

**CLOSES WITH SECURITY.**  
**LEAVES WITHOUT A TRACE.**



**MYNX CONTROL™**  
VASCULAR CLOSURE DEVICE

**Cordis**  
A Cardinal Health company

## Close with Confidence. Leave Nothing Behind.

The innovative design and predictable deployment of MYNX CONTROL™ Vascular Closure Device (VCD) delivers outstanding performance and control, for consistently secure arterial closures.

### The Science of Active Extravascular Sealing

MYNX® GRIP TIP      MYNX® Sealant



MYNX CONTROL™ VCD is comprised of two configurations of polyethylene glycol (PEG), for durable hemostasis.

#### Proven PEG Material

- **SAFE** No foreign-body reaction or scar tissue formation<sup>1</sup>
- **SYNTHETIC** Non-thrombogenic<sup>1</sup>
- **HYDROLYTIC DEGRADATION** Fully resorbs through hydrolysis—no enzymatic breakdown<sup>1</sup>

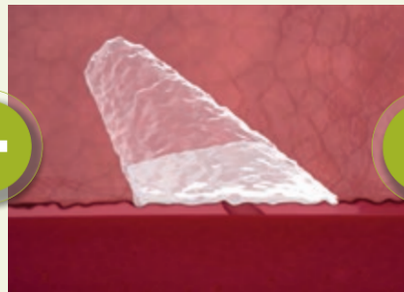
#### Dual-mode Active Sealing

**1** MYNX® GRIP TIP



- Activated by body temperature and pH
- Interlocks with contours of the vessel by actively attaching to the artery, for secure mechanical closure

**2** MYNX® SEALANT COLUMN



- Expands to 3-4 times its original size on contact with blood and subcutaneous fluids, creating a matrix structure for clot formation
- Provides further support for the MYNX® GRIP TIP

FULLY EXTRAVASCULAR CLOSURE



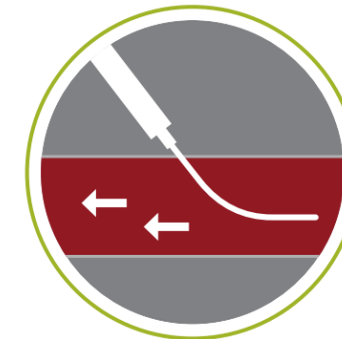
<sup>\*</sup>The sealant hydrolyzes within 30 days; the body's own healing mechanisms are the sole mechanism of action beyond 30 days. P. 2; MYNX CONTROL FDA Submission: # P040044/S079

## Secure Extravascular Closure in a wide range of clinical scenarios

With exceptional versatility, MYNX CONTROL™ VCD offers dependable closure with nothing left behind\*—even in cases where using a different vascular closure device might be unsuitable.



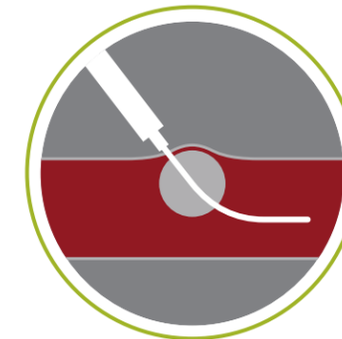
Safe for **bifurcations**<sup>2†</sup>



Useful on **antegrade** punctures<sup>3</sup>



No footplates, sutures, or metal implants to impede **reaccess**



Balloon **visualization** verifies position



<sup>†</sup>Confirm vessel size is  $\geq 5$  mm

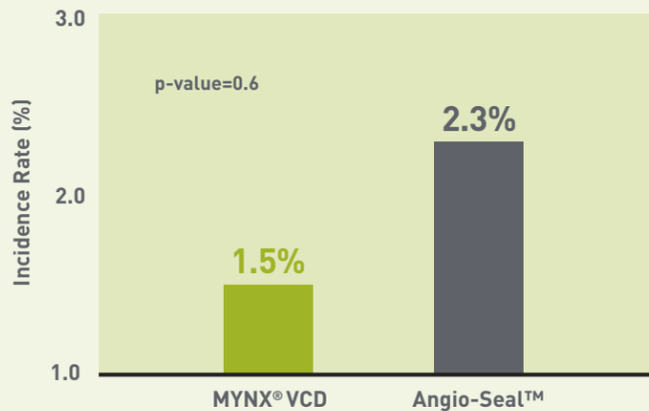
## + Safety by the Numbers.

