Heart rate complexity analysis in Brugada syndrome during physical stress testing

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Abstract. Symptoms such as ventricular arrhythmias in Brugada syndrome (BS) typically occur at rest, especially during sleep, suggesting that the autonomic nervous system (ANS) function may be relevant in the arrhythmogenesis of the disease. The aim of this work was to assess the ANS response captured by a non-linear heart rate variability (HRV) measure in 69 patients diagnosed with BS, who underwent a standardized physical stress test. Heart rate complexity (HRC) was evaluated by the power-law scaling analysis ($\beta$ slope) during rest, exercise, recovery and rest post-recovery, in order to discriminate between symptomatic and asymptomatic BS patients. Symptomatic patients showed a significant reduction in HRC in comparison to asymptomatic subjects, after exertion ($p=0.015$); during the whole recovery period ($p=0.023$), and, in particular within the passive recovery phase ($p=0.025$), as well as during rest post-recovery ($p=0.022$). Based on these results, symptoms could be associated with a lower ANS complexity during the stress test stages where parasympathetic activity is predominant. Therefore, the proposed HRV indicators could be of help in the risk stratification of asymptomatic patients.

Keywords: Brugada syndrome, heart rate dynamics, autonomic nervous system, risk stratification, stress test

1. Introduction

Brugada syndrome (BS) is an inherited disorder characterized by a typical electrocardiographic pattern presenting a distinct ST-segment elevation in right precordial leads, associated with a high risk for unexpected sudden cardiac death (SCD) due to malignant ventricular arrhythmias in absence of any apparent structural
cardiopathy. Since its description in 1992 as a new cardiac disorder (Brugada & Brugada 1992), BS has raised a great interest because of its high incidence, especially in far eastern countries, and its relation to sudden death in young adults and, less frequently, in infants and children. It has been estimated that BS are responsible for 4-12% of the total amount of SCD and for 20% of SCD in patients without structural cardiopathy (Brugada, et al. 1998, Antzelevitch, et al. 2005). Although several indicators have been assessed for the prediction of cardiac events in BS, studies based on the largest clinical series only proved prior symptoms and spontaneous type-1 Brugada ECG pattern as consistent predictors of ventricular fibrillation (VF) episodes (Probst, et al. 2010). Thus, risk stratification in BS still remains controversial, especially for asymptomatic patients. However, by relating previous symptomatology with the occurrence of cardiac events in BS, the information provided from new indicators in symptomatic and asymptomatic patients can be compared and related to the prediction of VF events.

Since ventricular arrhythmias in BS typically occur at rest and mainly at night, it is commonly assumed that vagal activity may play a determinant role in the arrhythmogenesis of this disorder (Matsuo, et al. 1999, Kies, et al. 2004). Indeed, previous works, focused on the analysis of the autonomic nervous system (ANS) function using positron emission tomography, have identified sympathetic dysfunctions on BS patients (Kies et al. 2004, Wichter, et al. 2002, Bigi, et al. 2008, Paul, et al. 2011). Also, many previous works have studied the autonomic function of BS patients through HRV analysis (Krittayaphong, et al. 2003, Hermida, et al. 2003, Kostopoulou, et al. 2010, Nakazawa, et al. 2003, Makimoto, et al. 2010, Tokuyama, et al. 2014, Pierre, et al. 2007). However, these works are based on linear, time and frequency-domain methods, being unable to capture the complex, non-linear effects on heart rate (HR) regulation. We hypothesize that the evaluation of HR dynamics on BS patients through non-linear fractal methods may provide additional information for risk stratification in this population.

Moreover, the autonomic response can be better assessed by stimulating the ANS through controlled maneuvers such as physical exercise. Exertion provokes a sympathetic activity increase and a parasympathetic inhibition, leading to higher HR. Conversely, post-exercise cardiodeceleration is adjusted by a continued increase in parasympathetic activity (Imai, et al. 1994), as well as a progressive sympathetic withdrawal (Savin, et al. 1982). Since traditional HR analysis at rest commonly fails to distinguish between symptomatic and asymptomatic BS patients, we explored HR changes during physical stress testing. Indeed, (Makimoto et al. 2010) published the first work reporting the interest of exercise to predict ventricular arrhythmias in BS patients, focusing on recovery after exertion.

