Decreased Heart Rate Variability in Individuals With Amyotrophic Lateral Sclerosis

Renata M M Pimentel, Hugo Macedo Jr, Vítor E Valenti, Fernando O Rocha, Luiz Carlos Abreu, Carlos Bandeira de M Monteiro, and Celso Ferreira

BACKGROUND: Amyotrophic lateral sclerosis (ALS) is considered a multisystem degenerative disease due to its autonomic dysfunction effects. Autonomic cardiac control disorders can be seen in ALS and influence the quality of life and the life expectancy of affected individuals. We evaluated heart rate variability in subjects with ALS and with variable lung capacity. METHODS: We conducted a prospective cross-sectional study performed in 42 subjects with ALS. The subjects were classified into 2 groups according to their FVC: (FVC > 50% of the predicted value [n = 19] and FVC < 50% of the predicted value [n = 23]). Heart rate was recorded at rest during spontaneous breathing by using a heart rate monitor. Linear indices in the time and frequency domains were analyzed, and non-linear analysis was performed by using Poincaré plots. RESULTS: The results showed a decrease of heart rate variability in the subjects with lower lung capacity and who needed ventilatory support. Qualitative analysis when using the plots supported the quantitative analysis, wherein the group with a lower lung capacity showed reduced heart rate variability. No significant differences were found in the other heart rate variability indices. CONCLUSIONS: The subjects with ALS and with decreased pulmonary capacity had reduced heart rate variability. Key words: motor neuron disease; amyotrophic lateral sclerosis; spirometry; autonomic nervous system; nervous system diseases; respiratory function. [Respir Care 0;0(0):1–. © 0 Daedalus Enterprises]

Introduction

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease characterized by the death of motor neurons,1 which eventually leads to respiratory failure and medulla oblongata dysfunction.2 There is recent evidence that an ALS diagnosis may be well established before the symptoms become apparent.3 According to the Brazilian Amyotrophic Lateral Sclerosis Association,3,4 a series of studies established that progressive bulbar palsy and primary lateral sclerosis are subtypes of motor neuron disease. Motor neuron disease in general is related to a degenerative process of the central nervous system, with a variable clinical spectrum, but these subtypes could present different associated clinical characteristics. Twenty percent of patients begin in the bulbar group, which most often presents with either dysarthria or dysphagia.4

The association of motor neuron death with mutations in the Cu and Zn superoxide dismutase (SOD1) genes5 is
in agreement with studies that demonstrated autonomic failure in subjects with ALS with a mutated SOD1 gene in neurons. Recent studies showed the existence of subclinical cardiovascular, gastrointestinal, sweating, and salivary and lacrimal dysfunctions, even in the early phases of ALS, which indicates abnormal sympathetic activity during the early stages of the disease with hyperhidrosis and a reduction in sweat production as the disease progresses. Furthermore, in advanced stages and in patients who are dependent on ventilatory support, apparent autonomic disorders may lead to circulatory collapse or sudden death. Thus, ALS is a degenerative multisystemic disease that results from related autonomic dysfunction.

Breathing modulates the autonomic flow to the heart and contributes to changes in heart rate variability because of the influence of breathing on the autonomic system, some studies have assessed heart rate variability while performing techniques used for respiratory therapy by integrating the management of cardiac care subjects. It is important to detect autonomic cardiac control disorders early because of the influence on survival and quality of life. Orthostatic hypotension, nocturnal blood pressure fluctuations, or falls and postural dizziness are increasingly reported. Studies reported decreased heart rate variability coincident with impaired salivary excretory function and gastrointestinal tract function, which indicate parasympathetic abnormalities, whereas other studies showed impaired cardiac sympathetic innervation. We believe that these associated disorders in different neurodegenerative diseases could affect the clinical course of ALS by increasing the risk of cardiac arrhythmia and sudden death.

