QT Interval Variability and QT-HP Coupling Strength in Amyotrophic Lateral Sclerosis Patients

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Abstract

In the recent years we have witnessed an increasing interest in studying cardiac control in amyotrophic lateral sclerosis (ALS) patients. The variability of the overall duration of cardiac electrical activity comprising depolarization and repolarization periods, namely the QT interval, could provide information about the cardiac control of ALS patients complementary to that derived from the variability of heart period (HP). In this study we evaluate first the HP and QT variabilities in 10 ALS patients at rest in supine position (REST) and during 75° head-up tilt (TILT). The QT interval was approximated as RTa and RTe wave peak and apex and end, respectively. HP was taken as the time distance between two consecutive R-wave peaks. Time and frequency domain markers were computed over HP, RTa and RTe beat-to-beat series. The RTa-HP and RTe-HP squared coherence was calculated as well. We found that time and frequency domain indexes derived from QT variability changed during TILT in the direction expected for a healthy population. Frequency domain HP variability markers showed a blunted response to TILT. RTa-HP and RTe-HP squared coherence did not vary during TILT. QT variability and QT-HP coupling markers in ALS patients showed an apparently normal response to TILT not fully mirrored by HP variability indexes.

1. Introduction

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease characterized by the loss of the upper and lower motoneurons. In the last years, signs and symptoms of autonomic nervous system involvement have been described in ALS, including subclinical dysfunction of cardiovascular, sudomotor, gastrointestinal, salivary and lacrimal regulations. At the cardiovascular control level, impaired baroreflex response and blunted response to postural stressor have been described [1-3]. Nevertheless, results about the autonomic nervous system involvement in ALS are still controversial. In addition, no clear relation between autonomic disturbances and patients’ clinical characteristics and prognosis have been identified. New approaches to the study of the cardiac control of ALS patients are needed to improve the characterization of ALS patients. The study of the variability of the time taken for ventricular depolarization and repolarization, approximated by the QT interval measured on the electrocardiogram (ECG), could provide new insight. This analysis has been demonstrated to be useful in describing the sympathetic drive directed to the heart, in particular to the ventricles, with prognostic meaning [4,5]. To the best of our knowledge, only two studies quantified the QT interval in ALS patients [6,7], but none of them monitored the QT variability and the coupling strength between QT and heart period (HP) variations.

In this study we test the hypothesis that the analysis of the QT variability can be performed in ALS patients and we evaluate the changes of QT variability during a postural challenge in ALS patients. Indexes of QT variability were examined with HP variability markers to assess potential additional information. QT-HP coupling strength was monitored as well.

2. Experimental protocol and analysis

2.1. Experimental protocol

We enrolled 10 ALS patients (2 males; age: 62.5±7.55 years; body mass index: 22.76±4.08 kg·m⁻²). Eight patients had spinal onset, while 2 patients had bulbar onset. The time from diagnosis was 27.9±16.38 months. The study was conducted at IRCCS Istituti Clinici Scientifici Maugeri, Milan, Italy, with the approval of the Local
Ethics Committee (CE1077). The study is in keeping with the principles of the Declaration of Helsinki and each subject signed a written informed consent.

All the subjects were instructed to avoid caffeinated beverages in the 24 hours preceding the test and reached the laboratory by wheelchair. During the experiment ECG (modified lead II) was acquired at 1000 Hz. Thoracic movements were monitored as well to estimate breathing rate. The signals were recorded for 10 minutes at rest in supine position (REST) and for 5 minutes during head-up tilt test at 75° (TILT). All the patients completed the experiments without signs of pre-syncope.

