Treat More with the Only 300 mm DCB*

Lutonix® Drug Coated Balloon family of PTA Catheters has the broadest size offering of all U.S. drug coated balloons (DCBs)*, with sizes on both .018" and .035", indications in AV and femoropoliteal lesions, and in lengths up to 300 mm on the .018" platform. Now with the longest DCB length available, Lutonix® 018 DCB offers a comprehensive size matrix to treat lesions without the need to exchange to an .035" guidewire.

Balloon Length (mm)

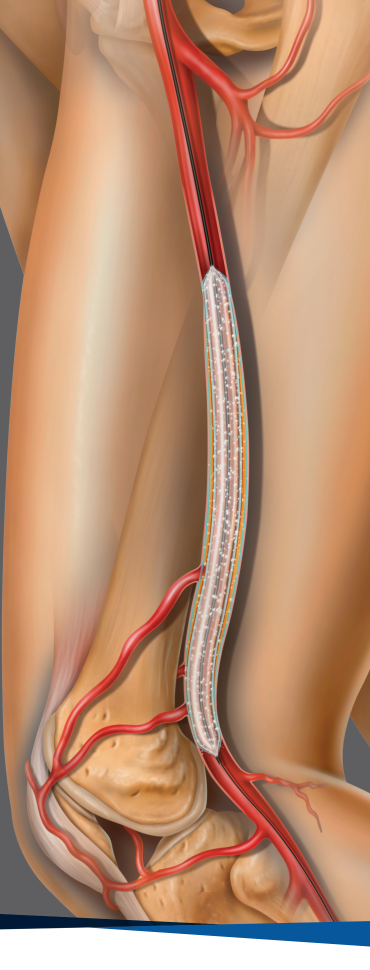
| | | 40 | 60 | 80 | 100 | 150 | 220 | 300 |
|-----------------------|---|----|----|----|-----|-----|-----|-----|
| Balloon Diameter (mm) | 4 | 4F | 4F | 4F | 4F | 4F | 4F | 5F |
| | 5 | 5F | 5F | 5F | 5F | 5F | 5F | 5F |
| | 6 | 5F | 5F | 5F | 5F | 5F | 5F | 5F |
| Ballo | 7 | 5F | 5F | 5F | 5F | 5F | 5F | |

| 40 | 60 | 80 | 100 | 150 | 220 | 300 |
|---------------|---------------|-------------|-----|-----|-----|-----|
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Lutonix® 018

Drug Coated Balloon PTA Catheter

* As of January 2020: on the US market





| Diameter (mm) | Length (mm) | Sheath Profile | 100 cm Catheter Length |
|---------------|-------------|----------------|------------------------|
| | 40 | 4F | □ LX181004404F |
| 4 | 150 | 4F | ☐ LX1810041504F |
| 4 | 220 | 4F | □ LX1810042204F |
| | 300 | 5F | ☐ LX1810043005F |
| | 40 | 5F | ☐ LX181005405F |
| 5 | 150 | 5F | ☐ LX1810051505F |
|) | 220 | 5F | ☐ LX1810052205F |
| | 300 | 5F | ☐ LX1810053005F |
| | 40 | 5F | □ LX181006405F |
| 6 | 150 | 5F | ☐ LX1810061505F |
| 0 | 220 | 5F | ☐ LX1810062205F |
| | 300 | 5F | ☐ LX1810063005F |
| 7 | 40 | 5F | ☐ LX181007405F |
| / | 150 | 5F | ☐ LX1810071505F |

| Diameter (mm) | Length (mm) | Sheath Profile | 130 cm Catheter Length |
|---------------|-------------|----------------|------------------------|
| | 40 | 4F | ☐ LX181304404F |
| | 60 | 4F | ☐ LX181304604F |
| | 80 | 4F | ☐ LX181304804F |
| 4 | 100 | 4F | ☐ LX1813041004F |
| | 150 | 4F | ☐ LX1813041504F |
| | 220 | 4F | ☐ LX1813042204F |
| | 300 | 5F | ☐ LX1813043005F |
| | 40 | 5F | ☐ LX181305405F |
| | 60 | 5F | ☐ LX181305605F |
| | 80 | 5F | ☐ LX181305805F |
| 5 | 100 | 5F | ☐ LX1813051005F |
| | 150 | 5F | ☐ LX1813051505F |
| | 220 | 5F | ☐ LX1813052205F |
| | 300 | 5F | ☐ LX1813053005F |
| | 40 | 5F | ☐ LX181306405F |
| | 60 | 5F | ☐ LX181306605F |
| | 80 | 5F | ☐ LX181306805F |
| 6 | 100 | 5F | ☐ LX1813061005F |
| | 150 | 5F | ☐ LX1813061505F |
| | 220 | 5F | ☐ LX1813062205F |
| | 300 | 5F | ☐ LX1813063005F |
| | 40 | 5F | ☐ LX181307405F |
| | 60 | 5F | ☐ LX181307605F |
| , | 80 | 5F | ☐ LX181307805F |
| 7 | 100 | 5F | ☐ LX1813071005F |
| | 150 | 5F | ☐ LX1813071505F |
| | 220 | 5F | ☐ LX1813072205F |

| I authorize the purchase of these products. | |
|---|--|
| | |
| PHYSICIAN NAME | |
| | |
| PHYSICIAN SIGNATURE | |
| | |
| | |
| REPRESENTATIVE'S NAME | |
| | |
| CONTACT PHONE NO. | |
| | |