MYNX<sup>®</sup> VCD has been clinically proven to reduce surgical complications, expedite recovery, shorten hospital stays, and increase patient comfort.<sup>2-7†</sup>

### Established Safety and Efficacy in Interventions

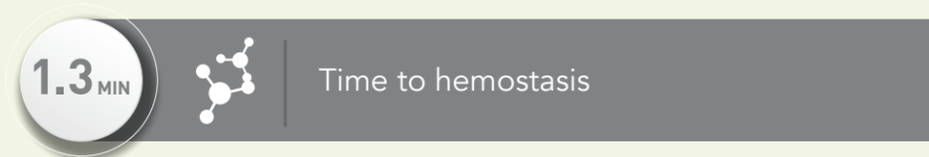
A single-center, multi-year comparative analysis involving **4,074** percutaneous coronary intervention (PCI) patients found MYNX<sup>®</sup> VCD to be equally safe and effective as Angio-Seal<sup>™</sup>, with no intra-arterial components left behind.<sup>4</sup>

#### Access-site bleeding and vascular injury<sup>4</sup>

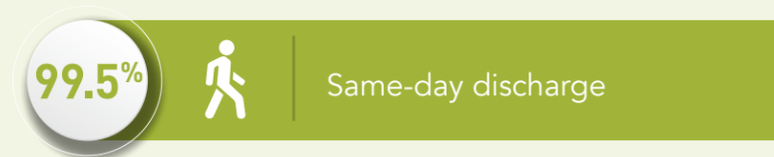


### Proven Safe in Clinical Trials and Real-world Use

In a prospective multi-center, non-randomized clinical trial (n=190) MYNX<sup>®</sup> VCD demonstrated:<sup>2,8</sup>



In a real-world cohort of 432 patients undergoing coronary angiography, MYNX<sup>®</sup> VCD demonstrated:<sup>5</sup>

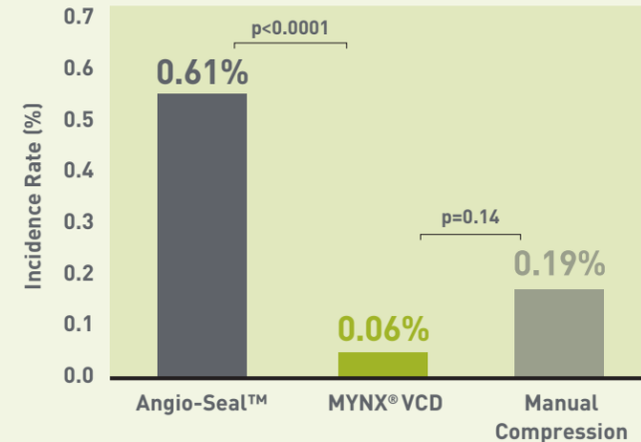


†Time to discharge eligibility as compared to manual compression. MATRIX Clinical Trial (IDE# G030182)

### Reduced Risk and Severity of Complications

In a retrospective, single-center review of **11,006** cardiac and peripheral vascular procedures, MYNX<sup>®</sup> VCD was proven to reduce the risk and severity of surgical complications following catheterization, compared to Angio-Seal<sup>™</sup> and manual compression.<sup>6</sup>