2.2. Beat-to-beat time series extraction

The HP was approximated as the temporal distance between two consecutive R-wave peaks detected over the ECG. QT interval was approximated as the temporal distance between the R-wave peak and the T-wave apex (RTa) and as the temporal distance between the R-wave peak and T-wave end (RTe) [8]. Both the apexes of R-wave and T-wave were fixed using parabolic interpolation [8]. The end of the T-wave was identified as the time when the first derivative, calculated on the T wave downslope, became smaller of the 30% of the maximum first derivative value [8]. The ith RTa and RTe measures always followed the ith HP, thus linking the ith RTa and RTe to the previous HP. The detections of the R-wave peaks, T-wave apexes and T-wave ends were visually checked and manually corrected in case of erroneous identifications. In case of ectopic beats, the series were corrected by means of linear interpolation. Corrections never exceeded 5% of the total values analyzed in each session. HP, RTa and RTe series of 256 consecutive values were selected randomly in each experimental session (i.e. REST or TILT). Examples of HP, RTa and RTe series at REST and during TILT are shown in Fig.1. We computed mean and variance of all series indicated, respectively, as \( \mu_{\text{HP}} \), \( \mu_{\text{RTa}} \), \( \mu_{\text{RTe}} \) and \( \sigma^2_{\text{HP}} \), \( \sigma^2_{\text{RTa}} \), \( \sigma^2_{\text{RTe}} \). Means were expressed in ms, while variances in ms².

2.3. Power spectral analysis

Parametric power spectral analysis was performed on HP, RTa and RTe series. The series were described by an autoregressive model, whose order was chosen according to Akaike information criterion. Power spectral density was estimated from the identified coefficient of the autoregressive model and the variance of the prediction error [9]. The power spectral density was decomposed into spectral components. Each component was classified as low frequency (LF, 0.04-0.15 Hz) or high frequency (HF, 0.15-0.5 Hz) according to their central frequency [9]. The sum of the LF components of HP, RTa and RTe was expressed in absolute units (i.e. ms²) and termed, respectively, as LFₐ₉ₜ₉, LFₐ₉ₜₐ and LFₐ₉ₜₑ₉. The LFₐ₉ₜ₉, LFₐ₉ₜₐ

<table>
<thead>
<tr>
<th>Index</th>
<th>REST</th>
<th>TILT</th>
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</thead>
<tbody>
<tr>
<td>( \mu_{\text{HP}} ) [ms]</td>
<td>871.19±138.73</td>
<td>776.41±82.57*</td>
</tr>
<tr>
<td>( \mu_{\text{RTa}} ) [ms]</td>
<td>269.48±20.13</td>
<td>260.26±19.32*</td>
</tr>
<tr>
<td>( \mu_{\text{RTe}} ) [ms]</td>
<td>342.17±19.66</td>
<td>328.89±10.9*</td>
</tr>
<tr>
<td>( \sigma^2_{\text{HP}} ) [ms²]</td>
<td>842.37±1058.57</td>
<td>531.5±536.84</td>
</tr>
<tr>
<td>( \sigma^2_{\text{RTa}} ) [ms²]</td>
<td>19.08±21.52</td>
<td>59.38±73.75*</td>
</tr>
<tr>
<td>( \sigma^2_{\text{RTe}} ) [ms²]</td>
<td>84.66±73.64</td>
<td>165.81±159.23*</td>
</tr>
</tbody>
</table>

REST: at rest in supine position; TILT: head-up tilt; HP: heart period; RTa: R-wave peak to T-wave apex; RTe: R-wave peak to T-wave end; \( \mu_{\text{HP}} \): HP mean; \( \mu_{\text{RTa}} \): RTa mean; \( \mu_{\text{RTe}} \): RTe mean; \( \sigma^2_{\text{HP}} \): HP variance; \( \sigma^2_{\text{RTa}} \): RTa variance; \( \sigma^2_{\text{RTe}} \): RTe variance. The symbol * indicates \( p<0.05 \) versus REST.
power can be considered to be a marker of the sympathetic and vagal modulation directed to the sinus node [9], while LF_{a,RTe} and LF_{a,RTe} powers to be markers of sympathetic modulation directed to the ventricles [4,5]. The sum of the HF components of HP was expressed in absolute units (i.e. ms^2) and labelled as HF_{a,HP}. The HF_{a,HP} power is a marker of the vagal modulation directed to the sinus node [9]. For HP series, LF and HF powers were also expressed in normalized units (LF_{nu,HP} and HF_{nu,HP}, respectively) as the ratio of LF_{a,HP} and HF_{a,HP} to \sigma_{a,HP}^2 minus the power of components with central frequencies below 0.04 Hz [10]. Normalization makes the LF_{nu,HP} index more linked to sympathetic modulation directed to sinus node than the LF_{a,HP} power [10].

