Leadless pacemakers:

A new technology in cardiac pacing

Learn about the nursing implications of these innovative devices.

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ONE of the most recent developments in cardiac pacing is the leadless pacemaker (LP). This device is one of the biggest advancements in pacemakers since they came into use for symptomatic bradycardia more than 50 years ago. (See Traditional pacemaker basics.) However, as with any new technology, a knowledge gap exists. Many nurses may not have encountered a patient with an LP and little nursing literature on the topic exists. Understanding this new technology will help you deliver safe care to patients who may be candidates for an LP or who may already have one.

LPs vs. traditional pacemakers

The LP was developed to avoid the complications associated with the leads and surgical pocket required when placing a traditional pacemaker. LPs, which are miniature devices, are implanted directly into the right ventricle via the femoral vein in the groin.

LPs are single-chamber devices, so currently they’re available only for a limited number of pacemaker candidates. The 2009 world survey by Mond and Proclemer provides prevalence data on the use of pacemakers by type. In the United States, just 20% of patients who required permanent pacemakers received traditional single-chamber devices, while the remaining 80% received traditional dual-chamber devices. (See Pacemaker indications.)

LP availability

Medtronic won Food and Drug Administration (FDA) approval to market the Micra LP on April 6, 2016. (See Pacemaker comparison.) The device is 25.9 mm long, 6.7 mm wide, and weighs 2 grams. It’s inserted with a catheter through the femoral vein with a 23 French introducer, advanced into the right ventricle, then attached to the en-
Traditional pacemaker basics

Traditional permanent pacemakers can pace a single chamber (atrial or ventricular), dual chambers (both atrial and ventricular), or biventricular chambers (both ventricles, specifically for the treatment of heart failure and cardiomyopathy).

Components
Pacemaker components include a battery, a pulse generator, and transvenous leads. The transvenous lead goes from the pulse generator through a central vessel to the chamber of the heart. One or two leads may be required depending on whether it's single- or dual-chamber pacing.

Placement
Before 2016, if a patient needed a pacemaker, the battery and pulse generator component usually were placed under the skin below the clavicle into a surgical pocket. The transvenous lead was then advanced through a central vein, usually the subclavian or cephalic vein, and attached into the heart chamber to deliver an electrical impulse to generate a paced heartbeat.

Complications
In a study by Udo, about 12% of patients who received permanent pacemakers experienced complications within 2 months, and around 9% experienced complications after 2 months. Complications during device implantation included pneumothorax and cardiac tamponade. After implantation, complications associated with the surgical pocket included hematoma, skin breakdown, and pocket infections. Complications associated with the transvenous leads can occur at any time.

Most of the complications that occurred 2 months after placement resulted from problems with the transvenous leads. They can cause venous obstruction from thrombosis and stenosis because of inflammatory and fibrotic changes that result from the lead position within the vein. In addition, leads can dislodge or break because of compression between the clavicle and first rib or if they're caught in soft tissue.

docardium. After it’s fixed to the inside of the right ventricle, the catheter and introducer are removed. Vascular closure of the femoral vein, using manual pressure, purse-string suture, or a vascular closure device, prevents bleeding at the groin site.

As of October 2017, Medtronic reports that more than 10,000 patients in 40 countries have a Micra LP. Currently, Micra LP is the only LP on the market. St. Jude Medical (now Abbott) conducted clinical trials of its Nanostim LP, but after 1,423 implants, the company halted implantation in October 2016 because of seven reports of battery issues that resulted in lost telemetry and pacing output. The manufacturer recommends device replacement for anyone with the Nanostim. Of those, six experienced bradycardia. No other adverse outcomes have been reported. Crotti reports that Abbott is now planning to focus on developing the dual-chamber leadless pacemaker, as this is where the greater need is. Micra’s battery life is expected to be 10 years or more.

LP candidates
Patients who require a VVIR (ventricle pacing, ventricle sensing, paced beat inhibition, rate modulation) for symptomatic bradycardia may be candidates for the LP. (Note: For pacing code information, visit americanrnursetoday.com/?p=52458.) Patients with existing pacemakers who’ve had surgical pocket or transvenous lead complications also may benefit from this technology.