Approaches using the power-law scaling exponent ($\beta$) obtained from the regression line of Power Spectral Density (PSD) vs. frequency plot on a bi-logarithmic scale have shown decreased complexity in patients with cardiovascular disorders and sudden cardiac death events (Huikuri, et al. 2000, Huikuri, et al. 1998, Lombardi, et al. 2001, Gomis,
et al. 2006, Maestri, et al. 2007, Tapanainen, et al. 2002). However, we are not aware of any study that has analyzed the HR dynamics’ complexity (HRC) of symptomatic and asymptomatic BS patients under conditions of physical activity and recovery.

Thus, the aim of this manuscript was to compare, according to symptomatic status, the autonomic cardiac control captured by this non-linear complexity indicator from RR series, in 69 BS patients, for each phase of a standardized physical stress test. The role of complexity assessment would be helpful in risk stratification of asymptomatic BS patients for whom the decision to implant an Implantable Cardioverter Defibrillator (ICD) based on the benefit/risk for the patient is particularly complex.

2. Methods

2.1. Study population

Sixty-nine Brugada syndrome patients underwent a physical exercise stress test while continuous ECG recordings were acquired. They were enrolled in 6 French centres: Rennes, Saint Pierre de la Réunion, Nantes, Bordeaux, Brest and La Rochelle. After approval by the ethics committee of each centre, all participants provided informed consent to take part in the study, which was performed in accordance with the clinical research and ethics recommendations. Patients’ age ranged from 19 to 73 years old (45.06 ± 12.79 years old) and 78.3% were males. Table 1 summarizes participants’ baseline characteristics, where no significant differences in age and gender between patients included in the symptomatic and asymptomatic groups can be noted. Sixteen patients had the following documented symptoms: syncope (56.25%), cardiac arrest (25%), dizziness (18.75%) and, less frequently, palpitations (12.50%) and nocturnal convulsions (6.25%).

ICD implantation had been performed in 15 of 53 (28.3%) asymptomatic patients, based on a positive EPS (Electrophysiological Study) test, whereas all symptomatic patients were ICD carriers. Since no relevant cardiac episodes occurred during the test, these devices caused no significant effects on the recordings. Among 54 patients (13 were symptomatic) in whom genetic screening was performed, an SCN5A mutation was found in 19 (35.2%), from whom 5 were symptomatic.

Table 1. Patients’ baseline characteristics.

<table>
<thead>
<tr>
<th></th>
<th>All patients (n=69)</th>
<th>Symptomatic (n=16)</th>
<th>Asymptomatic (n=53)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years old</td>
<td>45.06 ± 12.79</td>
<td>45.88 ± 15.31</td>
<td>44.81 ± 12.09</td>
<td>0.837</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>54 (78.3%)</td>
<td>13 (81.3%)</td>
<td>41 (77.4%)</td>
<td>0.743</td>
</tr>
<tr>
<td>ICD implantation, n (%)</td>
<td>31 (44.9%)</td>
<td>16 (100%)</td>
<td>15 (28.3%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Presence of SCN5A mutation, n (%)</td>
<td>19 (35.2%)</td>
<td>5 (38.5%)</td>
<td>14 (34.2%)</td>
<td>0.656</td>
</tr>
</tbody>
</table>
2.2. Signal acquisition and test

Patients participated in a clinical protocol recommended by the American Heart Association (Gibbons, et al. 2002) described as triangular stress test since the load was increased until it reached the patient’s maximum potential. It was performed on a cyclo ergometer (Ergoline 900 Egamed, Piestany, Slovakia) and divided in the following phases:

- Resting phase: supine rest for 8 minutes and rest sitting on the bike for 2 more minutes.
- Exercise phase:
  - Warm-up phase: for men, initial load of 50 watts (W); for women, initial load of 30 W, both for 2 minutes.
  - Incremental exercise phase: for men, initial load of 80 W for 2 minutes and then incrementing 20 W every 2 minutes; for women, initial load of 50 W increasing load 20 W every 2 minutes.
- After exercise phase:
  - Recovery phase:
    * Active recovery phase: for men, fixed load of 50 W; for women, fixed load of 30 W, both for 3 minutes.
    * Passive recovery phase: total cessation of effort for 3 minutes.
  - Rest post-recovery phase: supine rest until reaching 50 minutes of total recording, never exceeding 30 minutes of resting after recovery.

The purpose of the test was to reach at least 80% of the theoretical maximum heart rate, defined by the formula $MHR = 220 - age$ (Fox & Haskell 1968).

