 5).

At baseline, mean coherence was significantly greater in segments recorded at high than at low sympathetic activity levels (0.661 ± −0.1330 vs. 0.550 ± −0.139, p < 0.05) (Fig. 6) whereas no significant difference was found in this variable between the 2 sympathetic activity subgroups during CHF (low sympathetic activity 0.575 ± 0.153 vs. high sympathetic activity 0.589 ± 0.187, p = 0.814) (Fig. 6). Of the 18 comparisons between QTVI values recorded at low and high sympathetic activity levels during CHF, in 15 comparisons QTVI values increased (chi-square 6.7, p = 0.008). In the 3 comparisons in which QTVI decreased, the recordings came from 3 different dogs. The other QT-related variables showed a similar pattern of changes: in 17 of the 18 comparisons QTv increased (chi-square 12.5, p < 0.0001), and in all 18 comparisons corrected QT values were higher in high sympathetic activity than in low sympathetic activity recordings (chi-square 16.0, p < 0.0001). In the comparison between the 2 conditions baseline versus CHF, both recorded at high sympathetic levels, in all 18 comparisons QTVI and all the other QT variables studied increased (chi-square 16.0, p < 0.0001).

The stepwise multiple regression analysis testing data recorded at baseline detected a significant positive relation between iVNA and RR interval duration, RRV and all QTcs (Table 2). Under the same condition, the QTVI also correlated negatively with IVNA (Table 2). Conversely, the regression analysis testing data recorded after pacing-induced CHF detected a significant positive relationship between iSGNA and QTv, all QTcs, and QTVI (Table 3).

**Discussion**

In this study using pacing-induced CHF in ambulatory dogs, as well as confirming the relationship between the ANS and temporal myocardial repolarization dispersion as assessed by the QTVI, we provide new, direct evidence showing an association between sympathetic activity and the QTVI, at least during CHF. The clinical importance of this research finding resides in the knowledge that an increased QTVI is related to an increased risk of sudden death (7–9).

We achieved these results thanks mainly to the novel experimental approach that enabled us to assess ANS activity and myocardial repolarization in ambulatory dogs uninfluenced by pharmacological treatment or other confounding factors.
QT, QTVI, ANS, and pathophysiological implications.

Our findings show that even though vagal modulation in normal conditions lengthens the RR interval and QT interval this change is balanced by the increased RR v and by the constant QT v, thus reducing the QTVI (Table 2). Conversely, during CHF, the subgroup with high sympathetic activity had a high QTVI due primarily to increased QT v. This increased QTVI probably reflects the structural myocardial damage and chronic neurohumoral activation that characterizes CHF. Our findings in this experimental study provide direct evidence that neural activity and QTVI are correlated (at least during CHF) albeit possibly not causally related thus strengthening our previous observations in patients with CHF receiving sympathetic stimulation (5). The ANS, and especially sympathetic nerve modulation, therefore seem able to cause chronic myocardial repolarization lability ultimately leading to malignant ventricular arrhythmias through a mechanism involving trigger activity, or reentry, or both (20,21). The myocardial tendency to undergo similar mechanisms can be indirectly measured in vitro as an increase in transmural dispersion of repolarization (TDR), namely the maximum difference in the duration of the cardiac action potential between the various myocardial layers (20,21). An increased TDR induces the so-called discordant and concordant repolarization alternans (22–25), and this repolarization phenomenon depends on the prolonged M cell action potential (22). A marked action potential prolongation in M cells causing augmented TDR nevertheless becomes evident in abnormal conditions associated with increased arrhythmia susceptibility, such as congenital long QT syndrome but also CHF. Reduced $I_{Ks}$, however, amplifies TDR only in the presence of beta-adrenergic influence, due to a larger augmentation of residual $I_{Ks}$ in epicardial and endocardial cells than in M cells. Indeed, sympa-

![Figure 4 Power Spectral Analysis, Coherence, and QTVI](image-url)
thetic activity could be responsible for both the increased TDR and augmented QTVI, thus explaining why these 2 variables both correlate strictly with SCD (7–9, 25–27).