Some patients aren’t good candidates for an LP. They may require the more complex features and programmable technologies available in dual-chamber and biventricular pacemakers. Others may have a mechanical tricuspid valve, which prevents passing the LP into the right ventricle. Similarly, if a patient has a vena cava filter, the introducer can’t be inserted through the groin. (See LP contraindications.)

Nursing care
Most LP placement procedures take about 1 hour, and patients usually stay in the hospital overnight.

After device placement, assess the patient’s blood pressure, heart rate, cardiac rhythm, respiratory rate, and oxygen saturation; evaluate the groin site for bleeding, oozing, or hematoma; and assess circulation by palpation or Doppler signals distal to the procedure site. Monitor the patient every 15 minutes for 1 hour, every 30 minutes for 1 hour, every hour for 2 hours, and then follow the usual protocol unless assessments indicate more frequency is needed. Bedrest may be maintained for about 4 hours to allow for hemostasis at the groin site and to prevent bleeding complications. The appropriate provider will interrogate the pacemaker by placing a wand-like device over the pace-
Pacemaker indications

The 2008 Guidelines for Device-Based Therapy developed by the American College of Cardiologists, the American Heart Association, and the Heart Rhythm Society remain current today.

Permanent cardiac pacemakers are inserted to treat symptomatic bradycardia, which is a documented bradycardia syndrome responsible for symptoms such as syncope or near syncope, dizziness, confusion, fatigue, exercise intolerance, and heart failure.

Symptomatic bradycardia may be caused by sinus node dysfunction (SND) or atrioventricular (AV) node dysfunction (other conditions may cause symptomatic bradycardia but are beyond the scope of this article).

- **SND** (sometimes called sick sinus node) is common in patients in their 70s and 80s and is considered a result of sinus node aging. It can be further defined as:
  - persistent bradycardia (a slow heart rate for an extended period)
  - chronotropic incompetence (inadequate heart rate response to activity or exercise)
  - tachy-brady syndrome (alternating very fast and very slow heart rates, often atrial fibrillation).

- **AV node dysfunction** is represented by symptomatic bradycardia associated with advanced second- or third-degree heart block. It can be further defined as:
  - bradycardia associated with advanced second- or third-degree heart block with or without atrial fibrillation
  - escape rates less than 40 beats per minute
  - escape rhythms below the AV node
  - atrial fibrillation associated with bradycardia and pauses.

maker area to allow the pacemaker to communicate with a computer that evaluates device function, the battery, and any abnormal cardiac rhythms the patient may have experienced.

Notify the provider of any assessments that are out of normal range or unexpected, such as vital sign changes, chest pain or palpitations, abnormalities on the cardiac monitor, diminished pulses, and color change or temperature change of the extremity with the groin site. A failure to pace will appear on the cardiac monitor as an absent pacemaker spike when one is expected and as a heart rate lower than the rate set on the pacemaker. Note that if the patient’s native heart rate is higher than the demand rate on the pacemaker, then no pacing will occur and no pacemaker spikes will be seen on the monitor.

If you observe a failure to pace when pacing should occur, assess the patient clinically for symptoms of bradycardia and notify the provider or rapid response team. Interventions will depend on the patient’s clinical status and may include emergency measures for symptomatic bradycardia, such as atropine, dopamine, or epinephrine; a temporary pacemaker; continued observation; and bedrest. Other pacemaker rhythm disturbances include a failure to capture (the pacemaker fires, but doesn’t cause contraction), oversensing (pacemaker fires less frequently than it should), and undersensing (pacemaker fires more frequently than it should).

Alert the provider if you find evidence of hematoma, bleeding, or pain at the affected groin site. While awaiting evaluation from the provider, demarcate the area on the dressing, reinforce the dressing as necessary, and apply pressure if needed.

### Potential complications

At the start of the investigational study for both the Micra LP and Nanostim LP, the FDA identified potential complications, including right ventricle perforation, cardiac tamponade, pneumothorax, hemothorax, device dislodgement, device migration during implantation, infections requiring reoperation or extraction, and femoral vascular access complications (arteriovenous fistula, pseudoaneurysm, and excessive bleeding).

