CASE REPORT

Effective Pacing Intervention by Closed-loop Stimulation Using a Coronary Vein Lead in a Post-tricuspid Valve Replacement Patient

Yukitoshi Ikeya, Toshiko Nakai, Nobuhiro Murata, Masaki Monden, Akihito Ogaku, Koichiro Hori, Ryuta Watanabe, Masaru Arai and Yasuo Okumura

Abstract:
To avoid the negative effects associated with pacing, pacemakers are designed to achieve a pacing cadence as close to physiological pacing as possible. In closed-loop stimulation (CLS; a type of rate-responsive functionality used in pacemakers), the changes in impedance (which correlates with the contractility of the myocardium around the lead tip electrode) are tracked, and the paced heart rate is adjusted accordingly. We herein report a case in which we implanted a pacemaker in a post-tricuspid valve replacement patient. A ventricular lead positioned in the coronary vein exhibited good CLS functionality, and the patient’s dizziness and heart failure improved.

Key words: closed-loop stimulation, CLS, coronary vein lead, left ventricular pacing

Introduction
The design of pacemakers aims for pacing that is as physiological as possible in order to prevent the detrimental effects of pacing on the cardiac function and to improve the patient’s exercise tolerance. Biotronik pacemakers use a unique closed-loop stimulation (CLS) technology as a form of rate-responsive functionality.

CLS is a technology that extracts changes in impedance, which correlates with the contractility of the myocardium around the lead tip electrode, and converts the changes into a heart rate response in order to maintain the cardiac output, enabling more natural, physiological rate responsivity.

We herein report a case in which we placed the pacing lead and left ventricular (LV) lead [for cardiac resynchronization therapy (CRT)] of a pacemaker with CLS functionality in the patient’s coronary vein when implanting a single-chamber ventricular (VVI) pacemaker in a post-tricuspid valve replacement (TVR) patient with chronic atrial fibrillation (AF) with bradycardia. The lead, which was positioned on the epicardial side of the LV, exhibited good CLS functionality.

Case Report
The patient was a 79-year-old Japanese woman who visited the emergency room at our hospital for presyncope and nausea in June 2019. As the monitor showed bradycardia associated with chronic AF and because the patient’s subjective symptoms were persisting, she was admitted to the hospital.

The patient had undergone pulmonary vein isolation (PVI) for paroxysmal AF in 2010, which progressed to chronic AF. In addition, TVR was performed in March 2016 for tricuspid valve insufficiency. In other systems, endoscopic polypectomy for early gastric cancer and stent placement for lower extremity arteriosclerosis obliterans (ASO) were both performed in 2016. The patient’s medical history included chronic heart failure, chronic obstructive pulmonary disease, hypertension, chronic renal failure, pulmonary hypertension, and diabetes.

The physical findings at the initial visit included the following: body temperature 36.7 °C, blood pressure 182/80...
mmHg, pulse rate 45 beats per minute (bpm), irregular; and no heart murmurs or abnormal respiratory sounds. Her blood test results showed chronic anemia (Hb 9.4 g/dL) and renal dysfunction [blood urea nitrogen (BUN) 27.7 mg/dL, Cr 15 mg/dL]. Her thyroid function was within the normal range [thyroid-stimulating hormone (TSH), 2.32 μIU/mL; free triiodothyronine (FT3) 3, 2.51 pg/mL; free thyroxine (FT4) 4, 1.38 ng/dL]. There were no abnormalities in cardiac enzyme levels [creatinine kinase (CK), CK-muscle brain (MB), troponin I] and no changes in markers of inflammation, and the results of liver function tests were normal. NT-proBNP was high at 8,464 pg/mL.

Electrocardiography (ECG) revealed AF [heart rate (HR): 20-50 bpm] and complete right bundle branch block (QRS=144 ms). Inverted T waves were seen in leads II, III, aVF, and V1-4 (Fig. 1). Chest X-ray showed cardiomegaly (cardiothoracic ratio: 61%) and pulmonary congestion.

Echocardiography did not exhibit clear asynergy in LV wall motion: the LV ejection fraction (LVEF) was 69%, the LV end-diastolic dimension (LVDd) was 54 mm, the LV end-systolic dimension (LVDs) was 33 mm, the interventricular septal thickness at end-diastole (IVSTd) was 10 mm, and the posterior LV wall thickness (PWTd) was 10 mm. The findings indicating valvular disease included trivial tricuspid regurgitation (TR) (post-TVR), mild to moderate pulmonary regurgitation (PR) [PR end-pressure gradient (PG): 3 mmHg], right atrial and ventricular (RA and RV) enlargement, and left atrial (LA) enlargement with a left atrial diameter (LAD) of 47 mm.

