Noninvasive Electrocardiographic Imaging Diagnosis Of Brugada Syndrome

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Abstract

Brugada syndrome (BrS) is a well-known inherited heart disease with a high risk of a sudden cardiac death (SCD) in young male patients. Thy only way to make a diagnosis is standard ECG. The spectrum of repolarization changes between normal ECG and BrS diagnostic criteria, are called BrS-like ECGs. They occur due to nonspecific right bundle branch conduction disorders. Patients with this abnormal ECG are always a challenge for a doctor – is there a potentially fatal diagnosis or variant of normal ECG.

1. Introduction

Electrocardiographic phenomena of right bundle branch block (RBBB) with ST-segment elevation was first described in 1953 in a little boy with recurrent syncope [1]. His smaller sister with similar ECG changes and symptoms died at the age of 2 despite implanted pacemaker and antiarrhythmic drug therapy. In 1992 brothers Pedro and Joseph Brugada based on 8 clinical cases proposed new syndrome of ECG changes associated with sudden cardiac death [2]. At 2002, by group of experts [3], there were defined three types of BrS, depends on ST-T segment morphology, amplitude of elevation. Second consensus conference [4] conclude that only 1 type correlates with life threatening arrhythmias, while other types are nondiagnostic. All suspicious but nondiagnostic variants were called BrS-like and required further investigation.

In 1998 genetic tests revealed mutation in SCN5A gene [5]. Later 10 more genes were associated with BrS phenotype. Mostly they encode sodium channels, but there are some genes, encoded calcium and potassium channels as well. BrS is an autosomal dominant disease with incomplete penetrance, which means that even carriers of BrS genotype can never show typical phenotype and do not have the disease. Genetic tests go positive for causative mutation in about one third of all patients. In vast majority of patients, genetic cause of BrS stays unknown. Thus, clinical evaluation always goes first.

Patients with once registered 1 type BrS could have nondiagnostic phenotype on other day. Physiological conditions (autonomic tone) are thought to be the buffer between genotype and phenotype. There is a number of reports describing BrS ECG pattern appearance in people under different physiological conditions, such as exercise, fever, drug administration. Drug test with sodium channel blocker enables to unmask BrS in some patients. Its diagnostic value depends on drug used, mutation, autonomic tone, lead position [6]. Best profile of specificity and sensitivity showed ajmaline [7] and flecainide [8], in which positive predictive value was reported more than 90% and negative predictive value from 30 to 83% [9]. Other drugs with sodium blocking activity could reveal BrS phenotype. The full list of them is described on the website brugadadrugs.org. All patients with BrS should avoid these medicine, because sodium channel blockage may lead not only to ECG changes, but also to clinical manifestation with ventricular arrhythmias.

Mechanism of arrhythmias and ECG development many years was the matter of debate. Too independent groups after series of experiments in silico, reported opposite hypothesis: repolarization disorder hypothesis [10] and depolarization disorder hypothesis [11]. Real breakthrough was made after cardiac catheterization and electrophysiological study in BrS patient with recurrent ventricular fibrillation episodes. So called electrophysiological substrate was found [12,13]. The area of abnormal electrogramms was located in anterior aspect of right ventricular outflow tract (RVOT). The abnormalities of electrogramms were the following: low voltage, prolonged duration and fractionated late potentials beyond ventricular complex. The same characteristics of electrogramms in RVOT in different patients with BrS enables to identify the substrate, and later, to eliminate this substrate with radiofrequency ablation. The similar picture of arrhythmogenic substrate was obtained using noninvasive electrocardiographic imaging [14, 15]. They found low amplitude of electrogramms, fragmentation, slow discontinuous conduction, delayed RVOT activation and ST segment elevation and inverted T wave of reconstructed electrogramms.

The compromise between two confronting groups was achieved by providing confirmation of both repolarization and depolarization disorders. The huge step forward was described with eliminating of arrhythmogenic substrate and normalization of ECG and absence of ventricular arrhythmias in patient with
repetitive electrical storms (recurrence of VF more than 3 times per day) [16].

Prevalence in Asia of BrS in adults is relatively higher than in West population (0.15% vs 0.02%). It seems like there is some ethnical modulation of gene polymorphism, causing the disease [17]. Despite the low prevalence, BrS is responsible up to 20% of all sudden cardiac deaths in young population, with absence of structural heart disease [18]. The symptoms of the disease are the symptoms of ventricular arrhythmias. In case of nonsustained ventricular tachycardia or ventricular fibrillation, patient may develop dizziness, palpitations and syncope. In case of sustained VF patient develop cardiac arrest. Syncpe and spontaneous type 1 ECG pattern are risk factors of SCD, according to current guidelines. Cardioverter-defibrillator should be implanted in these patients for primary prevention reasons. We have conflicting data on other proposed risk factors. Risk evaluation and management of BrS patients remains the matter of debate. BrS clinical manifestation occurs on the forth decade of life, in healthy individuals. In the absence of symptoms, prognosis is favorable and annual death rate is expected to be less than 0.5% [17]. In case of syncope and documented VF episodes, annual recurrence rate is estimated to be 11% [19]. Cardiac arrest could be the first sign of BrS. Thereby diagnostic and risk evaluation are still needs further investigations.