At the completion of the investigational trial period, Reynolds and colleagues reported that the Micra was successfully implanted in 719 of 725 patients. Twenty-eight major complications occurred in 25 of the 725 patients (a rate of 4%). These complications included one with pulmonary embolism, one with deep vein thrombosis, five with events at the groin puncture site, 11 with cardiac injury (three with pericardial effusion), and eight with other cardiac issues (one with myocardial infarction, three with cardiac failure, one with pacemaker syndrome, and one each with metabolic acidosis, pre-syncope, and syncope).

Since the Micra has been approved by the FDA, Roberts and colleagues have been reporting on the progress of the LP in the real world and comparing their results with those of the investigational research. In this ongoing study, LPs have been implanted in 792 of 795 patients. Thirteen major complications occurred in 12 patients (a rate of 1.51%). These complications included one deep vein throm-
LP contraindications

Patients with the following conditions are not candidates for the leadless pacemaker (LP).

- Need for more complex pacemaker features and programmable technologies
- Pacemaker syndrome (a collection of signs and symptoms of heart failure, including dyspnea, hypotension, lightheadedness, and fatigue, caused by right ventricular pacing)
- Other implanted medical devices (including cardiac devices) that would interfere with the LP
- Mechanical tricuspid valve prosthesis
- Implanted vena cava filter
- Intolerance to any materials involved
- Sensitivity or allergy to heparin, to contrast media that can’t be adequately premedicated for, or to dexamethasone acetate

Discharge teaching

Before the patient is discharged from the hospital, the provider will evaluate the pacemaker for proper pacing, a chest x-ray will be ordered to verify device position, and a 12-lead electrocardiogram (ECG) will be performed to verify cardiac rhythm. Inform the patient that if he or she is feeling well, a return to usual activities can begin in 1 or 2 days. To allow for groin healing, the patient should avoid heavy lifting and squatting for 1 week.

Discuss the importance of attending follow-up appointments for a physical exam, ECG, and pacemaker adjustments as needed to pace the heart faster or slower or to deliver more or less energy to optimize therapy. Explain that the pacemaker should relieve the symptoms of bradycardia and that the patient should be able to complete activities of daily living more easily.

Instruct the patient to report symptoms of syncope, dizziness, confusion, fatigue, and exercise intolerance. These symptoms may indicate pacemaker malfunction or the need for setting adjustments.

Tell the patient to always carry his or her identification (ID) card, which includes important information about the type of pacemaker, serial number, and date and location of implantation.

Verify that the patient has the device patient manual, which includes information about the effects of electromagnetic interference (EMI). Although cell phones are described as having low or no risk, instruct the patient not to carry his or her cell phone in a shirt pocket. Explain to the patient that if he or she feels dizzy or experiences a rapid or irregular heartbeat while using an electrical device, to let go of the device and move away from it. The pacemaker should return to normal function away from the EMI. In addition, explain that the patient shouldn’t enter an area with a “no pacer” symbol.

Patients with an LP should talk with their cardiologist before undergoing medical procedures. Electrosurgery, electrocautery, external defibrillation, lithotripsy, or radiation therapy can cause interference with pacemaker functioning.

Medtronic describes its LP as being magnetic resonance imaging (MRI) conditional. “Conditional” refers to the MRI magnet strength, proper pacemaker functioning, duration of implant, and country in which the pacemaker was inserted. These conditions are subject to change and will evolve over time with this new device. The Medtronic MRI technician manual defines the conditions.

Although airport security metal detectors aren’t likely to affect an LP, Medtronic recommends that patients pass through the detectors at a normal pace but not linger in the archway. Advise the patient to immediately move out of range of the detector if he or she experiences fatigue or dizziness. If the metal detector is set off by the LP, the patient should show the ID card to security personnel and explain that the handheld detection wand should not be
Joy in the journey

All of us experience tragedy, sadness, and grief; they’re part of the human condition. If you’re wondering if finding joy and peace is possible under what appear to be impossible conditions, remember this: History is replete with ordinary humans rising to challenges of the day in extraordinary ways. They were able to unlock that part of themselves that gave them the strength and courage to carry on.

Nurses are extraordinary—don’t lose sight of the amazing work you do to improve the lives and comfort of the people you touch. It’s never too late to make a positive change in your life. (See Continue the journey.) If you take a few small steps in the direction you want to go, you’ll be amazed at the results.

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Selected references