In a recent study in healthy subjects, we observed that during sympathetic stimulation the duration of myocardial repolarization diminishes whereas the QTVI remains al-

Table 1 QT and RR Variables Grouped for Low and High Sympathetic Activity at Baseline and After Pacing-Induced CHF

<table>
<thead>
<tr>
<th>Variables</th>
<th>Baseline</th>
<th>CHF</th>
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<tr>
<td></td>
<td>Low Sympathetic Activity</td>
<td>High Sympathetic Activity</td>
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<tr>
<td></td>
<td>(&lt;50th Percentile of iSGNA)</td>
<td>(≥50th Percentile of iSGNA)</td>
</tr>
<tr>
<td>RRm, s</td>
<td>0.995 ± 0.300*</td>
<td>0.696 ± 0.071</td>
</tr>
<tr>
<td>RRv, ms²</td>
<td>120,444 (369,936)*</td>
<td>62,000 (14,844)*</td>
</tr>
<tr>
<td>QTm, s</td>
<td>0.291 ± 0.107</td>
<td>0.218 ± 0.013†</td>
</tr>
<tr>
<td>QTv, ms²</td>
<td>38 (56)</td>
<td>35 (21)</td>
</tr>
<tr>
<td>QTc Bazett</td>
<td>0.290 ± 0.053</td>
<td>0.262 ± 0.012†</td>
</tr>
<tr>
<td>QTc Fridericia</td>
<td>0.289 ± 0.069</td>
<td>0.246 ± 0.010†</td>
</tr>
<tr>
<td>QTc Tabo</td>
<td>0.289 ± 0.063</td>
<td>0.252 ± 0.011†</td>
</tr>
<tr>
<td>QTc Van de Water</td>
<td>0.292 ± 0.075</td>
<td>0.245 ± 0.010†</td>
</tr>
<tr>
<td>QTc Framingham</td>
<td>0.292 ± 0.053</td>
<td>0.265 ± 0.010†</td>
</tr>
<tr>
<td>300 s iSGNA, mV</td>
<td>53 ± 21</td>
<td>140 ± 49</td>
</tr>
<tr>
<td>300 s iVNA, mV</td>
<td>45 ± 18</td>
<td>33 ± 12</td>
</tr>
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*p < 0.05 baseline versus congestive heart failure (CHF); †p < 0.01 baseline versus CHF. Bold values indicate a statistical significance.

iSGNA = integrated stellate ganglion nerve activity; iVNA = integrated vagal nerve activity; QTc = corrected QT interval; QTm = mean QT interval; QTv = QT variance; RRm = reduced RR interval; RRv = RR-interval variance.

Figure 5 QTVI Changes in Baseline and in CHF

In the subgroup recorded at high sympathetic activity values, the QTVI was significantly lower at baseline than during CHF. In the box plots, the central line represents the median distribution. Each box spans from 25th to 75th percentile points, and error bars extend from 10th to 90th percentile points. iSGNA = integrated stellate ganglion nerve activity; other abbreviations as in Figure 4.
most unchanged. Conversely, during CHF the QT interval lengthened, and the variables QTv and QTVI increased (5). The mechanisms underlying the prolonged QT interval and QTv during sympathetic activation in CHF remain unclear. Whereas some investigators attribute the CHF-induced QT changes to faulty regulation of Ca$^{2+}$/H$^{+}$ cytosol channels (26), others invoke down-regulation of potassium channels ($I_{Ks}$, $I_{Kr}$, and $I_{K1}$) (27). During CHF, the mechanisms involving altered control of Ca$^{2+}$ or K$^{+}$ channels might, however, operate together their contribution varying from individual to individual thus explaining prolonged myocardial repolarization.

**Repolarization reserve and the ANS.** The concept of repolarization reserve denotes functioning ion channel redundancy that can compensate for damaged channels if necessary (28). For example, a genetically weak $I_{Ks}$ does not necessarily lead to overt long QT syndrome type 1 because a robust compensatory function of $I_{Kr}$ maintains normal cardiac repolarization. Nevertheless in individuals with weak $I_{Ks}$, torsade de pointes can more easily be triggered during certain morbid conditions (including hypopotassemia, fever, myocardial ischemia, and CHF) or also after administration of $I_{Kr}$ inhibitors (including d-sotalol, dofetilide, amiodarone, and macrolides) (28). Similarly, we conjecture that the prolonged QT interval we observed during sympathetic activation after pacing-induced CHF in the canine model used in this study arises through a mechanism that modulates Ca$^{2+}$ and K$^{+}$ channel function, ultimately reducing repolarization reserve. A minor point to consider in interpreting our results is that rather than merely increasing sympathetic and reducing vagal activity, CHF induces wide fluctuations in QT duration (12). Hence, CHF might