The autonomic nervous system regulates the cardiovascular and respiratory systems. However, there is little understanding of how ALS is involved in cardiac arrhythmias. Because patients with ALS often have predominantly respiratory symptoms, diagnosis and treatment of cardiac arrhythmias can be difficult. Moreover, because the clinical course is dependent on disease progression and worsening of muscle and respiratory deficiencies, it is necessary to examine autonomic cardiac control in patients with motor neuron disease and to associate these findings with respiratory capacity. Therefore, we aimed to evaluate heart rate variability in ALS with and without bulbar palsy in subjects with variable lung capacity.

Methods

STROBE Guidelines

Our study conformed to the STROBE (Standards of Reporting of Observational Studies in Epidemiology) guidelines. Our investigation contained details of the study design, setting, participants, variables, data sources, measurement, description of potential sources of bias, quantitative variables description, and statistical methods.

Population Study and Eligibility Criteria

This study was performed in 42 subjects diagnosed with ALS. The group of subjects was divided and classified according to their FVC. The protocol was applied during the assessment carried out during subject visits. Subjects with an FVC < 50% of the predicted were using noninvasive ventilation, usually at night. No subject had used invasive mechanical ventilation. As inclusion criteria to participate in this study, we considered all the patients in clinical follow-up at the Department of Neuromuscular Diseases at Department of Neuromuscular Diseases of the Federal University of São Paulo diagnosed with ALS and with the confirmation of characteristics in El Escorial supported by an electroneuromyographic evaluation. All the participants provided signed informed consent (subject or representative). Patients with congenital abnormalities, heart disease, and pulmonary malformations, and those patients using drugs such as antiarrhythmics, insulin, and medications that influence the autonomic nervous system were excluded.

Ethics Statement

The study was approved by the ethics committee in Research of the Federal University of São Paulo (CAAE 30799414.9.0000.5505).

QUICK LOOK

Current knowledge

Amyotrophic lateral sclerosis (ALS) is considered a multisystem degenerative disease, which can lead to several subclinical disorders linked to autonomic dysfunction. Changes in the autonomic nervous system are associated with cardiovascular dysfunctions. Because patients with ALS usually have predominantly respiratory symptoms, diagnosis and treatment of cardiac arrhythmias can be difficult.

What this paper contributes to our knowledge

Subjects with ALS with decreased lung capacity present reduced heart rate variability. We believe that this autonomic dysfunction may affect the clinical course of ALS and increase the risk of cardiac arrhythmia and sudden death.
Study Design and Setting

This was a prospective, observational, and cross-sectional study. Recruitment was carried out by the Brazilian Amyotrophic Lateral Sclerosis Association in connection with the Department of Neuromuscular Diseases of the Federal University of São Paulo.

Bias

We performed all protocols by using the same conditions to address potential sources of bias. Variables were collected under controlled temperature (21–25°C) and humidity (50–60%). Abstinence from caffeine, alcohol, cola beverages, chocolate (for 12 h), and exhaustive effort (24 h) was recommended to individuals. Heart rate variability was evaluated between 1:00 PM and 6:00 PM to standardize the interference of circadian rhythm. All procedures necessary for data collection were explained to the individuals, and the subjects were instructed to remain at rest during the data collection.

Initial Assessment and Experimental Protocols

Data collection started after obtaining informed consent from the subjects or their representatives. Heart rate was recorded for 20 min while the subject was seated at rest and spontaneously breathing. Subsequently, spirometry was performed while the subject remained seated by the same respiratory therapist through a portable spirometer (MIR Spirobank version 3.6 New Berlin, WI) coupled to a microcomputer that used the Ocean & WinSpiro software (New Berlin, WI) for Windows version 1.04® for analysis. The subjects were classified according to their FVC (>50% or <50% of the predicted value). Heart rate, breathing frequency, and oxygen saturation (SpO₂) were also monitored at rest.