#### Rate of surgical repair<sup>6</sup>

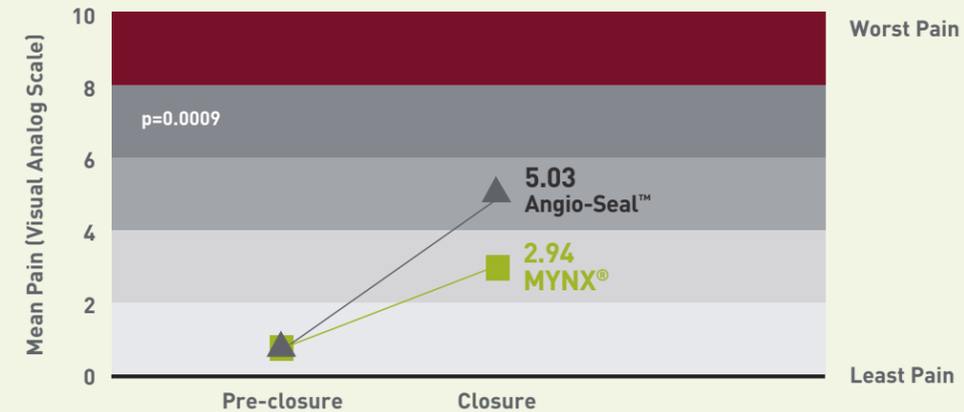


- 10x fewer secondary surgeries than Angio-Seal<sup>™</sup><sup>6</sup>
- 3x fewer secondary surgeries than manual compression<sup>6</sup>
- MYNX<sup>®</sup> VCD complications did not involve embolism or artery damage, worsening of peripheral vascular disease, or necessitate device removal<sup>6</sup>

### Increased Patient Comfort

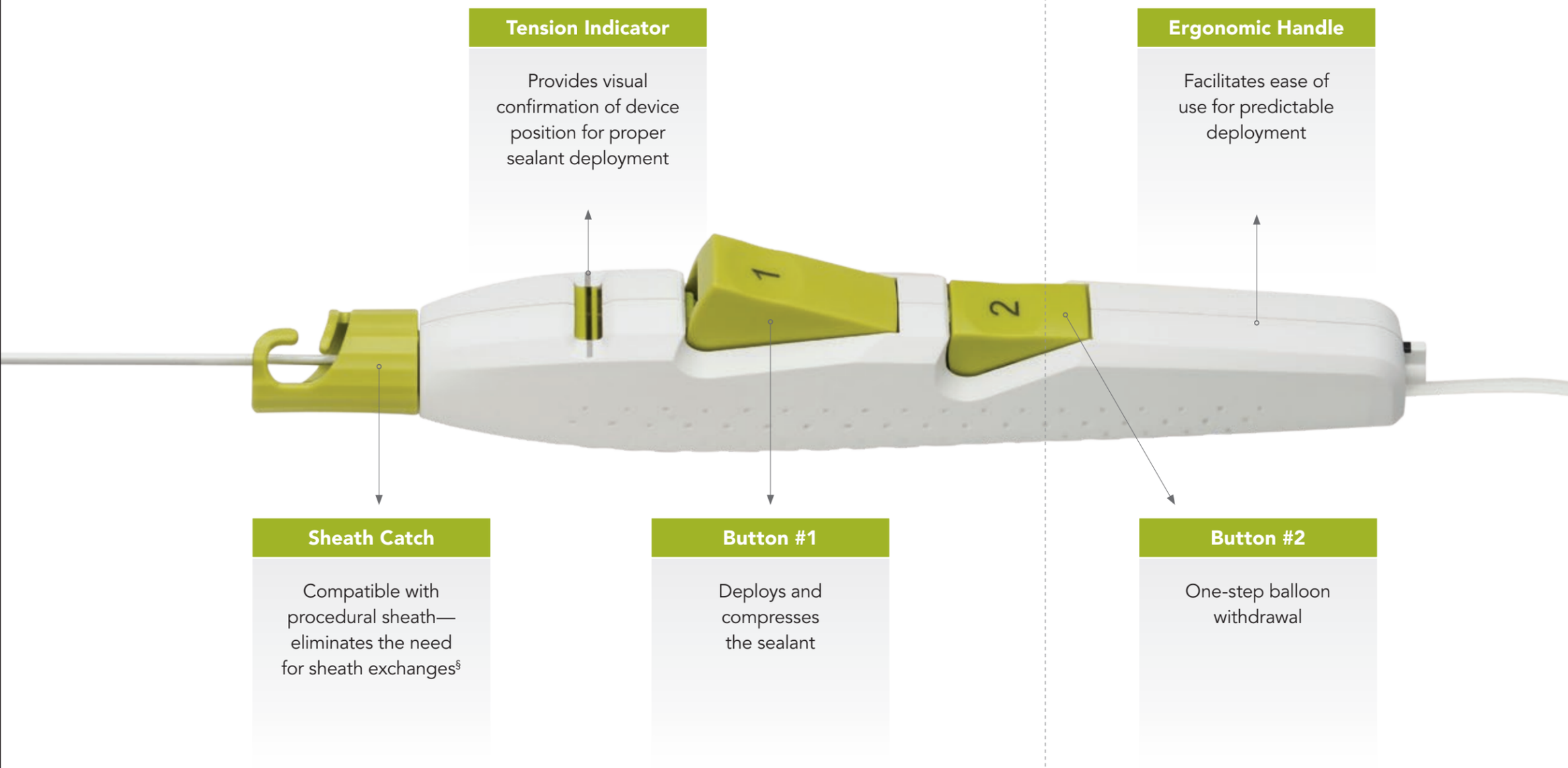
In a blinded, randomized clinical study, pain at closure and pain increase from baseline to close were significantly lower for MYNX<sup>®</sup> VCD than Angio-Seal<sup>™</sup>.<sup>7</sup>

