Choose the PTA Balloon That Delivers the Versatility You Need

Choose the balloon that offers versatile sizing options designed for flexibility in tortuous anatomy and delivers better trackability and pushability compared to a competitive 0.014” PTA balloon. Choose the ULTRAVERSE® 014 PTA Balloon Dilatation Catheter.

- GEOALIGN® Marking System
- ULTRA-CROSS™ Dual Layer Hydrophilic Coating
- Reinforced Inner Lumen
- CHECKER™ Flex Points
- Dual Marker Band

** 4x120 mm ULTRAVERSE® 014 – N=5; 4x120 mm NanoCross™ 014 N=5. p<.05. Data on file. Bench test results may not necessarily be indicative of clinical performance. Different tests may yield different results.

ULTRAVERSE® 014
PTA Balloon Dilatation Catheter
**ULTRASURE® 014 PTA Balloon Dilatation Catheter**

**Indications for Use:** ULTRASURE® 014 PTA Dilatation Catheter is recommended for use in percutaneous transluminal angioplasty (PTA) of the renal, popliteal, tibial, femoral, and peroneal arteries. These catheters are also used for use in coronary arteries.

**Contraindications:** None known.

**Warnings:**
2. Do not use if sterile barrier is opened or damaged. Do not reuse, reprocess or resterilize.
3. This device has been designed for single use only. Reusing this medical device bears the risk of cross-patient contamination as medical devices – particularly those with long and small lumina, joints, and/or crevices between components – are difficult or impossible to clean once body fluids or tissues with potential pyrogenic or microbial contamination have had contact with the medical device for an indeterminable period of time. The residue of biological material can promote the contamination of the device with pyrogens or microorganisms which may lead to infectious complications. Cleaning, reprocessing and/or resterilization of the present medical device increases the probability that the device will malfunction due to potential adverse effects on components that are influenced by thermal and/or mechanical changes.
4. To reduce the potential for vessel damage, the inflated diameter and length of the balloon should approximate the diameter and length of the vessel just proximal and distal to the stenosis. When the catheter is exposed to the vascular system, it should be manipulated while under high-quality fluoroscopic observation. Do not advance or retract the catheter unless the balloon is fully deflated. Resistance is felt during manipulation, determine the cause of the resistance before proceeding. Applying excessive force to the catheter can result in tip breakage or balloon separation. Do not exceed the RBP recommended for this device. Balloon rupture may occur if the RBP rating is exceeded. To prevent over pressurization, use of a pressure monitoring device is recommended. After use, this product may be a potential biohazard. Handle and dispose of in accordance with acceptable medical practices and applicable local, state and federal laws and regulations.
5. The safety and effectiveness of the device has not been established, or is unknown, in vascular regions other than those specifically indicated.

**Precaution:** Refer to accessory IFU for potential access site warnings, precautions, and adverse events. Carefully inspect the catheter prior to use to verify that catheter has not been damaged during shipment and that its size, shape and condition are suitable for the procedure for which it is to be used. Do not use if product damage is evident. The ULTRASURE® 014 PTA Balloon Dilatation Catheters shall only be used by physicians trained in the performance of percutaneous transluminal angioplasty. The minimal acceptable sheath French size is printed on the package label. Do not attempt to pass the PTA catheter through a smaller size introducer sheath than indicated on the label. Do not remove the guide wire in situ to insert contrast through the wire lumen or perform a wire exchange. If the wire is removed while the balloon catheter is situated in tortuous anatomy, the risk of kinking the catheter is increased. Use the recommended balloon inflation medium (25% contrast medium/75% sterile saline solution). It has been shown that a 25/75% contrast / saline ratio has yielded faster balloon inflation / deflation times. Never use air or other gaseous medium to inflate the balloon. If resistance is felt during post procedure withdrawal of the catheter, the catheter is recommended to remove the balloon catheter and guidewire / introducer sheath as a single unit, and replace the previously used balloon catheter with a new balloon. Exercise caution when removing the device. Do not continue to use the balloon catheter if the shaft has been bent or kinked. Do not excessively bend, twist, or alter the shape of the device as it may compromise the integrity of the hydrophilic coating. Prior to re-insertion through the introducer sheath, the balloon should be wiped clean with wet gauze and rinsed with sterile normal saline. Avoid excessively wiping the coated portions of the device, or wiping with dry gauze, as this may damage the hydrophilic coating. Ballon re-wrapping should only occur while the balloon catheter is supported with a guide wire or stylet. This device is coated with a hydrophilic coating at the distal segment of the sheath and the balloon. Please refer to the Directions for Use section for further information on how to prepare and use this device to ensure it performs as intended. Failure to abide by the warnings in this labeling might result in damage to the hydrophilic coating, which may require intervention or result in serious adverse events. In order to activate the hydrophilic coating, it is recommended to wet the ULTRASURE® catheter with sterile saline solution immediately prior to its insertion in the body. Using different media other than the recommended solution could affect the hydrophilic coating and its performance. The BioLinx® Marking System is designed to be used as an additional reference tool to accompany the interventionist standard operation procedures. The use of fluoroscopic imaging is recommended following positioning of the catheter to the target lesion and prior to balloon deployment. Avoid using alcohol, antiseptic solutions, or other solvents to pre-treat the device because this may cause unpredictable changes in the hydrophilic coating which could affect the device safety and performance. Avoid pre-soaking devices for extended periods, as this may impact the hydrophilic coating performance. It is recommended to consider the use of anti-coagulants, anti-platelet agents, and/or vasodilators in conformance with the accepted standard of practice or institutional guidelines surrounding peripheral endovascular procedures.

**Potential Adverse Reactions:** The complications that may result from a peripheral balloon dilatation procedure include: · Additional intervention · Allergic reaction to drugs or contrast medium · Anemia or pseudoanemia · Arrhythmias · Compartment Syndrome · Embolization · Hematoma · Hemorrhage, including bleeding at the puncture site · Hypotension/hypertension · Inflammation · Occlusion · Pain or tenderness · Pneumothorax or pneumomediastinum · Septic infection · Shock · Short-term hemodynamic deterioration · Stroke · Thrombosis · Vessel dissection, perforation, rupture, or spasm.

Please consult package insert for more detailed safety information and instructions for use.