Pacemaker Placement for You or Your Child

What is a pacemaker?
A pacemaker is a small device that corrects your child’s heart rhythm. Your child’s heart has a natural pacemaker called the **sinus node** — a cluster of cells that sends electrical impulses to make the heart beat. The electrical impulses move down through the heart, and are sent to the heart’s pumping chamber through the **atrioventricular node**. If the sinus node or atrioventricular node doesn’t work properly, an artificial pacemaker can keep your child’s heart beating at the right pace.

**Types of pacemakers**
There are two types of pacemakers:
- An **on-demand pacemaker** is set for a certain heart rate, and turns on only when your child’s heartbeat slows below this rate.
- A **rate-responsive pacemaker** can monitor breathing, blood temperature, and other factors. It uses these to change the rhythm based on how active your child is.

**Parts of a pacemaker**
A pacemaker has two parts:
- A **pulse generator** — the battery and circuits that create low-energy electrical pulses. The generator is a little larger than a quarter and less than ¼-inch thick.
- **Leads** — thin insulated wires that deliver the electrical pulses to your child’s heart.
  - A single-chamber pacemaker uses one lead to send pulses to the heart’s upper or lower chamber.
  - A dual-chamber pacemaker uses two leads: one in the upper chamber and one in the lower chamber.
  - A biventricular pacemaker has three leads. They are placed in the right upper chamber, right lower chamber, and left lower chamber.

**Why does my child need it?**
A pacemaker is used to correct:
- Damage to the sinus node that makes it fire slowly
- A problem with electrical signals moving through the heart
- A slowed or irregular heartbeat caused by heart failure or by heart medications your child must take (such as beta blockers)

These problems can cause symptoms such as dizziness, lightheadedness, shortness of breath, fatigue, or fainting.

Note: if you are an adult having this procedure at Primary Children’s Hospital, please substitute “you” for “your child” in the text.
How should we prepare?
There are several things you can do to get ready for the procedure and make it go more smoothly.

- **Arrange for time off work or school.** Your child can return to work or school when the doctor says it’s okay, usually after a week or so.

- **Tell your child’s doctor about medications and allergies.** Along with prescription medications, include over-the-counter drugs, herbs, and vitamins.

- **Follow the doctor’s directions about medications.** Your child may need to stop taking certain blood thinners before the procedure.

- **Arrange for a ride.** You or someone else needs to give your child a ride to and from the hospital.

- **Have your child fast (no food or drink) for 6 to 8 hours before the procedure.** If the procedure is in the morning, your child should not eat or drink anything after midnight the night before.

**What happens before?**
When your child arrives, here’s what will happen:

- You will fill out some paperwork and your child will change into a hospital gown.

- An IV (intravenous) line may be placed in your child’s arm or hand. Your child may be given medicine by mouth or through a mask before the IV is placed.

- Blood may be drawn for lab tests.

- The left or right side of your child’s chest might be shaved, if necessary, and cleaned.

- Your child will be moved to the cardiac cath lab. The room may feel cool, but your child will be covered with sterile drapes and a blanket.

- Your child will lie on her back. Your child will be given medication through the IV to make her sleep through the procedure.

- Your child’s arms may be secured at her sides, because it is important for the arms to be still during the procedure.

**What happens during?**
This procedure usually takes several hours. Here’s what will happen:

1. **Monitoring.** Devices will be attached to keep track of your child’s heart rate, blood pressure, and breathing.

2. **Incision.** The doctor will make a small incision (cut) in the skin. This makes a “pocket” for the pulse generator.

3. **Placing one or more leads.** The doctor will insert a needle into a vein, and will insert each lead using the needle. Using x-ray guidance, the leads will be threaded through the vein into your child’s heart.

4. **Testing the leads.** The medical team will take electrical measurements to make sure each lead is in the correct place. Each lead might be moved, and retested each time, until it is perfectly positioned.

5. **Attaching the pulse generator.** The pulse generator will be connected to the leads and will be inserted under your child’s skin.

6. **Programming the pacemaker.** The pacemaker will be set to the rate your child’s heart needs. The medical team might also adjust other settings.

7. **Closing the incision.** Your child’s doctor will close the surgical cut with a few stitches, and your child will be moved to recovery.

**What happens after?**

- Your child will probably stay in the hospital overnight so your healthcare team can monitor her heartbeat.

- The next morning your child will have a device check to make sure the leads and generator are working well.

- Your child will get a pacemaker ID card that identifies the type of pacemaker she has, when it was placed, and who performed the procedure. Your child will get a permanent card in the mail in a few weeks. Make sure your child has the ID card at all times.