#### Less pain than Angio-Seal<sup>™</sup>



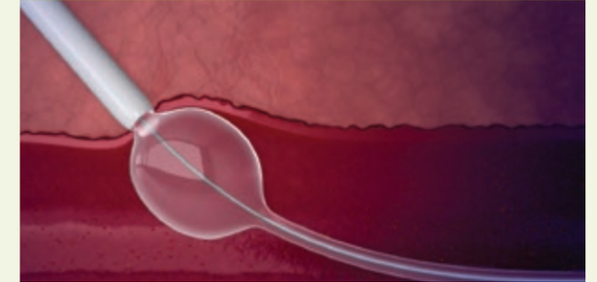
**✓ Made for Predictable Deployment.  
Designed for Ease of Use.**

The next-generation MYNX CONTROL™ Vascular Closure Device (VCD) deployment system is purpose-designed to enhance safety and deliver reliable performance.



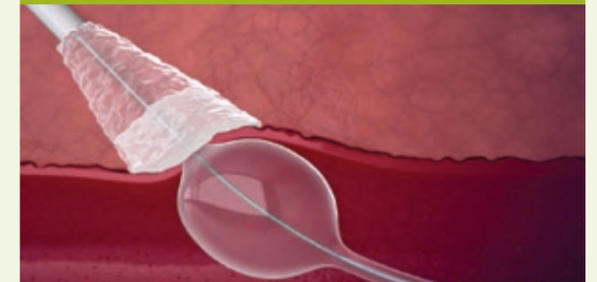
**Procedure Steps**

**1. DEPLOY THE BALLOON**



Achieve temporary hemostasis and position at the arteriotomy.

**2. PLACE THE SEALANT**



The MYNX® GRIP TIP securely adheres to the artery and MYNX® Sealant fills the tissue tract.

**3. REMOVE THE DEVICE**



Platelets and blood cells collect inside the sealant's porous matrix.

**4. FINAL RESULT**



The sealant dissolves within 30 days leaving nothing behind but a healed artery.

§MYNX CONTROL™ VCD is incompatible with Medtronic Input® Introducer (11 cm) sheaths, Cook Check-Flo® Performer® Introducer sheaths, and procedural sheaths longer than 12 cm in effective length.