Variables and Data Sources

Instantaneous RR intervals were recorded with a digital telemetry system, which consisted of a transmitter placed on the subject’s chest and a heart rate monitor. This equipment was previously validated to capture the heart rate for heart rate variability analysis.¹⁹ Oxygen saturation was determined by pulse oximetry (DX2010, Dixtal, São Paulo, SP, Brazil) connected to the subject’s second finger through an appropriately sized sensor. The saturation of hemoglobin oxygen was recorded after stabilization as the most constant value during the second minute.

After placing the heart rate monitor, the participants remained seated, at rest. This position was chosen because many of the subjects were in a wheelchair throughout the day. In addition, most patients do not remain in the supine position due to diaphragmatic muscle weakness, which may result in discomfort and cause a significant change in heart rate variability. We respected the initial 5-min rest interval to stabilize the heart rate. For analyzing heart rate variability data, we selected consecutive stable RR intervals, then filtering was performed through Microsoft Excel (Microsoft, Redmond, WA) to eliminate artifacts. The first 1,000 intervals of beats were chosen, and only series with >95% sinus beats were included in the study.²⁰ The software used for heart rate variability evaluation was Kubios heart rate variability version 2.0 (Helsinki, Finland).

Outcome Measures

We analyzed the time and frequency domain heart rate variability indices and non-linear methods through Poincaré plots. For heart rate variability analysis in the frequency domain, we used spectral components of low frequency (0.04–0.15 Hz) and high frequency (0.15–0.40 Hz) in ms² and normalized units (nu). Spectral analysis was calculated by using the fast Fourier transform algorithm. The analysis in the time domain was performed through the mean ± SD of normal RR intervals and the root of the square successive differences between adjacent normal RR intervals. We used the Kubios heart rate variability analysis software for the linear analysis of heart rate variability.²¹

The RR triangular index was calculated based on the construction of the density histogram of normal RR intervals and was obtained by dividing the histogram integral interval (ie, the total number of RR intervals) by the maximum density distribution (modal frequency of RR intervals) measured on a discrete scale with boxes of 7.8125 ms (1/128 s).²² The triangular interpolation of RR intervals constitutes the width of the baseline distribution as measured by the base of a triangle; the distribution of all RR intervals and the difference of least squares were used to determine the triangle.²²

The heart rate variability Poincaré plot was analyzed from the RR intervals based on non-linear dynamics.²³,²⁴ This analysis was a 2-dimensional graphic representation of the correlation between consecutive RR intervals, in which each interval was plotted against the next interval.²³,²⁵,²⁶ For the quantitative analysis of the Poincaré plots, the following indices were calculated: SD of the instantaneous variability beat by beat (SD1), SD for long-term of continuous RR intervals (SD2), and SD1/SD2.²⁷ Qualitative analysis (visual) of the Poincaré plots was performed through the analysis of the figures formed by the attractor of the plot, which were described by Tulppo et al.²⁸

1. A figure in which an increase in the dispersion of RR intervals beat by beat is observed, with an increase in RR intervals, characteristic of a normal plot.
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Table 1. Distribution of Study Subjects by Age, Sex, Baseline Heart Rate, Oxygen Saturation

<table>
<thead>
<tr>
<th>Variable</th>
<th>FVC &gt; 50%</th>
<th>FVC &lt; 50%</th>
<th>P</th>
<th>Cohen’s d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean ± SD y</td>
<td>56.5 ± 9.8</td>
<td>58.7 ± 9.2</td>
<td>.47</td>
<td>NA</td>
</tr>
<tr>
<td>Males, n</td>
<td>8</td>
<td>12</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Females, n</td>
<td>11</td>
<td>11</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Heart rate, mean ± SD beats/min</td>
<td>80.11 ± 10.03</td>
<td>87.3 ± 10.44</td>
<td>.03</td>
<td>0.7</td>
</tr>
<tr>
<td>Oxygen saturation, mean ± SD %</td>
<td>96.41 ± 2.12</td>
<td>94 ± 3.1</td>
<td>.008</td>
<td>0.9</td>
</tr>
</tbody>
</table>

NA = not applicable

2. A figure with little global dispersion beat by beat and without increased dispersion of RR intervals in the long term.

