

Patient Device Information

COPY-CORDIS-23025 Rev.1



Cordis S.M.A.R.T. CONTROL® Vascular Stent System

What Cordis S.M.A.R.T. CONTROL® Vascular Stent System is Used For

The Cordis S.M.A.R.T. CONTROL® Vascular Stent System is for use in patients with atherosclerosis disease of peripheral arteries, including iliac and superficial femoral arteries.

Product Description

The Cordis S.M.A.R.T. CONTROL® Vascular Stent is a self-expanding stent made from nickel titanium alloy (nitinol).

Active Ingredient

- Nitinol

About Your Procedure

You may be given a mild sedative to help you relax, but you will not be put to sleep. There are two reasons for this. Firstly, most people find they experience little to no discomfort from the procedure. Secondly, your doctor may need to ask you to take a deep breath while X-rays are being taken, to improve the quality of the pictures.

Your procedure will be performed in a Cardiac Catheterization Lab or an Interventional Radiology Lab. You will lie on an X-ray table, and an X-ray camera will move over your body during the procedure. The staff will monitor your heart by attaching several small patches to your chest and using a specialized monitor.

The blood vessel at the top of your thigh is the most common site for catheter insertion and requires a very small skin incision. The area will be shaved and cleaned with an antiseptic, and you will be given a local anesthetic to numb the area. This incision will allow an introducer sheath (short tube) to be inserted into your femoral artery (the main artery of the thigh, supplying blood to the leg). Your doctor will then insert a guiding catheter (a long flexible tube) into the introducer

sheath and advance it to the peripheral or biliary artery disease. A flexible guide wire is then advanced through the guiding catheter to the narrowing in the artery. This helps carry all the necessary devices required during the stenting procedure.

Additional options for catheter insertion include an arm artery (brachial artery) on the inside of your elbow and the wrist (radial artery).

After the catheters are advanced, your doctor will inject fluid (contrast dye) through the guiding catheter into your artery to view the narrowing. Your doctor will watch the injection on an X-ray monitor, much like a TV screen. While these X-rays are being taken, your doctor may ask you to take a deep breath and hold it for a few seconds. You may also be asked to cough after the X-ray picture is completed, to help speed the removal of the contrast dye from the arteries.

The balloon tip is threaded into the narrowed area and inflated to push the plaque to the side and widen the vessel.

A small metal mesh tube (stent) may be placed in the newly opened vessel. The expanded stent provides support that helps prevent the artery from narrowing again. The stent may be coated in a drug that is released slowly over time to help prevent restenosis. The filter, sheath, catheter and balloon are removed. Pressure is applied to the small catheter insertion site to prevent bleeding.

When the procedure is done, you lie still in one position while pressure is applied to the site to stop bleeding. You generally won't have stitches, but a dressing is applied to cover the small incision site. You'll then be taken to the recovery area.

Caution

Patients with allergic reactions to nitinol (nickel titanium) may suffer an allergic response to this implant.

Before undergoing implantation of any stent, speak with your doctor if you plan to have any type of surgery

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that may require you to stop taking antiplatelet or blood thinning medications.

Adverse Effects

The risks of using the Cordis S.M.A.R.T. CONTROL® Vascular Stent System are similar to those that are associated with other standard stent procedures. For more information regarding the risks, consult your doctor.

Device Interaction and MRI Information

If you require a magnetic resonance imaging (MRI) scan, tell your doctor or MRI technician that you have a stent implant. The Cordis S.M.A.R.T. CONTROL® Vascular Stent has been shown to be MRI conditional.

Post-Procedure Care

You will be asked to lie flat for four to six hours following the procedure and to not bend your leg or arm, depending on which area your doctor used to insert the catheters. Pressure will also be placed on the area.

A vascular closure device may be used to seal the incision site in your groin or arm. You will be allowed to get up and walk around sooner if this type of device is used.

The catheter site may remain tender, swollen and bruised for a few days. There may be a small area of discoloration or a small lump in the area of the puncture. You may take acetaminophen (Tylenol, others) in the recommended dose as needed for discomfort, or other medication as prescribed by your doctor.

You may need to avoid strenuous activity and heavy lifting for 24 hours after the procedure. Please talk to your doctor about any limitations in your activities following your procedure.

Take All Medications as Instructed

After you leave the hospital, your doctor will instruct you to take a daily dose of Aspirin and another blood thinning antiplatelet drug. Your doctor will tell you how long you should continue taking these antiplatelet

drugs. It is very important that you take these medications exactly as your doctor instructs you:

- Follow your medication schedule exactly to avoid possible complications after you receive your stent. Do not miss any doses.

Call your doctor if you cannot keep taking your medications because of side effects such as rash, bleeding, or upset stomach.

Do not stop taking your prescribed medications unless you are instructed to do so by the doctor who performed your stent procedure.

Daily Activities

Day of discharge

- No driving

Modify activity for a minimum of 3 days

- No heavy lifting of anything over 5 pounds or 2.3 kgs (equivalent to a 1/2 gallon or 1.9 liters of milk)
- No pushing or pulling
- No vigorous activity or straining
 - Avoid stairs unless necessary: if necessary, take them slowly.
 - Coughing, sneezing, or straining for a bowel movement: support your groin by pressing with your palm on top of the dressing/ bandage
 - Sexual activity: check with your doctor
- No strenuous exercise
- Avoid driving unless necessary

Talk to your doctor about returning to work, which depends on your type of work, your procedure, and any medication you may be taking.

Precaution

Any serious incident that occurs in relation to the device should be reported to Cordis US Corp. and to the Therapeutic Goods Administration (TGA).

Cordis Corp.
14201 North West 60th Avenue
Miami Lakes, Florida 33014, USA
email: anz-product-complaints@cordis.com

TGA:
<http://www.tga.gov.au/reporting-problems>

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
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Discharge Information


You will be discharged to the care of your procedural doctor or primary doctor. You should be able to return to your normal activities soon.

Your doctor will ask you to return for follow-up visits. The first visit is usually two to four weeks after your stents are implanted, with follow-up visits every six months for the first year. Be sure to keep all appointments for follow-up care, including blood tests.

Legal Manufacturer

Cordis Corp. 
14201 North West 60th Avenue
Miami Lakes, Florida 33014, USA

For healthcare professionals only.

IMPORTANT INFORMATION: Prior to use, refer to the "Instructions for Use" supplied with these devices for indications, contraindications, side effects, suggested procedure, warnings, and precautions. As part of the Cordis policy of continuous product development, we reserve the right to change product specifications without prior notification. Please contact your Cordis representative for additional product availability information.
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