Case Report

Pacing in the right atrium may induce atrial dyssynchrony and subsequent AF [1]. The percentage of atrial pacing can lead to hemodynamic alterations, and changes in the shape and velocity of intra-atrial, interatrial, and atrioventricular conduction, as well as alter cellular properties. An association between %AP and AF may be possible because of differences in baseline variables. Because all of the patients in this study had this feature turned on, we are not able to confirm this possibility. Pacing strategies targeting the intra- and interatrial conduction tracts [2,3] and algorithms aimed at minimizing atrial pacing should be further examined.

In our case report, we presented a female patient, 64 years old, complaining of syncope, pre-syncope, and fatigue on usual efforts. It has been ablated to atrium-ventricular node reentrant tachycardia (AVNRT) and is in use of sotalol hydrochloride 160 mg twice daily for the treatment of ventricular premature beats. She has hypertension, type 2 diabetes mellitus, dyslipidemia, treated with losartan 100 mg daily, metformin hydrochloride 850 mg twice a day, and rosuvastatin 10 mg daily. She also has negative myocardial scintigraphy for ischemia, normal carotid Doppler and echocardiogram, normal 24-hour Holter monitoring showing heart rate (minimum, average and maximum) of 58-78-122 bpm, sinus rhythm, 6633 polymorphic ventricular premature beats, 2 episodes of non-sustained ventricular tachycardia composed by 3 beats fast as 128 bpm. Cardiac nuclear magnetic resonance image showing left ventricular ejection fraction 60%, an end-diastolic diameter of 48 mm, and without necrosis or ischemia. She underwent electrophysiologic study that showed dysfunction of the distal electrical conduction, with HV interval of 78 ms. Faced with this situation, the dual chamber pacemaker was indicated.

During the positioning of the right ventricular lead in the middle interventricular septum, no change was noted, being fixed and having the following parameters: R wave = 12.0 mV, impedance = 620 Ω, and threshold: 0.5 V@0.5 ms. However, during the right atrial lead positioning observed by a posterior-anterior view (Figure 1) no acceptable sense and threshold were registered in the several positions (the “bald” right atrium): presenting P wave measures < 0.5 mV and thresholds > 3.5 V@1.0 ms. Given this problem, we opted to insert the atrial lead into the coronary sinus, positioning it in the in the highest position possible, fixing it in this position and measuring the left atrial parameters through the coronary sinus observed by a posterior-anterior view and left-anterior oblique view (Figure 2 A and B), which were the following: P wave = 3.2 mV, impedance = 850 Ω, and threshold: 0.9 V@0.5 ms. The leads were connected to DDDR generator, and the incision was sutured and closed in layers. The pacemaker was finally programmed with the following atrial and ventricular parameters, respectively: bipolar sense presenting P wave = 3.4 mV, impedance = 670 Ω, left atrial threshold = 0.9 V@0.5 ms, and unipolar output = 2.5V@0.5 ms; bipolar sense presenting R wave = 14.2 mV, impedance = 590 Ω, ventricular threshold = 0.4 V@0.5 ms, and and bipolar output = 2.0V@0.5 ms.

In this case, we can conclude that despite the difficulties encountered to position the lead in the right atrium, we found an alternative way to sense and pacing the left atrium, by the intra coronary sinus approach.

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References


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