After admission to the hospital, a β-blocker (bisoprolol fumarate, 0.625 mg) that the patient had been taking was withdrawn, and she was followed-up with diuretic dose adjustments. However, as the patient frequently had a heartbeat pause lasting 5-8 seconds associated with lightheadedness and since her bradycardia did not improve, permanent pacemaker implantation was performed. Because the patient’s AF had become chronic, a VVI pacemaker was used. An LV lead for CRT - instead of an RV lead - was inserted in the coronary sinus (CS) and placed on the LV side (CS lateral branch), since the patient had previously undergone TVR, although a biological tissue valve had been used.

At the implantation, the R-wave sensing was 12.1 mV, the ventricular pacing threshold was 0.6 V at 0.4 ms, and the lead impedance was 526 Ω. All measurements were within the normal range. The pacemaker implanted this time was a model with a CLS function (Edora 8 SR-T; Biotronik, Berlin, Germany), and the settings of VVI-CLS were a lower rate of 60 ppm and sensor rate of 120 ppm. After the pacemaker implantation, the doses of the patient’s oral medications, including the β-blocker and diuretics, were adjusted once more. Her lightheadedness and heart failure signs both improved, and the patient was discharged 10 days after the implantation.

An inspection of the pacemaker and ECG Holter monitoring after the patient’s pacemaker implantation indicated that even though pacing was being performed by a lead placed on the epicardial side of the coronary vein, the CLS functionality was good, as indicated by an increased pacing rate (Fig. 2) and the following data: total heart beats (/day), 100,108; maximum HR (bpm), 121; minimum HR (bpm), 59; average HR (bpm), 70; and longest RR interval (sec), 1.22 (Table). Post-pacemaker implantation chest X-ray performed after the patient’s discharge did not show pulmonary congestion (Fig. 3).

**Discussion**

We implanted a VVI pacemaker in a post-TVR patient
with AF and bradycardia. As the patient had previously undergone TVR, although a bioprosthetic valve was used, a V-lead was placed in her left ventricle using an LV lead for CRT. A pacemaker with a CLS function was selected, which worked with an LV lead.

To achieve a rate-responsive function, sensors other than the CLS were used in the present case, including an accelerometer that responds to body movements and has a superior rate response during exercise at a high activity level, and a minute ventilation sensor that monitors respiration and is suitable for responding to slow exercise and changes in mental activity.

The present patient had AF with bradycardia as her underlying heart condition; she had undergone TVR surgery and had a history of hospitalization for heart failure. The use of β-blockers for her heart failure treatment worsened the bradycardia, and the patient developed heart failure again. The pacemaker implantation stabilized the patient’s heart rate, and the dose adjustments of oral drugs, including the β-blocker, alleviated her heart failure.

As noted above, in patients who have had bradycardia or develop bradycardia or chronotropic incompetence induced by a β-blocker, backup pacing at a physiological rate is important for preventing heart failure. CLS uses a sensing algorithm that can detect variations in cardiac contractility and respond to a decrease in the blood pressure by increasing the heart rate (1). Several research groups reported that CLS pacing is beneficial for patients with recurrent vasovagal syncope who demonstrate a cardio-inhibitory response on the head-up tilt table test (HUTT) (1-4). Another report showed that CLS is effective during states of emotional stress in patients with AF and chronotropic incompetence (5).

For these reasons, CLS was considered appropriate in this patient, who was 79 years old with an activity level that was not high and who had an unstable heartrate associated with AF and exhibited signs of heart failure. CLS is normally used with endocardial right ventricular leads, and whether or not the sensor functions well in epicardial pacing with a lead positioned in the coronary vein, as in our patient’s case, was unclear. However, we selected Biotronik’s Edora 8 SR-T model because previous reports noted that CLS was capable of heart rate modulation in pediatric patients when connected to an epicardial lead (6, 7); in addition, we considered that, should CLS prove ineffective, the use of the rate response function of the accelerometer included in this model would be a useful contingency plan.

The CLS function was useful for LV pacing with an LV lead placed in the coronary vein of our adult patient. Our experience in this patient’s case suggests the possibility of
capitalizing on the advantages of CLS in postoperative dialysis patients and patients who require a surgically placed epicardial lead.

Author’s disclosure of potential Conflicts of Interest (COI).
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