The standard 12-lead ECG recording was collected and analyzed by the central board (Centre Hospitalier Universitaire de Rennes). It was acquired with the Holter monitor (ELA medical, Sorin Group, Le Plessis Robinson, France) and continuously recorded at rest (from 10 minutes before warm-up phase beginning), during incremental exercise, recovery (6 minutes) and rest post-recovery. Since each test differed in the incremental exercise and rest post-recovery periods, the duration for these phases was fixed to 7 and 10 minutes, respectively, corresponding to the smallest duration found in the clinical series. Sampling frequency was set at 1000 Hz. Heart rate was continuously displayed and blood pressure measured with the sphygmomanometric method at rest, every two minutes during exercise, maximum effort and recovery.

RR series were obtained using a noise-robust wavelet-based algorithm for QRS complex detection and subsequent R-wave peak location (Dumont, et al. 2010). Since the result is a non-uniformly sampled RR series, a cubic splines interpolation was applied to signals in order to obtain regularly sampled data at a rate of 4 Hz.
2.3. Power-law Scaling Analysis

The RR series in healthy subjects fluctuates in a complex way with a non-uniform pattern that has been associated with fractal behavior and, on a broadband spectrum, with a 1/f-like distribution (inverse power law). The $\beta$ index measures the power-law behavior as the slope of the regression line calculated by the least-squares method on the linear portion of PSD against frequency plots on a bi-logarithmic scale. Several studies have employed the frequency band 0.0001-0.01 Hz to analyze long-term fluctuations when evaluating sudden cardiac death and myocardial infarction (Huikuri et al. 1998, Tapanainen et al. 2002). However, for short-term recordings, this technique has also been applied using the 0.003-0.1 Hz band (Lombardi, et al. 2000).

PSD was estimated using the Burg method (Burg 1975) with an optimized order for each signal, based on the Broersen’s combined information criterion (Broersen 2000), to obtain the autoregressive model parameters. Since we analyzed short segments of the RR series, the scaling exponent $\beta$ was extracted from the frequency band 0.003-0.1 Hz (-2.5 to -1 on a logarithmic scale).

2.4. Statistical Analysis

Comparisons between symptomatic and asymptomatic patients during different phases of the test were evaluated by Mann-Whitney U non-parametric tests. The analysis was made using the commercially available statistical program SPSS 21.0 (IBM, Chicago, USA) and setting the level of significance at $p < 0.05$.

Binary logistic regression models were applied to variables that significantly differed between groups in order to predict the probability of experiencing symptoms, as a function of HRC measures.

3. Results

Table 2 summarizes the $p$-values obtained when comparing the $\beta$ slope analyzed across
the different phases of the test between symptomatic and asymptomatic BS patients, as well as the R-squared (R²) as a fitting quality measure between the PSD and its fitted model associated with each phase. In general, linear models reached an acceptable goodness of fit, where R² > 0.86. The lowest fitting qualities were found for exercise and after recovery phases, which were then analyzed in smaller segments with higher R², always greater than 0.92.

**Table 2.** P-values and R² obtained when comparing symptomatic and asymptomatic patients along the test.

<table>
<thead>
<tr>
<th></th>
<th>β</th>
<th>R²</th>
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<tbody>
<tr>
<td>Resting</td>
<td>0.320</td>
<td>0.91</td>
</tr>
<tr>
<td>Exercise</td>
<td>0.418</td>
<td>0.86</td>
</tr>
<tr>
<td>Warm-up</td>
<td>0.135</td>
<td>0.92</td>
</tr>
<tr>
<td>Incremental exercise</td>
<td>0.065</td>
<td>0.94</td>
</tr>
<tr>
<td>After exercise</td>
<td>0.015*</td>
<td>0.94</td>
</tr>
<tr>
<td>Recovery</td>
<td>0.023*</td>
<td>0.95</td>
</tr>
<tr>
<td>Active recovery</td>
<td>0.118</td>
<td>0.96</td>
</tr>
<tr>
<td>Passive recovery</td>
<td>0.025*</td>
<td>0.96</td>
</tr>
<tr>
<td>Rest post-recovery</td>
<td>0.022*</td>
<td>0.89</td>
</tr>
</tbody>
</table>

*p < 0.05, Mann-Whitney U test.

Figure 2 displays an example of fit for a symptomatic and an asymptomatic patient after exertion, so as to represent the fitting quality, as well as the β slope decay experienced by the symptomatic case with respect to the asymptomatic patient.

**Figure 2.** Linear fit for β extraction after exercise in a symptomatic (left) and in an asymptomatic (right) BS patient.
Symptomatic patients showed a significant reduction in HRC, as estimated by the $\beta$ slope, in comparison to asymptomatic subjects after exertion ($p=0.015$); during recovery ($p=0.023$), and more specifically during the passive recovery phase ($p=0.025$), and at rest post-recovery ($p=0.022$). Figure 3 reflects how symptomatic patients showed significantly decreased values in both recovery and post-recovery phases, with respect to asymptomatic subjects.