Kubios Heart Rate Variability software Version 2.0 was used to assess these indices.21 Vital capacity was measured by spirometry. The subjects were classified into 2 groups according to their lung capacity (FVC). Individuals with ALS were distributed into the spirometry group <50% (FVC < 50% predicted [n = 23]) and the spirometry group >50% (FVC > 50% predicted [n = 19]). Based on these parameters, the subjects with FVC < 50% predicted required ventilatory support. In neuromuscular diseases, for example, ALS, an FVC < 50% of the predicted value is an indication for ventilatory support with 2 pressure levels, even in the absence of respiratory symptoms.29-32

Study Size

To determine the sample size, previous knowledge was required based on the mean ± SD of normal RR intervals index in a study by Moreno et al.33 A sample size of 18 participants per group was stipulated by a hypothesis test (2-tailed), with a 5% level of significance and 80% power.

Statistical Analysis

Qualitative variables were analyzed by using the Fisher exact test or the chi-square test. The data are presented as box plots. We used the Shapiro-Wilk normality test to evaluate the distribution. For parametric distributions, we applied the unpaired Student t test; whereas, for nonparametric distributions, we applied the Mann-Whitney test. All findings with P < .05 were considered statistically significant. To measure the magnitude of the difference between the groups, the effect size was calculated by using Cohen’s d for significant differences. A large effect was considered for values > 0.9, moderate for values between 0.9 and 0.5, and small for values between 0.5 and 0.25.

Table 2. Frequency Domain Indices

<table>
<thead>
<tr>
<th>Index</th>
<th>FVC &gt; 50%, mean ± SD</th>
<th>FVC &lt; 50%, mean ± SD</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>LF, ms²</td>
<td>451.4 ± 513.0</td>
<td>296.6 ± 341.3</td>
<td>.29</td>
</tr>
<tr>
<td>HF, ms²</td>
<td>101 ± 104.8</td>
<td>88 ± 151.9</td>
<td>.41</td>
</tr>
<tr>
<td>LF, nu</td>
<td>80.4 ± 14.1</td>
<td>79.6 ± 11.7</td>
<td>.62</td>
</tr>
<tr>
<td>HF, nu</td>
<td>19.4 ± 14.1</td>
<td>20.2 ± 11.6</td>
<td>.55</td>
</tr>
<tr>
<td>LF/HF</td>
<td>477.8 ± 1,944.3</td>
<td>680.5 ± 3238.1</td>
<td>.48</td>
</tr>
</tbody>
</table>

LF = low frequency
HF = high frequency
nu = normalized unit

Results

Population Characteristics

Forty-two subjects diagnosed with motor neuron disease were evaluated: 47.5% were female and 52.5% were male subjects. The mean age was 57.7 y. The subjects had been diagnosed with motor neuron disease on average, 3 y earlier, and 17 subjects had bulbar disease characteristics. Data regarding age, sex, baseline heart rate, and oxygen saturation of the groups are shown in Table 1. We noted that the resting heart rate was higher, whereas SpO₂ was lower in the FVC <50% group. In relation to the lung capacity of the 2 groups, the FVC <50% group was mean ± SD, 38.06 ± 0.10%, whereas the FVC >50% group was mean ± SD, 73.54 ± 0.097%.

Linear Methods

We found no significant differences between the groups in the frequency domain analysis (Table 2).

Geometric Heart Rate Variability Indices

With regard to the geometric indices, we observed increased SD2 in the FVC >50% group (moderate effect size) (Table 3). Shown in Figure 1, is an example of a Poincaré plot from one subject in the FVC >50% group.
rate in the FVC rate variability modulation. We also found a higher heart
ventilatory support, which indicated reduced overall heart
rate in the FVC group (Fig. 1A) and one subject in the FVC <50% group (Fig.
1B). We noted less dispersion in the points of subjects
with reduced lung capacity. We performed a correlation
analysis to verify whether there is an association between
respiratory function and autonomic control of heart beats
in both groups (Table 4). There was no significant corre-
lation between FVC and heart rate variability in both groups.