### 2.4. Squared coherence analysis

The assessment of the coupling strength between HP and RTa, or RTe, was performed by exploiting the squared coherence function K^2. It represents the degree of linear correlation between the two series as a function of the frequency. K^2 was calculated as the ratio between the square cross-spectrum modulus between HP and RTa, or RTe, to the product of the power spectra of RTa, or RTe, and HP series. The estimation of the cross-spectrum was grounded over the identification of a bivariate autoregressive process and the model order was set to 10 [5]. K^2 was sampled in correspondence of the weighted central frequency of the HP components in LF and HF bands, where the weights were the powers of the components [5]. Markers in the LF and HF bands were indicated as K^2_{RTa-HP,LF}, K^2_{RTe-HP,LF}, K^2_{RTa-HP,HF}, K^2_{RTe-HP,HF}. K^2_{RTa-HP,LF} and K^2_{RTe-HP,LF} ranged from 1 (full correlation) to 0 (null correlation) and were dimensionless.

### Table 2. HP and QT power spectral markers

<table>
<thead>
<tr>
<th>Index</th>
<th>REST</th>
<th>TILT</th>
</tr>
</thead>
<tbody>
<tr>
<td>LF_{nu,HP} [ms^2]</td>
<td>125.72±181.21</td>
<td>121.4±152.69</td>
</tr>
<tr>
<td>LF_{nu,HP} [nu]</td>
<td>46.47±18.83</td>
<td>52.95±26.08*</td>
</tr>
<tr>
<td>HF_{a,HP} [ms^2]</td>
<td>91.32±95.65</td>
<td>66.64±64.4</td>
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<tr>
<td>HF_{nu,HP} [nu]</td>
<td>48.51±17.32</td>
<td>39.51±23.07</td>
</tr>
<tr>
<td>LF_{a,RTa} [ms^2]</td>
<td>4.81±5.32</td>
<td>14.93±17.18*</td>
</tr>
<tr>
<td>LF_{a,RTe} [ms^2]</td>
<td>20.59±16.76</td>
<td>54.65±69.05*</td>
</tr>
</tbody>
</table>

REST: at rest in supine position; TILT: head-up tilt; HP: heart period; RTa: R-wave peak to T-wave apex; RTe: R-wave peak to T-wave end; LF: low frequency; HF: high frequency; LF_{nu,HP}: absolute power of HP series in LF band; HF_{a,HP}: absolute power of HP series in HF band; HF_{nu,HP}: normalized power of HP series in HF band; LF_{a,RTa}: absolute power in LF band of RTa variability; LF_{a,RTe}: absolute power of RTe series in LF band. The symbol * indicates p<0.05 versus REST.

### Table 3. QT-HP coupling strength parameters

<table>
<thead>
<tr>
<th>Index</th>
<th>REST</th>
<th>TILT</th>
</tr>
</thead>
<tbody>
<tr>
<td>K^2_{RTa-HP,LF}</td>
<td>0.24±0.18</td>
<td>0.17±0.12</td>
</tr>
<tr>
<td>K^2_{RTe-HP,LF}</td>
<td>0.19±0.26</td>
<td>0.13±0.12</td>
</tr>
<tr>
<td>K^2_{RTa-HP,HF}</td>
<td>0.14±0.1</td>
<td>0.14±0.15</td>
</tr>
<tr>
<td>K^2_{RTe-HP,HF}</td>
<td>0.14±0.07</td>
<td>0.12±0.08</td>
</tr>
</tbody>
</table>

REST: at rest in supine position; TILT: head-up tilt; HP: heart period; RTa: R-wave peak to T-wave apex; RTe: R-wave peak to T-wave end; K^2: squared coherence; LF: low frequency; HF: high frequency; K^2_{RTa-HP,LF}: K^2 between RTa and HP series in the LF band; K^2_{RTe-HP,LF}; K^2 between RTa and HP series in the HF band; K^2_{RTa-HP,HF}; K^2 between RTe and HP series in the LF band; K^2_{RTe-HP,HF}; K^2 between RTe and HP series in the HF band.