2. Materials and Methods

We examined 10 consistent patients with BrS-like ECGs. It was a prospective 2 center study, lasting for 3 years (Petrovsky National Research Centre of Surgery, Moscow, Russia and Mannheim University Clinic, Mannheim, Germany). All patients were asymptomatic. One with family history of BrS and positive genetic screening for SCN5A mutation (c.1233del). Predominantly males (n=8), age median was 37±11.4 y. The aim of our study was to determine the difference between RBBB and BrS in patients with Brugada-like ECG noninvasively.

We perform noninvasive ECGI using AMYCARD system (EP Solution SA, Yverdon-les-Bains, Switzerland), which allows to reconstruct epicardial and endocardial unipolar electrograms noninvasively, using heart/torso anatomy from computing tomography of cardiac magnetic resonance imaging and body surface ECG of 224 channels.

3. Results

Standard 12-lead and polyECG recordings were made at the baseline conditions and during drug challenge, aimed to unmask typical BrS ECG pattern. In half (n=5) of our group test was positive and obtained ECG changes met diagnostic criteria of BrS, while in others sodium channel blockage does not impact on ECG.

![Picture 1. Drug-induced 1 type BrS ECG: before and after drug challenge.](image1)

![Picture 2. Morphology of epicardial and endocardial electrograms in drug-induced type 1 patient: A and B – before drug challenge; C and D – after drug challenge.](image2)

While baseline standard ECG looks very similar in all 10 patients (ST segment elevation < 2 mV with a following negative T wave), morphology of local unipolar electrograms in right ventricle outflow tract (RVOT) differ a lot. At baseline condition patients with positive drug test, have morphological anomalies inherent to BrS patients in RVOT. Such as ST segment elevation >2 mV, fragmentation (more than three distinct peaks) and low voltage. Median (Me) of amplitude of ST segment elevation on the epicardium was 2.3 mV in BrS patients versus 0.1 mV in patients with negative drug test (p=0.008), and 2.6 mV vs 0.2 mV on endocardium. Fragmentation of electrograms was registered in all BrS patients on the epicardial surface (n=5). The amplitude of electrograms was lower on the epicardium surface in BrS patients Me=0.1 mV vs Me=3.6 mV in healthy individuals, but the difference was not statistically significant (table 1).
4. Discussion

ICD implantation for a long time stays the golden standard of care of BrS patients [20]. In presence of syncopes or after aborted cardiac arrest, this was the only way of SCD prevention. Antiarrhythmic drugs do not show any effect on ventricular arrhythmias in these patients.


Discussing with patient his or her prognosis and death rate is a hard talk to make. Especially in situation, when patient have spontaneous ECG pattern, but no risk factors. This is a grey zone, whether SCD occurs and when would it happened? Some of patients prefers double confidence and forces the doctor to the unnecessary ICD implantation. Device is technically lifesaving, but quality of life is compromised with inappropriate shocks.

But nowadays we have promising experience of arrhythmia substrate elimination using invasive or noninvasive navigation. This approach improves quality of life, and practically healing patients. The question is, should we ablate asymptomatic patients or not. Is it safe enough to perform cardiac catheterization and epicardial ablation in people with annual risk of SCD 0.5%?

The other question is diagnostic evaluation. Can we define patients with BrS-like ECG between healthy individuals and sick people? If we can, are there any additional factors for risk stratification?

With our study we find the answer for one of these questions. Using ECGI alone or in combination with standard drug test we can improve diagnostics of BrS.

5. Conclusion

All BrS patients has zone of abnormal electrograms in epicardial and endocardial aspects of RVOT, irrespective to the current ECG picture. Coved ST-segment elevation of rEGs, magnitude ≥ 2 mV might be strong independent predictor of BrS in patients with nondiagnostic ECG

References

Table 1. Characteristics of electrogramms in RVOT in patients with BrS like ECG

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Heart surface</th>
<th>BrS-like ECG, n=10</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Positive drug test, n=5</td>
<td>Negative drug test, n=5</td>
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<tr>
<td></td>
<td></td>
<td>Me (Q1-Q3)</td>
<td>Min-Max</td>
</tr>
<tr>
<td>Surface area of ST segment elevation, mm²</td>
<td>Epi</td>
<td>1208 (1125-1643)</td>
<td>0-2683</td>
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<tr>
<td></td>
<td>Endo</td>
<td>1935 (1087-3483)</td>
<td>0-4582</td>
</tr>
<tr>
<td>ST segment elevation amplitude, mV</td>
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<td>0,1-3,7</td>
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<td>Endo</td>
<td>2,6 (2,1-2,7)</td>
<td>0,4-3,6</td>
</tr>
<tr>
<td>Surface area of electrogramms fragmentation, mm²</td>
<td>Epi</td>
<td>1422 (1232-1554)</td>
<td>1023-1607</td>
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<td></td>
<td>Endo</td>
<td>0 (0-204)</td>
<td>0-2386</td>
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<tr>
<td>Mean voltage of electrogramms, mV</td>
<td>Epi</td>
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<td>0,1-0,7</td>
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<td>5,6 (2,3-6,5)</td>
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