Idiopathic intrafascicular reentrant left ventricular tachycardia in an elite cyclist athlete

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Abstract
A 32 year-old Caucasian male, an elite athlete, was admitted to the emergency department because of a sudden onset of palpitations which had lasted more than 12 hours and were associated with chest discomfort. He had a two-year history of recurrent stress-induced palpitations. He denied either episodes of syncope or any family history of sudden death. Physical examination was normal. He had no evidence of structural heart disease. The electrocardiography (ECG) documented during the event supported the diagnosis of idiopathic reentrant left ventricular tachycardia. Ventricular tachycardia ablation was successful. This case demonstrates that a careful physical examination and correct ECG diagnosis can lead to an appropriate arrhythmia management. (Cardiol J 2009; 16, 6: 564–567)

Key words: idiopathic left ventricular tachycardia, athlete’s heart apparent structural normal heart

Case report

Identification
A 32 year-old Caucasian Brazilian male, elite athlete (cyclist) in ectomorph biotype.

Complaint and duration
Regular heart palpitations, associated with atypical chest pain, began two days prior to the visit to the emergency department. Our patient complained of recurrent rapid palpitations triggered by exertion (walking short distances or climbing stairs), associated with dizziness and diffuse chest discomfort, described as “burning” with moderate intensity, without radiation. He denied nausea, vomiting or diaphoresis. The symptoms were intermittent; they stopped for hours and always recurred with exertion. At the beginning, the palpitations lasted from 30 minutes to one hour. But at the time of presentation in the emergency room, they had already lasted for 12 hours without interruption.

Our patient reported that progressive symptoms of palpitations started two years ago. They occurred always after training and lasted for around 20 minutes. The symptoms were always less intense than the current ones. He denied syncope, dizziness, or any symptom of low output during these episodes.

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Received: 7.02.2009 Accepted: 11.02.2009
Personal history: Competitive athlete, cyclist, covering on average 400 km per week; training from three to four hours per day. He denies any cardiovascular risk factor.

Family history: Noncontributory.

Physical examination: Blood pressure 110/70 mm Hg, heart rate 150 bpm, respiratory rate 26 rpm, jugular venous pulse presents with irregular cannon “a” waves and absence of stasis.

Pulmonary system: Lungs clear to auscultation bilaterally.

Cardiovascular system: The ictus cordis was palpable at the fifth left intercostal space, on midclavicular line. The first heart sound displayed a varying intensity. The second sound was normal. No murmurs.

Abdomen: Soft, non-tender, non-distended. Bowel sounds present. No hepatosplenomegaly.

Limbs: Pulses present on all four extremities, good peripheral perfusion, without edema.

The electrocardiography (ECG) recorded during an episode of palpitation is shown in Figures 1–3.

**Figure 1.** Electrocardiography recorded during palpitations showing following features: the ectopic beats showing right bundle branch block pattern with QRS axis shifted to right and extreme left axis deviation; rS pattern is shown in leads I, and aVL and qR in lead III, indicating infranascicular re-entrant left ventricular tachycardia re-entry was facilitated via left anterior fascicle; the QRS is relatively narrow or borderline-broad (between 120 ms and 140 ms) with RS interval (from the beginning of R wave until the nadir of S wave) < 80 ms in all precordial leads. In V1 biphasic pattern (qR) and notched R wave showing left R peak (second) taller than the right one (first): the ‘rabbit ear clue’ sign in V1 [4], and RS ratio < 1 in V6.
Transthoracic echocardiogram and cineventriculography were normal. The ventricular repolarization abnormalities, shown in the ECG immediately after the event (Fig. 4), and the complaints of chest pain, led to a cineangiography.

**Electrophysiology study**

**Programmed ventricular stimulation (PVS)**
During PVS, there was induction of tachycardia of wide QRS, right bundle branch block (RBBB) pattern, with upward axis of 420 ms. Absence of ventricle-atrial conduction.

**Percutaneous radiofrequency catheter ablation (PRCA)**
Left ventricular mapping looking for the earliest ectopy in relation to QRS. PRCA was conducted with thermo-controlled applications (60°C, 30 s) in the middle septal region of the anterior wall. After waiting for 30 minutes, there was no recurrence of arrhythmia, even after PVS with up to three extra-stimuli and on isoproterenol infusion.

![Figure 2](image1.png)
**Figure 2.** Initial excursion of negative QRS complexes (Vi)/terminal excursion of QRS complexes (Vt) < 1 (Vi/Vt < 1) [5].

![Figure 3](image2.png)
**Figure 3.** Capture and fusion beats are observed (atrioventricular dissociation). Conclusion: idiopathic infr fascicular re-entrant left ventricular tachycardia.

![Figure 4](image3.png)
**Figure 4.** A resting electrocardiography recorded immediately after ventricular tachycardia event showing typical features of athletic heart: increased QRS amplitude in V4–V5 associated with asymmetric T wave inversion. The T wave amplitude is low in inferior limb leads.
Discussion

Idiopathic intrafascicular reentrant left ventricular tachycardia (ILVT) or verapamil-sensitive ventricular tachycardia [1] is the second most common form of idiopathic VT. ILVT is usually seen in young males without apparent structural heart disease, featuring paroxysmal palpitations and RBBB morphology. QRS axis shifting depends on the fascicle involved. Characteristically, extreme left axis deviation is seen in left anterior fascicular tachycardia, whereas left axis (–45 to –60) deviation is noted with left posterior fascicular tachycardia. Zipes et al. [2] described a triad characteristic for ILVT: induction with atrial pacing, RBBB pattern with extreme left axis deviation, and no structural heart disease.

Idiopathic ventricular tachycardia (VT) encloses a heterogeneous group of tachycardias that may result from multiple cellular electrophysiological mechanisms [3]. It is characterized by two predominant forms. The most common form originates from the right ventricular outflow tract and presents as repetitive monomorphic VT or exercise-induced VT. The tachycardia is adenosine-sensitive and is thought to be caused by cAMP-mediated triggered activity. The other major form seen in this case is verapamil-sensitive intrafascicular re-entrant tachycardia, which most often originates in the region of the left posterior fascicle. Both forms of idiopathic VT can be readily treated with PRCA.

Conclusions

Recognizing ILVT and its unique characteristics can lead to a correct diagnosis and facilitate appropriate therapy.

Acknowledgements

The authors do not report any conflict of interest regarding this work.

References