**Figure 3.** Group average results for $\beta$ during resting, exercise, recovery and post-recovery phases; for symptomatic and asymptomatic BS patients (*$p<0.05$).

Table 3 collects the odds ratio (OR), 95% confidence intervals (CI) and $p$-values associated with each univariate model applied to variables that significantly differed between groups.

<table>
<thead>
<tr>
<th></th>
<th>Odds Ratio</th>
<th>95% CI</th>
<th>$p$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta$, recovery</td>
<td>0.180</td>
<td>0.028 - 1.146</td>
<td>0.069</td>
</tr>
<tr>
<td>$\beta$, passive recovery</td>
<td>0.230</td>
<td>0.029 - 1.812</td>
<td>0.163</td>
</tr>
<tr>
<td>$\beta$, rest post-recovery</td>
<td>0.097</td>
<td>0.012 - 0.803</td>
<td>0.030*</td>
</tr>
<tr>
<td>$\beta$, after exercise</td>
<td>0.189</td>
<td>0.036 - 0.992</td>
<td>0.049*</td>
</tr>
</tbody>
</table>

* $p < 0.05$, Mann-Whitney U test.

During the rest post-recovery phase, $\beta$ turned to be an independent predictor in univariate analysis with a 9.7% ($p=0.030$) decreased risk per one point increase. After
exercise, $\beta$ was also a significant predictor of symptoms, decreasing risk by a 18.9% per each point increase ($p=0.049$).

Finally, table 4 summarizes the mean ± standard deviation values for symptomatic and asymptomatic patients and summarizes $p$-values obtained in each significant case, showing higher values of HRC for asymptomatic BS patients.

Table 4. Mean ± standard deviation, for symptomatic and asymptomatic patients, and associated $p$-values of significant variables.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Symptomatic (n=16)</th>
<th>Asymptomatic (n=53)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta$, recovery</td>
<td>-1.866 ± 0.258</td>
<td>-1.685 ± 0.356</td>
<td>0.023</td>
</tr>
<tr>
<td>$\beta$, passive recovery</td>
<td>-1.773 ± 0.292</td>
<td>-1.655 ± 0.288</td>
<td>0.025</td>
</tr>
<tr>
<td>$\beta$, rest post-recovery</td>
<td>-1.853 ± 0.207</td>
<td>-1.651 ± 0.334</td>
<td>0.022</td>
</tr>
<tr>
<td>$\beta$, after exercise</td>
<td>-1.706 ± 0.335</td>
<td>-1.468 ± 0.423</td>
<td>0.015</td>
</tr>
</tbody>
</table>

4. Discussion

In this study, non-linear HRV indicators were compared in 69 BS subjects who took part in a standardized physical stress test, in order to discriminate between symptomatic and asymptomatic patients. Although previous works have already analyzed the autonomic response of BS patients, this study presents, to our knowledge, the first evaluation of HR dynamics captured by short-term power-law scaling analysis, under conditions of physical activity and recovery.

This HRV marker has been employed to assess HRC in other cardiac disorders, such as chronic heart failure or transient myocardial ischemia, demonstrating decreased values of complexity in worse cardiac health conditions (Huikuri et al. 2000, Huikuri et al. 1998, Lombardi et al. 2001, Gomis et al. 2006, Maestri et al. 2007, Tapanainen et al. 2002).

The results obtained show a significant reduction in $\beta$ after exertion, during recovery and rest post-recovery in symptomatic patients, with respect to asymptomatic patients. Therefore, subjects with higher risk showed lower complexity in HR dynamics, as already published for other cardiac pathologies.

Those statistically significant variables were then included in univariate logistic regression models in order to estimate their discrimination power. After recovery, $\beta$ turned to be an independent predictor of symptomatology, supporting the idea that decreased complexity could be related to a greater risk in Brugada syndrome. After exercise, $\beta$ was also a significant predictor, relating a decreased power-law scaling exponent to a worse prognosis.

It should be noted that although other indicators described in the literature assessing HRV non-linear properties were analyzed, differences between symptomatic and asymptomatic BS patients were not as significant as those captured by the $\beta$ slope.
Despite DFA, the Hurst exponent, Sample Entropy and Higuchi’s Fractal Dimension capture similar HRV properties (Acharya, et al. 2006, Baumert, et al. 2007), in our clinical series, only the $\beta$ slope demonstrated a potential impact on the prediction of risk in BS patients.