## Closes with Security. Leaves Without a Trace.

MYNX CONTROL™ Vascular Closure Device (VCD) integrates dual-mode active sealing and resorbability with a next-generation delivery system to maximize predictability, safety, and ease of use.



**SECURE  
CLOSURE**



**SAFETY AND  
PATIENT COMFORT**



**EASE OF  
USE**

### Ordering Information

The MYNX CONTROL™ VCD includes:

- (1) MYNX CONTROL™ VCD including balloon catheter and integrated polyethylene glycol sealant
- (1) 10 mL locking syringe

| SIZE    | ORDER NUMBER |
|---------|--------------|
| 5F      | MX5060       |
| 6F / 7F | MX6760       |

To order the MYNX CONTROL™ VCD in the United States contact your local Cordis sales rep or customer service at 800.327.7714. To learn more visit [cordis.com/mynx](http://cordis.com/mynx).

**REFERENCES:** **1.** Scheinert D, Sievert H, Turco MA, et al. The safety and efficacy of an extravascular, water soluble sealant for vascular closure: Initial clinical results for Mynx™. *Cathet Cardiovasc Intervent.* 2007 Oct;70:627-633. **2.** MYNX Control Vascular Closure Device Instructions for Use. **3.** Pruski MJ Jr, Blachut AM, Konkolewska M, et al. MynxGrip for closure of antegrade puncture after peripheral interventions with same-day discharge. *Vasc Endovasc Surg.* 2017 Feb;51(2):67-71. **4.** Baker NC, Escarcega RO, Lipinski MJ, et al. Active versus passive anchoring vascular closure devices following percutaneous coronary intervention: a safety and efficacy comparative analysis. *J Interv Cardiol.* 2016 Feb; 29(1): 108-112. **5.** Hutchings D, Hayat A, Karunakaran A, Malik N. Success, safety, and efficacy of the Mynx femoral closure device in a real-world cohort: single-center experience. *J Invasive Cardiol.* 2016 Mar;28(3): 104-108. **6.** Noor S, Meyers S, Curl R. Successful reduction of surgeries secondary to arterial access site complications: a retrospective review at a single center with an extravascular closure device. *Vasc Endovascular Surg.* 2010 Jul;44(5):345-349. **7.** Fargen KM, Hoh BL, Mocco J. A prospective randomized single-blind trial of patient comfort following vessel closure: extravascular synthetic sealant closure provides less pain than a self-tightening suture vascular compression device. *J NeuroInterv Surg.* 2011 Sep; 3(3): 219-223. **8.** MATRIX Clinical Trial (IDE# G030182).

**INDICATIONS FOR USE:** MYNX CONTROL™ VCD is indicated for use to seal femoral arterial 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.

**PRECAUTIONS:** MYNX CONTROL™ VCD should only be used by a trained licensed physician or healthcare professional. MYNX CONTROL™ VCD should not be used in patients with a known allergy to PEG. MYNX CONTROL™ VCD should not be used with sheaths longer than 12 cm effective length or incompatible sheaths listed in Table 9 of the Instructions for Use.

**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 RESTERILIZE. MYNX CONTROL™ VCD is for single use only. The catheter is loaded with a single hydrogel sealant. Reuse of the device would result in no delivery of hydrogel sealant. Do not use MYNX CONTROL™ VCD if the puncture site is located above the most inferior border of the inferior epigastric artery (IEA) and/or above the inguinal ligament based upon bony landmarks, since such a puncture site may result in a retroperitoneal hematoma/bleed. Perform a femoral angiogram to verify the location of the puncture site. Do not use MYNX CONTROL™ VCD if the puncture is through the posterior wall or if there are multiple punctures, as such punctures may result in a retroperitoneal hematoma/bleed.

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

**IMPORTANT INFORMATION:** Prior to use, refer to the Instructions for Use supplied with this device for indications, contraindications, side effects, suggested procedure, warnings, and precautions.

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