- **Going home:** When your child has recovered adequately from sedation and the procedure, the nurse will send you and your child home with instructions from the cardiologist.
What should I do after we go home?
When you go home, you’ll need to take care of your child’s incision, report problems, help your child gradually increase her activity, and adjust to a few basic restrictions (see “Life with a Pacemaker” on the last page).

Life with a pacemaker
Along with a better heartbeat, a pacemaker brings a few other changes to your child’s life. Follow these guidelines to help your child’s pacemaker work safely and effectively.

Take care of the incision.
• Keep the incision clean and dry. Your child may shower after 48 hours, but cover the incision with plastic so the water doesn’t hit the incision. Don’t immerse the incision in water for the first ten days or until it heals. Don’t scrub the site. Avoid using creams, ointments, or lotions on the site.
• Wear loose clothing around the site. You may place a gauze pad over the pulse generator to reduce rubbing on the incision.
• Expect some soreness for the first few days and slight swelling for about two to four weeks.

Call the doctor about problems.
Call the doctor if your child experiences any of these:
• Redness, swelling, or drainage around the wound
• A wound that separates or isn’t healing
• A fever over 101°F
• Very fast or slow heartbeat
• Swelling in her hands or ankles
• Hiccups that won’t go away

Have your child gradually increase activity.
• Follow the doctor’s instructions about keeping your child’s arm still for the first few days. Your child might need to wear a sling for a while. This may help reduce the pain.
• Check with your child’s doctor about activity during the first 6 to 8 weeks, and about when your child can return to work or school. Ask specifically about:
  – Lifting objects or driving
  – Activities that involve your child raising her arms, such as golfing, bowling, tennis, swimming, or diving
  – Activities that could bump or jar the pacemaker site, such as contact sports

Let people know
• Make sure your child has her device ID card at all times. The ID will give healthcare providers important information in an emergency. It will also be helpful if the pacemaker sets off an alarm.
• Tell healthcare providers that your child has a pacemaker before any procedures that involve needles or incisions (cuts).
• Tell your child’s dentist. The dentist can avoid using devices that produce electromagnetic fields that can interfere with the device.

Protect the pulse generator
Follow these guidelines:
• Avoid letting anything hit or rub the device. Make sure your child is careful about contact sports or other activities that may jar the pulse generator under his skin.
• Avoid strong electromagnetic fields. Your child should stay away from:
  – High-intensity power lines or radio towers
  – Magnetic resonance imaging (MRI) equipment
  – Combustion motors — don’t let your child lean over the hood of a running car, or touch the spark plug or distributor on a running car or lawn mower
  – Radio transmitters, such as CB radios, ham radios, or antennas used to control toys
Monitoring and maintenance

Here’s how the pacemaker will be maintained:

- **Follow-up appointments.** Your child’s healthcare provider will ask you to set up follow-up appointments. To check the pacemaker, your child will have various kinds of tests. For example, a pacemaker programmer — like a small portable computer — will check the device using a wand placed over the pacemaker. The doctor might also adjust the pacemaker’s settings.

- **Checking the pacemaker over the phone.** A transmitter might be used at home to send pacemaker signals to your doctor.

- **Replacing the battery.** The average battery life is 5 to 10 years. Follow-up appointments will tell your child’s healthcare provider if this is needed.

- **Replacing leads.** In rare cases, the leads can become cracked. Your child’s healthcare provider will check the leads and replace them if needed.

Talking with your child’s doctor before this procedure

The table below lists the most common potential benefits, risks, and alternatives for this procedure. Other benefits and risks may apply in your child’s unique medical situation. Talking with your child’s healthcare provider is the most important part of learning about these risks and benefits. Don’t be afraid to ask questions. All of your questions or concerns should be addressed before you agree to a recommended procedure.

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<tr>
<th>Potential benefits</th>
<th>Risks and potential complications</th>
<th>Alternatives</th>
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| • Relief of symptoms, such as dizziness or fainting | • Bleeding or infection where the pacemaker’s pulse generator was inserted  
• Problems related to the anesthetic  
• Nerve or blood vessel damage  
• Problems caused by electronic devices | • There is currently no alternative to a pacemaker if your child has a slowed heartbeat.  
• If your child’s heartbeat sometimes goes too fast or your heart muscle is weakened, your doctor may consider an implantable cardioverter device (ICD) instead of a pacemaker. |
| • Better supply of oxygen to the body, which can relieve shortness of breath or fatigue | | |

Also available in Spanish.