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Lead to lead interaction leading to left bundle branch area pacing lead failure

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Conflict of Interest:
Pugazhendhi Vijayaraman reports being a speaker and consultant for, and receiving research and fellowship support, from Medtronic; being a consultant for Abbott, Biotronik, and Boston Scientific; and patent for an HBP delivery tool

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AV node ablation and conduction system pacing utilizing left bundle branch area pacing (LBBAP) is an excellent option for patients with atrial fibrillation and rapid ventricular rates refractory to medical therapy.\cite{1,2} Patients with severely reduced left ventricular systolic function benefit from implantable cardioverter-defibrillator for prevention of sudden cardiac death.\cite{3} Biventricular pacing using coronary vein lead has been shown to reduce heart failure hospitalization and mortality compared to right ventricular pacing in patients undergoing AV node ablation\cite{4}, but may be limited by anatomical challenges, non-physiologic biventricular activation, phrenic nerve stimulation and/or high pacing thresholds. While His bundle pacing is an excellent option in these patients, it can be technically challenging and be associated with unexpected late threshold rise.\cite{5} In addition to better sensing, lower and more stable long-term thresholds are an advantage with LBBAP compared to His bundle pacing.\cite{6-8} While LBBAP has been shown to be safe in multiple observational studies, concern regarding the long-term integrity of the LBBAP lead remains due to a significant portion of the lead being buried deep in the interventricular septum. We report a case of LBBAP lead failure due to interaction with the defibrillator lead near the septal insertion site.

Case Report

A 68-year-old man with hypertension, atrial fibrillation refractory to medical therapy and multiple ablations, non-ischemic cardiomyopathy with left ventricular ejection fraction of
25-30% was referred for AV node ablation. He underwent AV node ablation and uncomplicated biventricular implantable cardioverter-defibrillator (ICD) placement utilizing 3830 SelectSecure R lead for LBBAP (LV port), Sprint Quattro R ICD lead in right ventricular apical septum and 5076 CapSureFix R Novus lead (Medtronic Inc, Minneapolis, MN) in the right atrial appendage. First, the His region was identified using double curved His Delivery sheath (C315 Sheath) guided by local electrocardiogram, His capture and fluoroscopy. The system, lead and sheath, was then advanced approximately about 2 cm distally towards right ventricular apex and rotated counterclockwise to maintain perpendicular orientation to septum. At this point, the lead was advanced into the septum with rapid clockwise rotations while monitoring for triggered ventricular beats, changes in impedance, current of injury and local electrocardiogram amplitude and injury. The rotations were repeated until LBBAP was confirmed. LBBAP lead demonstrated anodal capture threshold at 3V and nonselective left bundle capture at 0.5V at 0.4msec. Device was programmed to pace from LBBAP lead and RV lead with LV-RV delay of 80 ms. Patient lacked adequate internet coverage and refused remote monitoring. He had regular follow-up in the device clinic and cardiology clinic. On follow-up, he noted improvement in exercise capacity with improvement in NYHA functional class from III to II. A remarkable improvement was noted in left ventricular ejection fraction from 25-30% at baseline to 49% at 12-month follow-up.

Twenty months after implant, he was seen in heart failure clinic for dyspnea on exertion and fatigue. Acute rise in impedance and threshold with loss of capture of LBBAP lead
was recorded on device check (Figure 1A), suggestive of lead fracture. Both the right ventricular defibrillator and atrial leads showed stable threshold and impedance (pacing and shocking) parameters. The patient was scheduled for LBBAP lead extraction and implantation of new LBBAP lead. Pre-procedural cine-fluoroscopy of the system (Supplemental Video 1) revealed lead-lead interaction with the defibrillator coil sliding against the LBBAP lead near the ring electrode. Fluoroscopic images (Figure 1B) showed a sharp bend in the 3830 SelectSecureR lead where the defibrillator coil appears to make contact. The LBBAP lead was easily extracted with manual traction and a new 3830 SelectSecureR lead was placed deep in the mid-septum away from the defibrillator coil with non-selective left bundle branch capture threshold of 0.6V @ 0.4 msec. The patient tolerated the procedure well without any complications.