The autonomic nervous system is governed by complex interactions resulting from feedback loops of non-linear systems, that enable the organism to adapt to abnormal physiological conditions, such as metabolic variations, stress or disease. According to previous works, a reduction in the complexity of physiological control systems occurs with age and disease (Goldberger, et al. 2002, Rabinovich & Abarbanel 1998, Magrans, et al. 2010). Fractal power-law index $\beta$ decreased significantly in symptomatic patients after exertion and recovery, when parasympathetic activity is predominant. Indeed, (Makimoto et al. 2010) analyzed the parasympathetic reactivation in BS patients by computing heart rate recovery (HRR) as the heart rate decay in the first minute after peak exercise cessation, concluding that a higher parasympathetic activity was related to the occurrence of cardiac events in BS. Thus, recovery stages seem to unmask significant autonomic differences related to symptomatology, also captured by the non-linear HRV indicators here presented.

The results found in our work indicate a loss of cardiac rhythm complexity after exertion and recovery in symptomatic BS patients. This may be explained by the fact that a decreased organism’s adaptability leads to a reduced complexity of the underlying biological control systems (Smith, et al. 2005). Moreover, this altered autonomic cardiac system dynamics when vagal tone is predominant confirms previous works that highlight the relevant role of autonomic imbalance in the genesis of life-threatening cardiac arrhythmias in BS (Matsuo et al. 1999, Kies et al. 2004). Indeed, polymorphic ventricular arrhythmias (PVA) and sudden cardiac death (SCD) mainly occur at rest and especially during sleep, when parasympathetic activity is predominant.

Data on cardiac ANS analyzed by positron emission tomography has confirmed that BS patients display some autonomic dysfunction, with increased presynaptic norepinephrine (NE) recycling and thus a reduction in the concentration of NE at the synaptic cleft. This imbalance is thought to promote arrhythmogenicity by decreasing intracellular levels of 3-5-cyclic adenosine monophosphate (cAMP) and thus disrupting the subsequent signaling pathway with potential implication for arrhythmogenesis (Wichter et al. 2002, Kies et al. 2004). This may lead to an enlarged transmural electrical dispersion that results in an increased probability of ventricular arrhythmias. In addition, the autonomic imbalance might be aggravated with the presence of physiological downregulation of adrenergic activity, justifying the occurrence of PVA and SCD at rest or during sleep (Priori, et al. 2000, Yan & Antzelevitch 1999). Other studies based on the full stomach test demonstrated that vagal stimulation induced by a large meal not only increased distinctive Brugada-like ECG changes but also lead to prognostic tendencies, since patients with a positive full stomach test were more frequently related to life-threatening events than patients with a negative test result (Miyazaki, et al. 1996, Chalvidan, et al. 2000).
Thus, findings on the present study further support this hypothesis and provide additional evidence for the role of autonomic imbalance in the pathophysiology of the disease. Although we did not find significant differences between groups in metrics characterizing the parasympathetic component that directly relate the loss of complexity with an increase in vagal tone, this study highlights the importance of physical stress testing in order to unmask relevant autonomic changes that could not be captured at rest, before the test.

Therefore, measurement of HRV not only by conventional but also by new non-linear methods during physical stress testing could be a helpful tool for documenting neural regulation alteration related to symptoms, which can have a potential impact on the progress of risk stratification in BS patients.

However, a relevant limitation of the study is that the clinical value of the analyzed indicators can only be proved if a close relationship between HRV variations and ventricular arrhythmias is identified. Here, symptomatic patients had documented symptoms but since no cardiac events of ventricular origin were induced during the study; we cannot directly relate HRC alterations in symptomatic patients to a greater risk for suffering symptoms.

5. Conclusion

We compared non-linear HRV indicators in 69 BS patients who took part in a standardized physical stress test in order to distinguish between symptomatic and asymptomatic groups.

Statistically significant differences were found during and after recovery, showing lower HR dynamics’ complexity in symptomatic patients under conditions of recovery and rest after recovery.

Although this study is based on a relatively small population of 69 subjects and, thus, conclusions on symptomatic status predictions cannot be inferred, the identified differences between symptomatic and asymptomatic BS patients indicate relevant tendencies of clinical importance. Therefore, significant HRV results could be of potential help in risk stratification of asymptomatic patients for whom the decision to implant a cardioverter defibrillator is still contentious.

Acknowledgments

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Heart rate complexity in Brugada syndrome during stress


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