CASE REVIEW

Left CLI with Rest Pain / Right PTAB

Case Demographics

Age (years): 63 Gender: Male

Lesion Characteristics

Calcification: n/a Length: 234mm Runoff: 2 vessel Type (TASC): D

Clinical Presentation

- CAD with CABG, HTN, CKD stage II, HF, PVD
- Current smoker
- Hx successful right leg PTAB with the DETOUR System
- Left leg severe pain including rest pain
- Left ABI: 0.58

Procedural Information

The patient was treated with PTAB using the DETOUR System. Three TORUS Stent Grafts were placed to bypass the lesion in the following sizes: 6.0mm x 200mm, 6.7mm x 150mm, and 6.7mm x 200mm.

Anatomy

Pre

Post







INDICATIONS FOR USE: The DETOUR™ System is indicated for use for percutaneous revascularization in patients with symptomatic femoropopliteal lesions from 200mm to 460mm in length with chronic total occlusions (100mm to 425mm) or diffuse stenosis >70% who may be considered suboptimal candidates for surgical or alternative endovascular treatments. The DETOUR™ System, or any of its components, is not for use in the coronary and cerebral vasculature.

CONTRAINDICATIONS: The DETOUR™ System is contraindicated in patients with:

- · A distal common femoral artery (CFA) <7 mm in diameter.
- Increased risk of deep vein thrombosis (DVT), such as patients with a recent history of DVT, thrombophilia, and disseminated malignancy.
- · Untreated flow-limiting aortoiliac occlusive disease.
- · Lack of patent single vessel tibial runoff to ankle.
- Known coagulopathy, bleeding diathesis, or thrombocytopenia that cannot be medically managed.
- Known hypersensitivities, allergies or contraindications to: Nitinol; PTFE; aspirin; heparin; antiplatelet; anticoagulant or thrombolytic therapy; or contrast media that cannot otherwise be medically managed.

Refer to Instructions for Use for more information concerning Indications, Contraindications, Specific Anatomic Considerations, Warnings, Precautions, and Adverse Events.

CAUTION: Federal (USA) law restricts this device to sale by or on the order of a physician.

NOTE: Not all product components are available in every country. Please consult with your Endologix representative to confirm product availability.

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