Gross inspection of the extracted LBBAP lead showed a break in insulation and conductor where the lead was in contact with the defibrillator coil (Figure 2A). Detailed analysis of the lead showed wear and tear through the outer insulation, outer coil, ethylene-tetrafluoroethylene (ETFE) on the tip conductor, and the tip conductor (Figure 2B). Source of lead abrasion was determined to be the defibrillator coil. Longitudinal and 3-dimensional schematic (Figure 2C) of the SelectSecureR lead provides a reference to compare the damage on the various components of the lead. He reported significant improvement in functional status (NYHA II from NYHA III) on post implant follow up at 3 months. Repeat echocardiogram showed normalization of ejection fraction.
Discussion

The 3830 SelectSecure® lead consists of a Platinum/Iridium ring electrode connected to an outer coil conductor, and ETFE jacketed cable tip conductor, silicone inner insulation tubing and polyurethane outer insulation tubing. It is a co-axial, bipolar, steroid eluting, lumen-less, fixed-screw pacing lead approved for selective site pacing and His bundle pacing. No cases of intracardiac lead failure due to lead-lead interaction of this lead have been reported so far. To our knowledge, this is the first report of 3830 SelectSecure® lead failure due to intracardiac lead to lead interaction when used for conduction system pacing. The LBBAP lead in the initial implantation was located more posteriorly (Figure 3A) with the lead body proximal to the ring electrode coming in close contact with the defibrillator lead coil (Figure 1B). The septum likely acted as a fulcrum for the LBBAP lead and with cardiac motion, the lead body continuously rubbed against the defibrillator lead coil resulting in progressive abrasion and ultimately lead failure. We hypothesize the effective area of the insulation breach was not large enough to cause measurable drop in impedance before the rise in impedance due to lead failure from interaction.

The challenges of extracting the LBBAP lead in the deep septal location are currently unknown. It is unclear if extensive fibrosis will occur in this location along the length of the tip and the ring electrode intraseptally and around the insertion site, during long-term follow-up. In our case, the LBBAP lead was only 20 months old and was easily extracted with gentle traction. There was no significant fibrosis at the lead tip or
insertion site. Reimplantation of a new LBBAP lead was done at a more anterior part of the septum (Figure 3B) to maintain adequate distance between the hinge point of the LBBAP lead and the ICD lead coil. Our case report highlights the possible risk of lead-lead interaction of the LBBAP lead with the defibrillator lead. Care should be taken to maintain adequate distance between the hinge-point of the LBBAP lead and the defibrillator coil.


Figure 1. Impedance and threshold trend. A: Impedance and threshold rise in 3830 SelectSecureR lead, followed by open circuit and loss of capture. B: Fluroscopic image demonstrating the defibrillator lead coil in contact with the proximal end of the ring electrode of 3830 SelectSecureR lead. A sharp bend in the 3830 SelectSecureR lead is seen where the defibrillator lead coil appears to make contact.

Figure 2. Macroscopic image of the lead fracture. A: Apparent visible wear and tear on the 3830 SelectSecureR lead after explanation. B: Wear through the outer insulation, outer coil, ETFE on the tip conductor, and the tip conductor. C: Longitudinal and 3-dimensional schematic of the 3830 SelectSecureR lead (Images used with permission from Medtronic, plc © 2022).

Figure 3. Chest X-rays and electrocardiogram of initial and final lead position. A: Chest X-ray showing the posterior location of the LBBAP lead after initial implantation with proximity of the ring electrode to the defibrillator lead coil. B: Chest X-ray showing a more anterior location of the LBBAP lead after re-implantation with adequate distance from the defibrillator lead. C: Pacing morphology of initial implant with anodal capture and nonselective Left septal capture D: Pacing morphology of new implant with Anodal capture and nonselective Left bundle selective capture.
Figure 1

A

LV Tip-Ring impedance

Fracture?

Electrode interface change

LV Tip-Ring pacing threshold

Last 80 weeks (max/min/avg per week) | Last 15 days |

B

Quatro defibrillating coil appears to contact the 3830 proximal of ring electrode

Sharp bend in path of 3830
Figure 2
Figure 3

A

B

C

D
KEY TEACHING POINTS

- Conduction system pacing (CSP) with left bundle branch area pacing (LBBAP) is emerging as an alternative strategy to traditional coronary sinus pacing for cardiac resynchronization therapy (CRT). Low and stable long-term thresholds make LBBAP an excellent option for physiologic pacing.

- Long-term safety profile, lead integrity, lead to lead interaction and risk of extraction of LBBAP leads need to be determined. LBBAP lead is well-anchored, deep into the interventricular septum, which increases the risk of lead-lead interaction from constant friction when placed near a defibrillator lead.

- We report the first case of a LBBAP lead failure due to interaction with defibrillator lead. Adequate distance between the defibrillator lead and the LBBAP lead insertion site needs to be maintained at the time of implantation to avoid lead-lead interaction and potential lead failure.