Lutonix® 018 Drug Coated Balloon PTA Catheter

Indications for Use: The Lutonix® 018 Drug Coated Balloon PTA catheter is indicated for percutaneous transluminal angioplasty, after appropriate vessel preparation, of de novo, restenotic, or in-stent restenotic lesions up to 300 mm in length in native superficial femoral or poplited arteries with reference vessel diameters of 4-7 mm. The Lutonix® 018 Drug Coated Balloon PTA catheter is indicated for percutaneous transluminal angioplasty, after pre-dilatation, for treatment of stenotic lesions of dysfunctional native arteriovenous dialysis fistulae that are 4 mm to 7 mm in diameter and up to 80 mm in length.

Contraindications: The Lutonix® Catheter is contraindicated for use in: Patients who cannot receive recommended anti-platelet and/or anticoagulant therapy (SFA). Women who are breastfeeding, pregnant or are intending to become pregnant or men intending to father children over the next to two years. It is unknown whether paclitaxel will be excreted in human milk and there is a potential for adverse reaction in nursing infants from paclitaxel exposure. Patients judged to have a lesion that prevents complete inflation of an angioplasty balloon or proper placement of the delivery system.

exposure: * raterial guaged to that a resion that prevents complete inhation of an angiophasy balloon of proper placement of the delivery system.

Warnings: A signal for increased risk of late mortality has been identified following the use of paclitaxel-coated balloons and paclitaxel-eluting stents for femoropopliteal arterial disease beginning approximately 2.3 years post-treatment compared with the use of non-drug coated devices. There is uncertainty regarding the magnitude and mechanism for the increased late mortality risk, including the impact of repeat paclitaxel device exposure Inadequate information is available to evaluate the potential mortality risk associated with the use of paclitaxel-coated devices for the treatment of other diseases/conditions, including this device indicated for use in arteriovenous dialysis fistulae. Physicians should discuss this late mortality signal and the benefits and risks of available treatment options with their patients. Contents supplied STERILE using ethylene oxide (EQ) process. Do not use if sterile barrier is damaged or opened prior to intended use. - Do not use after 'Use by' date. - Do not use if product damage is evident. - The Lutonix" Catheter is for use in one patient only; do not reuse in another patient, reprocess or resterilize. Risks of reuse in another patient, reprocessing, or resterilization includer. Compromising the structural integrity of the device and/or device failure which, in turn, may result in patient injury, illness or death. - Creating a risk of device contamination and/or patient infection or cross-infection, including, but not limited to, the transmission of infectious disease(s) from one patient to another. Contamination of the device may lead to patient injury, illness or death. - Do not exceed the Rated Burst Pressure (RBP) recommended for this device. Balloon rupture may occur if the RBP rating is exceeded. To prevent overpressurization, use of a pressure monitoring device is recommended. - Use the recommended balloon inflation mediu

Precautions: General precautions: • The Lutonix® Catheter should only be used by physicians trained in percutaneous interventional procedures. • Consideration should be given to the risks and benefits of use in patients with a history of non-controllable allergies to contrast agents. • The safety and effectiveness of the Lutonix® Catheter have not been established for treatment in cerebral, carotid, coronary, or renal vasculature. • For SFA application, the safety and effectiveness of using more than four Lutonix drug coated balloons or a maximum drug coating quantity of approximately 15.1 mg paclitaxel in a patient has not been clinically evaluated. • For AV Fistula application, the safety and effectiveness of using multiple Lutonix drug coated balloons that deliver greater than 7.6 mg paclitaxel in a patient has not been clinically evaluated.

Potential adverse events which may be associated with a peripheral balloon dilatation procedure include, but are not limited to the following: Additional intervention: Allergic reaction to drugs, excipients or contrast medium: Amputation/loss of limb (SFA): Aneurysm or pseudoaneurysm: Arrhythmias: Embolization: Hematoma Hemorrhage, including bleeding at the puncture site: Hypotension/hypertension: Inflammation: Loss of permanent access (AVF): Occlusion: Pain or tenderness: Pneumothorax or hemothorax (SFA): Sepsis/infection: Shock: Steal Syndrome (AVF): Stroke: Thrombosis: Vessel dissection, perforation, rupture, or spasm

Although systemic effects are not anticipated, refer to the Physicians' Desk Reference for more information on the potential adverse events observed with paclitaxel.

Potential adverse events, not described in the above source, which may be unique to the paclitaxel drug coating include, but are not limited to the following: Allergic/immunologic reaction to the drug coating (paclitaxel) - Alopecia - Anemia - Blood product - transfusion - Gastrointestinal symptoms - Hematologic dyscrasia (including leukopenia, neutropenia, thrombocytopenia) - Hepatic enzyme changes - Histologic changes in vessel wall, including inflammation, cellular damage, or necrosis - Myalgia/Arthralgia - Myelosuppression - Peripheral neuropathy

Please consult product labels and instructions for use for indications, contraindications, hazards, warnings and precautions. $P_{X^{\circ n}}$