### 2.5. Statistical analysis

The significance of the differences of HP variability, QT variability and K^2 indexes between the two experimental conditions (i.e. REST and TILT) was tested via paired t-test, or Wilcoxon signed rank test when appropriate. Data were reported as mean±standard deviation. A p<0.05 was always considered as significant.

### 3. Results

Table 1 shows the results of the analysis of HP, RTa and RTe variability in the time domain. \mu_{HP}, \mu_{RTa} and \mu_{RTe} decreased during TILT compared to REST. \sigma^2_{RTa} and \sigma^2_{RTe} significantly increased during TILT compared to REST, while \sigma^2_{HF} did not change. Table 2 shows the results of the power spectral analysis of HP, RTa and RTe variability. LF_{a,RTa} and LF_{a,RTe} powers significantly increased during TILT in ALS patients. Only the LF_{nu,HP} increased during TILT, while none of the other HP variability spectral markers changed significantly. Table 3 shows the results of the K^2 analysis between HP and RTa, or RTe, in both LF and HF bands. Both RTa-HP and RTe-HP coupling strength remained stable across the experimental condition and this result held regardless of the frequency band.

### 4. Discussion

The main novelty of this study is the assessment of QT variability at REST and during TILT in ALS patients, together with the more traditional evaluation of HP variability. We found that QT variability can be analyzed in ALS patients and might represent a complementary tool with respect to the HP variability to investigate the cardiac control in ALS. Indeed, results showed that \mu_{RTa} and \mu_{RTe} decreased during TILT, as the consequence of the shortening of the HP and the QT-RR relationship. As observed in healthy subjects [5,11], the variance of QT, approximated by \sigma^2_{RTa} and \sigma^2_{RTe}, and the QT variability
power in LF band, approximated by LF$_{a,RTa}$ and LF$_{a,RTc}$. Increased during TILT as a consequence of the sympathetic activation induced by the orthostatic challenge. We conclude that the typical response of QT variability to TILT is preserved in our group of ALS patients. This result is in agreement with [7] who did not observe abnormalities in the QT interval of ALS patients. Remarkably, ALS patients were characterized by a stable degree of correlation between HP and QT variability during TILT. This result is at difference with healthy subjects [5,12]. Given the low level of QT-HP coupling observed in this study even at REST, this finding might indicate that the QT-HP relation is impaired in ALS patients and this impairment might favor the QT dispersion. It is worth noting that an increase of QT dispersion was observed in the terminal stages of ALS compared to initial stages and this increase was taken as a hallmark of an increased risk of sudden death [6].

As to the HP variability, ALS individuals showed a limited response to TILT compared to QT variability. Indeed, while the decrease of μ$_{HP}$ and the increase of LF$_{a,HP}$ power in reaction to TILT was found, a significant decrease of σ$_{HP}$ and HF$_{a,HP}$ power was not observed. This peculiar response of HP variability to TILT was already documented in [1,2]. This blunted response of HP variability to TILT in connection with the preserved QT variability and QT-HP coupling strength suggest that the analysis of QT variability might provide information complementary to that of HP variability.

5. Conclusion

In this study, we evaluated first the QT variability in ALS patients during an orthostatic challenge. We found that the sympathetic modulation directed to the heart, as quantified by QT variability analysis, increased in response to the postural change. However, modifications of the HP variability parameters were blunted and the expected reduction of the degree of association between HP and QT with TILT was absent. However, even though the response of QT variability to postural maneuver is preserved [5,11], the lack of comparison with age-matched control group does not allow us to fully understand the magnitude of this response. Moreover, the preservation of QT-HP coupling with TILT stresses the need of studying the QT-HP relationship via, for example, ad-hoc modeling approaches [11,12].

In general, the comparison between findings derived from QT and HP variability series allows us to conclude that the QT variability in ALS patients might provide information complementary to that given by the exclusive analysis of HP variability and might offer a more precise picture about the cardiac control of ALS patients.

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References


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