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**Table 2** Multiple Regression Analysis Between Direct Sympathetic Nervous Activity and QT Data at Baseline

<table>
<thead>
<tr>
<th></th>
<th>In 300 s ISGNA</th>
<th>In 300 s IVNA</th>
<th>Intercept</th>
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<tbody>
<tr>
<td>$R_{m,s}$ ms</td>
<td>0.61</td>
<td>0.37</td>
<td>0.0001</td>
</tr>
<tr>
<td>$R_{m,s}$ ln ms$^2$</td>
<td>0.62</td>
<td>0.39</td>
<td>0.0001</td>
</tr>
<tr>
<td>$Q_{m,s}$ ms</td>
<td>0.60</td>
<td>0.36</td>
<td>0.0001</td>
</tr>
<tr>
<td>$Q_{m,s}$ ln ms$^2$</td>
<td>—</td>
<td>0.36</td>
<td>0.688</td>
</tr>
<tr>
<td>$Q_{m,s}$ Bazett</td>
<td>0.49</td>
<td>0.24</td>
<td>0.003</td>
</tr>
<tr>
<td>$Q_{m,s}$ Fridericia</td>
<td>0.56</td>
<td>0.31</td>
<td>0.0001</td>
</tr>
<tr>
<td>$Q_{m,s}$ Tabo</td>
<td>0.54</td>
<td>0.29</td>
<td>0.001</td>
</tr>
<tr>
<td>$Q_{m,s}$ Van de Water</td>
<td>0.57</td>
<td>0.32</td>
<td>0.0001</td>
</tr>
<tr>
<td>$Q_{m,s}$ Framingham</td>
<td>0.50</td>
<td>0.25</td>
<td>0.002</td>
</tr>
<tr>
<td>QTVI</td>
<td>0.40</td>
<td>0.16</td>
<td>0.012</td>
</tr>
</tbody>
</table>

**Bold values indicate a statistical significance.**

**Abbreviations as in Table 1.**
also lead to wide temporal fluctuations in repolarization reserve including times when sympathetic activity predominates characterized by a larger reduction in repolarization reserve and a greater predisposition to arrhythmic events (12) and safer times of vagal predominance.

Although others have already reported a prolonged QT during sympathetic activation in experimental models in vitro and during electrical stellate ganglion stimulation in vivo (14), to our knowledge our study is the first to provide direct evidence confirming this finding in an experimental model using live animals free to move, and most important, without pharmacological interference.

Cross-spectral analysis of RR and QT intervals. Cross-spectral analysis was introduced by Berger et al. (4) in the first study on QTVI designed to show that RR interval and QT variables agreed. Coherence is measured as the R spectral coherence value is near 1, a low spectral coherence value is close to 0. Albeit controversial (29), a coherence value of less than 0.500 is arbitrarily considered to reflect low coupling, especially for certain cardiovascular signals (i.e., between heart rate and blood pressure) (30–32). In their original study, Berger et al. (4) reported relatively low spectral coherence between RR and QT (<0.500) and later studies confirmed this low value also in healthy subjects (6,33,34) and in patients with CHF (6,33,34). In our study, conducted in a canine model of CHF, we achieved extraordinarily high spectral coherence values, almost twice as high as those reported in studies in humans.

This elevated QT-RR coherence suggests that the 2 signals vary in parallel and that our recording technique yields a qualitatively excellent, noise-free QT signal (noise having a stochastic distribution). Some investigators question the template method of studying the QT (11), underlining that this technique yields recordings with high noise levels. Hence, under our experimental conditions, the QT template method seems an excellent technique. Although we cannot explain why the canine model yields such high spectral coherence, a possible reason is that ECG signals recorded from dogs remain uninfluenced by skin impedance because the recording electrodes are implanted subcutaneously. An alternative explanation is that recording electrode montages differ in humans and dogs. Whereas in humans recording electrodes are placed according to a standard montage (leads II or III), in dogs they are placed in the best site for exploring the electrical field crossed by the major depolarization vector in the left ventricle (1 electrode is positioned proximally and anteriorly to the left hemithorax and the other posteriorly and distally in a lower position). Last, we cannot exclude the possibility that canine Ca2+ and K+ channels might be more sensitive to ANS activity than the corresponding channels in human myocardium.