Discussion

Our main finding indicated a significant decrease
(P < 0.01) in the SD2 index in the subjects who needed
ventilatory support, which indicated reduced overall heart
rate variability modulation. We also found a higher heart
rate in the FVC <50% group, which reinforced the im-
paired autonomic function; whereas \( S_{\text{pO}_2} \) was decreased in
the same group, which supported the worst pulmonary
condition in this group. In this sense, our findings sup-
ported that breathing modulates autonomic flow and con-
tributed to changes in heart rate variability.10

Spirometric measurements showed that, in most cases,
regardless of motor neuron failure, FVC decreased with
progression of the disease.34 The restrictive lung pattern in
ALS is characterized by an increased residual volume with
reduced total lung capacity.35 Moreover, inspiratory and
expiratory pressures are also reduced, and pressure
decreases according to the disease evolution. Previous stud-
ies reported that dyspnea is generally associated with di-
aphragmatic dysfunction, respiratory pulse, and a delay in
or abolition of diaphragm responses to stimulation of the
phrenic nerve or cortical stimulation.34,36 These results were
consistent with previous observations of predominantly
phrenic nerve neuropathologic abnormalities in subjects
with ALS.34,36

The presence of FVC < 50% predicted in motor neuron
disease requires bi-level ventilatory support, even in the
absence of respiratory symptoms.30,32 Verification of heart
rate variability is important to detect autonomic dysfunc-
tion disorders in patients with motor neuron disease to
avoid sudden death or other conditions that lead to a de-
crease in life expectancy.12 Autonomic dysfunction also
supports the concept that ALS is a degenerative disease of
multiple systems.6

Autonomic dysfunction (hyperhidrosis, alterations in sal-
ivary and gastrointestinal tract function) usually leads to
subclinical symptoms. Patients in more-advanced stages of
disease that requires ventilatory support often show auto-
nomic nervous system disorders. Such responses are marked
by fluctuations in blood pressure and heart rate, and can
occur in clinically stable patients with ventilatory support.
This response is known as the autonomic storm and is
characterized by a hypertensive crisis with systolic blood
pressure >250 mm Hg and tachycardia.6 These autonomic
symptoms are similar to baroreflex failure; they indicate a
poor prognosis and may result in sudden death.6 In this
situation, careful assessment and individual treatment are
strongly indicated, although adequate therapeutic ap-
proaches have not been well established. Autonomic dys-
function may be clinically obvious and devastating, espe-
cially at an advanced stage when ventilatory support is
required.

Our results showed no significant correlation between
heart rate variability and pulmonary capacity, which is
supported by previous studies (Table 4).15,37 We hypothe-
size that the absence of significant differences in the time
(mean ± SD of normal RR intervals, percentage of adja-
cent RR intervals with a difference of duration of >50 ms,
and the root of the square successive differences between
adjacent normal RR intervals) and frequency (high fre-
quency, low frequency, and low frequency to high fre-
quency ratio) domains is because geometric indices are
more sensitive at detecting changes. In this sense, non-
linear behavior is predominant in human systems due to
their complex and dynamic nature. Human physiology can-
not be accurately described by linear methods. Chaos the-
ory describes elements that manifest behaviors that are
extremely sensitive to initial conditions but that are diffi-
cult to repeat and are deterministic.20

The analysis of non-linear heart rate variability methods
has been gaining increasing interest, because there is evi-
dence that the mechanisms involved in cardiovascular reg-
ulation probably interact in a non-linear way. This new
sight into abnormalities in heart rate behavior under var-
ious conditions may provide additional prognostic infor-
mation when compared with traditional methods. How-
ever, its clinical utility requires further research with large
populations.38

