ORDERING INFORMATION

THE MYNXGRIP® Vascular Closure Device includes:
(1) Balloon catheter with integrated sealant
(1) 10ml locking syringe

To order the MYNXGRIP® Vascular Closure Device in the United States contact your local Cordis sales rep or customer service at 800.327.7714. To learn more visit cordis.com/mynx

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INDICATIONS FOR USE

The MYNXGRIP® Device is indicated for use to seal femoral arterial and femoral venous access sites while reducing times to hemostasis and ambulation in patients who have undergone diagnostic or interventional endovascular procedures utilizing a 5F, 6F or 7F procedural sheath.

WARNINGS

Do not use if components or packaging appear to be damaged or defective or if any portion of the packaging has been previously opened. DO NOT REUSE OR RERESTERILIZE. The MYNXGRIP® Device is for single use only. The balloon catheter is loaded with a single hydrogel sealant. Reuse of the device would result in no delivery of hydrogel sealant. Do not use the MYNXGRIP® Device if the puncture site is located above the most inferior border of the inferior epigastric artery (IEA) (for arterial application) and/or above the inguinal ligament based upon osseous landmarks, since such a puncture site may result in a retroperitoneal hematoma/bleed. Perform a femoral angiogram or venogram to verify the location of the puncture site. Do not use the MYNXGRIP® Device if the puncture is through the posterior wall or if there are multiple punctures, as such punctures may result in a retroperitoneal hematoma/bleed.
WHY COMPROMISE?
The MYNXGRIP® Vascular Closure Device provides secure vascular closure without the trade-offs.

Built on the proven Mynx platform, the MYNXGRIP® Vascular Closure Device offers the security of mechanical closure combined with the safety of an extravascular sealant. The MYNXGRIP® Device offers a patient-friendly closure option with no sutures, clamping, or metal implants and dissolves within 30 days leaving nothing behind but a healed artery.

DEPLOYMENT STEPS

DEPLOY BALLOON
- Achieve temporary hemostasis and position at the arteriotomy

PLACE THE SEALANT
- The Grip Tip securely adheres to the artery and the sealant fills the tissue tract

REMOVE THE DEVICE
- Platelets and blood cells collect inside the sealant’s porous matrix

FINAL RESULT
- The sealant dissolves within 30 days leaving nothing behind but a healed artery

Security of mechanical closure, Safety of an extravascular sealant.

SEALANT SCIENCE
FEATURING GRIP TECHNOLOGY

THE SEALANT
The sealant in the MYNXGRIP® Device consists of Polyethylene Glycol (PEG), a water-soluble, bio-inert, non-thrombogenic polymer, and is comprised of two configurations of PEG, the Grip Tip and the sealant.

GRIP TECHNOLOGY
Once the sealant enters the tissue tract, the body’s temperature and pH level cause the Grip Tip to soften and securely adhere to the vessel wall, effectively gripping the artery and providing active closure.

The sealant’s porous structure absorbs blood and subcutaneous fluids. The sealant swells three to four times its original size, filling the tissue tract.

The sealant actively adheres to the artery while expanding and filling the tissue tract, providing a durable hemostasis and a platform for natural healing.