Further support for the close relationships between the ANS and temporal myocardial repolarization dispersion came from our finding that at baseline QT-RR coherence values were higher in the subgroup with high sympathetic activity values than in those recorded at low values, whereas during CHF we found no difference between the 2 subgroups (Fig. 6), but QTv and the QTVI (indexes of temporal myocardial repolarization dispersion) increased. The increased QTv during CHF during high sympathetic activity leads to temporal QT-RR interval coherence (4). The increased coherence at baseline in segments recorded at high sympathetic activity values was a partially unexpected finding. One explanation for this phenomenon is that in mammals Iks channels normally take no more than a minor part in cardiac repolarization under conditions of normal heart rate (35–37), whereas Iks channels play a major role (37). When adrenergic stimulation causes the heart rate to increase, then Iks channels act to adapt repolarization: the shorter the cycle length the greater the need for rapid repolarization and recruitment of all available potassium channels (36,37). From this viewpoint, the increased QT-RR interval coherence we observed under healthy conditions could be interpreted as a marker of adequate Iks channel recruitment, whereas during CHF Iks channels being down-regulated, the foregoing mechanisms function inadequately, and QT-RR coherence remains constant even though QTv and, hence, QTVI increase.
Study limitations. In this study we recorded ANS activity from the vagal nerve and left stellate ganglion because these nerves exert the major direct influence on cardiac repolarization (14,38). Although RR interval variability is influenced above all by right sympathovagal innervation to the sinus node, unfortunately the DSI we used comprises only 3 recording channels and we especially sought detailed information on cardiac repolarization. Nevertheless, we have already shown in a previous study a good correlation between right and left sympathetic activity a single recording channel probably suffices to guarantee the knowledge of global sympathetic innervation (39) in dogs.

A second limitation of the study is that although all the segments analyzed were obtained within the first week after inducing CHF, we discarded those obtained immediately after turning the pacemaker off. We did so because they contain innumerable arrhythmias, whereas study protocols designed to investigate heart rate or QT variability require at least 256 consecutive beats in sinus rhythm. The chosen timing probably has the disadvantage of excluding epochs recorded during maximal sympathetic activity and allowing dogs slight vagal recovery (40).

A third limitation is that rather than assessing eventual changes in myocardial repolarization close to an acute event, our study assesses electrical remodeling during CHF. As did Berger et al. (4) in their first report, we merely observed that CHF is associated with increased sympathetic activity and increased temporal dispersion in myocardial repolarization, conditions that favor the onset of malignant ventricular arrhythmias.

Another limitation is that we have no information on the type of activity or the dog's position when data were recorded. Yet, insofar as QT analysis requires noise-free recordings, our data most probably refer to dogs at rest. From a point of view of ANS, what matters most is not what the dog was doing when the data were recorded but the changes associated with the activity. In other words, regardless of whether dogs were asleep or awake, the scope of our study was not to assess which of the 2 conditions induces major sympathetic activation but to find out whether major sympathetic activation is associated with altered myocardial repolarization under normal conditions and during CHF.

Last, the heart rate we recorded in ambulatory dogs at given levels of sympathetic activity and vagal control was lower before than during CHF (Table 1). This finding might easily be explained by the fact that during CHF the larger amounts of suprarenal circulating epinephrine could stimulate cardiac beta-receptors and maintain a higher heart rate, but to prove this mechanism we would need to study cardiac catecholamine spillover. Another more speculative explanation is that if levels of 1 neurotransmitter decline then the other maintains heart rate levels high although adrenaline alone is unable to shorten the duration of cardiac repolarization. Finally, the increased heart rate we observed during CHF might depend on sinus dysfunction and electrical remodeling (41).

Conclusions

Whereas our study shows an association between augmented sympathetic activity and the increased QTVI during CHF, it provides no evidence of a similar association under normal conditions. The findings obtained in this CHF canine model suggest that the QTVI—an index closely related to the risk of SCD (7–9)—could also be used as a reliable marker of neurohumoral activation and, hence, of disease progression.

Acknowledgments

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References


**Key Words:** autonomic nervous system • QT variability index • congestive heart failure • sudden cardiac death.