Heart rate variability was analyzed from the RR inter-
vals through Poincaré plots based on non-linear dynam-
ics.23,24 The Poincaré plot is a 2-dimensional graphic rep-
resentation of the correlation between consecutive RR
intervals in which each interval is plotted against the next
interval.23,25,26 The Poincaré plot can be analyzed qualita-

Table 3. Geometric Indices of Heart Rate Variability

<table>
<thead>
<tr>
<th>Index</th>
<th>FVC &gt; 50%, mean ± SD</th>
<th>FVC &lt; 50%, mean ± SD</th>
<th>P</th>
<th>Cohen’s d</th>
</tr>
</thead>
<tbody>
<tr>
<td>RRtri</td>
<td>11.28 ± 6.01</td>
<td>8.7 ± 4.6</td>
<td>.08</td>
<td>ND</td>
</tr>
<tr>
<td>TINN</td>
<td>186.17 ± 84.03</td>
<td>144.13 ± 81.31</td>
<td>.14</td>
<td>ND</td>
</tr>
<tr>
<td>SD1</td>
<td>11.4 ± 6.0</td>
<td>9.9 ± 8.3</td>
<td>.15</td>
<td>ND</td>
</tr>
<tr>
<td>SD2</td>
<td>56.0 ± 22.6</td>
<td>38.6 ± 18.9</td>
<td>.01</td>
<td>0.83</td>
</tr>
</tbody>
</table>

RRtri = triangular index
TINN = triangular interpolation of RR intervals
SD1 = SD of instantaneous beat-by-beat variability
SD2 = long-term SD of continuous RR intervals
DECREASED HEART RATE VARIABILITY IN SUBJECTS WITH ALS

Fig. 1. Visual pattern of the Poincaré plot observed in one subject with FVC > 50% (A) and another with FVC < 50% (B).

Table 4. Correlation Between FVC and Heart Rate Variability in the Both Groups

<table>
<thead>
<tr>
<th>Parameter</th>
<th>FVC &gt; 50%</th>
<th></th>
<th></th>
<th>FVC &lt; 50%</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>RR interval</td>
<td>r</td>
<td>P</td>
<td></td>
<td>r</td>
<td>P</td>
<td></td>
</tr>
<tr>
<td>SDNN</td>
<td>−0.34</td>
<td>.15</td>
<td></td>
<td>−0.24</td>
<td>.26</td>
<td></td>
</tr>
<tr>
<td>Heart rate</td>
<td>−0.37</td>
<td>.11</td>
<td></td>
<td>0.18</td>
<td>.40</td>
<td></td>
</tr>
<tr>
<td>RMSSD</td>
<td>−0.06</td>
<td>.80</td>
<td></td>
<td>−0.21</td>
<td>.33</td>
<td></td>
</tr>
<tr>
<td>pNN50</td>
<td>−0.1</td>
<td>.66</td>
<td></td>
<td>−0.1</td>
<td>.64</td>
<td></td>
</tr>
<tr>
<td>RRtri</td>
<td>−0.26</td>
<td>.27</td>
<td></td>
<td>−0.29</td>
<td>.17</td>
<td></td>
</tr>
<tr>
<td>TINN</td>
<td>−0.33</td>
<td>.16</td>
<td></td>
<td>−0.24</td>
<td>.25</td>
<td></td>
</tr>
<tr>
<td>LF, ms²</td>
<td>−0.23</td>
<td>.34</td>
<td></td>
<td>−0.33</td>
<td>.11</td>
<td></td>
</tr>
<tr>
<td>HF, ms²</td>
<td>−0.15</td>
<td>.53</td>
<td></td>
<td>−0.14</td>
<td>.49</td>
<td></td>
</tr>
<tr>
<td>HF, nu</td>
<td>0.18</td>
<td>.44</td>
<td></td>
<td>0.05</td>
<td>.81</td>
<td></td>
</tr>
<tr>
<td>LF/HF</td>
<td>−0.06</td>
<td>.80</td>
<td></td>
<td>−0.15</td>
<td>.48</td>
<td></td>
</tr>
<tr>
<td>SD1</td>
<td>−0.06</td>
<td>.80</td>
<td></td>
<td>−0.2</td>
<td>.34</td>
<td></td>
</tr>
<tr>
<td>SD2</td>
<td>−0.39</td>
<td>.09</td>
<td></td>
<td>−0.22</td>
<td>.30</td>
<td></td>
</tr>
</tbody>
</table>

