Late complication of heart stimulation — lead abrasion in pacemaker pocket

Krzysztof Boczar¹, Barbara Małecka¹-², Andrzej Ząbek¹, Kazimierz Haberka¹, Jacek Lelakowski¹-²

¹Department of Electrocardiology, John Paul II Hospital in Krakow, Poland
²Institute of Cardiology, Jagiellonian University Medical College in Krakow, Poland

An 81-year-old male patient was hospitalised due to recurrence of syncope 4 years after DDD pacemaker (PM) implantation. The DDD pacing system contained Vitatron C60DR PM and 2 Vitatron Crystalline BP leads: an atrial one implanted by left subclavian venipuncture, and a ventricular one implanted by left cephalic venesection. In this PM-dependent patient, periodic inhibition of pacing up to 2 s was recorded in Holter electrocardiogram. Pacing inhibition occurred although bipolar sensing was programmed. In the PM check-up, moving of the patient’s left upper limb induced short-cycle abnormal signals, so called ‘crackles’, simultaneously in atrial and ventricular channels (Fig. 1). The remaining parameters i.e. pacing thresholds and lead impedance, were correct. Lead damage was diagnosed and a decision to perform transvenous lead extraction (TLE) was made. Because this patient had left ventricular ejection fraction lowered to 30% in the course of ischaemic cardiomyopathy, we decided to perform simultaneous implantation of a new implantable cardioverter-defibrillator (ICD) to prevent sudden cardiac death. Before the TLE procedure, venography of left-side venous inflow was performed and occlusion of left brachiocephalic vein with advanced collateral circulation was observed (Fig. 2). Nevertheless, a 150 cm-long guidewire Cordis 0.035 was successfully introduced into the heart. Firstly an ICD lead was implanted with the help of a set containing a long Peel-Away Curved Sheath made by Cook, designed for such a condition of blood vessels. The presence of the ICD lead in the right ventricle was used for temporary pacing during the removal of the damaged leads. Both leads were extracted using simple traction. Then we went along the narrowing of the inflow vein once again using a Medtronic Attain Command set designed for left ventricular lead implantation, and we implanted a new atrial lead. On examination of the extracted leads, we observed the presence of some abrasion in the external silicone lead insulation, with lead unsealing and metal wire exposure (Fig. 3). The damage distance from the distal end of the leads was compared to a chest X-ray conducted before TLE (Fig. 4). Measurements were performed using the DICOM system. This allowed the identification of sites of damage of the leads in the PM pocket where lead loops were located near the PM can, and where they remained in mutual contact (Fig. 4). Abrasion of endocardial silicone leads due to mutual friction is an important clinical problem. In this process, silicone insulation gradually becomes thinner until it completely breaks. This can have a negative influence on the functioning of the pacing system. It seems justified to take into account the abrasion of lead insulation in cases of pacing inhibition in the presence of bipolar sensing leads.

Address for correspondence:
Andrzej Ząbek, MD, MSc, Department of Electrocardiology, John Paul II Hospital in Krakow, ul. Prądnicka 80, 31–202 Kraków, Poland, e-mail: andrzej_j_z@poczta.onet.pl

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Figure 1. ‘Crackles’ in ventricular IEGM = VEGM (circles) and in atrial IEGM = AEGM (rectangles), inhibiting pacing for 1750 ms with lack of rhythm — lead II (arrow). Artifacts in leads I and III — wandering of isoelectric line while moving left upper limb

Figure 2. Collateral circulation with the inflow by right brachiocephalic vein (arrows)

Figure 3. Silicone insulation breakage in macrophotography

Figure 4. Chest X-ray — postero-anterior projection; A — atrial lead; V — ventricular lead; C — the site of mutual contact of leads in the pocket. In the circle can be seen magnification of the site C