SDNN = SD of all NN intervals
RMSSD = square root of the mean of the sum of the squares of differences between adjacent NN intervals
pNN50 = percentage of adjacent RR intervals with a difference of duration of >50 ms
RRtri = triangular index
TINN = triangular interpolation of RR intervals
LF = low frequency
HF = high frequency
nu = normalized units
SD1 = SD of instantaneous beat-by-beat variability
SD2 = long-term SD of continuous RR intervals

These methods describe the complex fluctuations of the rhythm and can separate structures of non-linear behavior in a temporal series of heartbeats more adequately than linear methods. It has been speculated that heart rate variability analysis based on non-linear, dynamic methods can provide valuable information for physiologic interpretations of heart rate variability and assess the risk of sudden death. Pinto et al. reported that a low coefficient of variation of heart rate is a potential prognostic marker in ALS and other forms of motor neuron disease, and may lead to sudden death. Our results indicated a decrease in SD2 in subjects with greater respiratory impairment (heart rate variability < 50%), which required ventilatory support. The SD2 index represents both the vagal and sympathetic modulation of HR. Reduced SD2 was suggested to be an indicator of cardiac mortality and morbidity. Taken together, we reinforced that declined SD2 in patients with ALS and with lower pulmonary capacity was related to a worse prognosis.

For visual, qualitative, analysis of the Poincaré plot, it is clear that the group with FVC < 50% predicted presented a more-flattened form with less dispersion of RR intervals, which indicated decreased heart rate variability. In this sense, the plot supported the quantitative analysis of SD2, which showed reduced values and worse autonomic function in the ALS group with FVC < 50%. The theories of non-linear systems have been progressively applied to interpret, explain, and predict the behavior of biological phenomena. These parameters have been found to be good predictors of morbidity and mortality in the clinical context, despite the need for deeper scientific analysis, with expressive samples and prolonged follow-up. Such studies may be useful in research and treatment of heart disease.
Disturbances in cardiac autonomic control in patients with ALS with and without progressive bulbar palsy can influence survival and quality of life because this is associated with the onset of a hypertensive crisis, sudden cardiac death, and cardiovascular collapse, all leading to a decrease in life expectancy.\textsuperscript{13,14,44} Also worth mentioning is that the associations tested through Pearson and Spearman procedures were not statistically significant. We believe that patients with ALS and with lower FVC are influenced by physiologic variables not evaluated, including cytokines,\textsuperscript{45} antioxidant status,\textsuperscript{46} and α-amylase.

This study showed that heart rate variability may help to provide information regarding the lung function of patients with ALS. However, because lung function was measured only by spirometry, our results should be considered with caution. Heart rate variability changes induced by autonomic change could provide important data about the interaction between heart rate autonomic control and FVC in patients with ALS. Thus, we suggest that further studies of this procedure are needed. A limitation of the present study is that we did not monitor breathing pattern and tidal volume during heart rate recording. The above-mentioned variables mainly influence the high frequency band.\textsuperscript{22} During heart rate variability analysis, all the subjects were breathing spontaneously.

Conclusions

Our study showed that subjects with ALS and with reduced lung capacity presented impaired autonomic regulation of the heart rate. We suggest that impaired lung capacity was associated with autonomic dysfunction in subjects with ALS. This finding is important for the clinical team to plan therapies and treatment based on patients’ lung and autonomic function.

ACKNOWLEDGMENTS

We thank the reviewers for the anonymous constructive review on our manuscript and for providing interesting ideas for future projects.

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