

GET THE #1 PERIPHERAL STENT* EVERFLEX™ STENT WITH ENTRUST™ DELIVERY SYSTEM

Simple. Predictable. Precise.

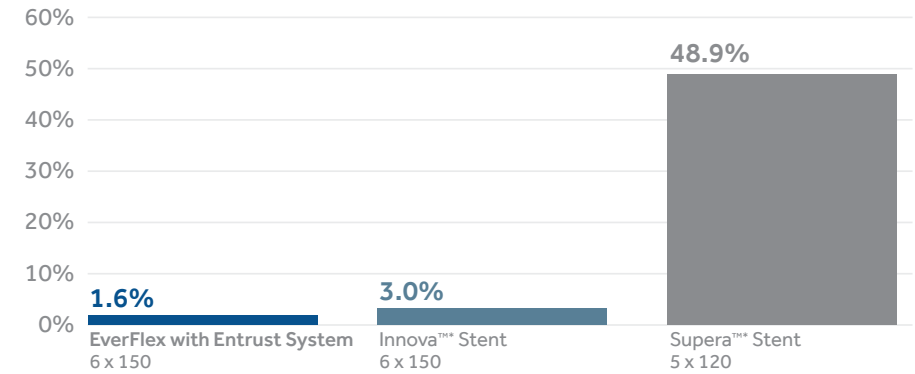
Trusted Performance

You asked for simple deployment with reduced variability — and we delivered.

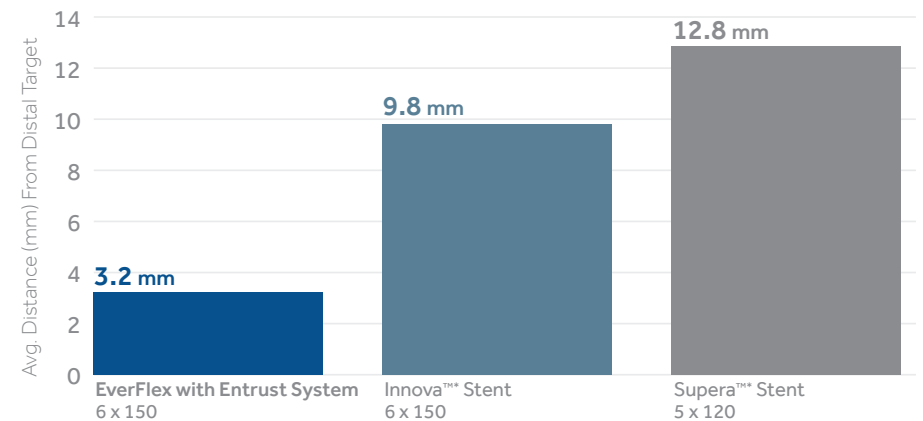
- 5 F low profile
- 0.035" guidewire compatibility
- Triaxial shaft design
- 150 cm catheter lengths

The EverFlex stent with Entrust delivery system offers the performance you've come to expect. The evidence is in the outcomes.

PERCENT STENT FORESHORTENING



DEPLOYMENT DISTANCE FROM TARGET



Test data on file at Medtronic. Results are not indicative of clinical performance.

REDESIGNED TIP

Tip attached to outer catheter eliminates risk of tip catching the stent upon removal of delivery system

5 F DELIVERY SYSTEM

- Low profile may allow for:
 - Smaller puncture site
 - Less time applying pressure¹
 - Quicker ambulatory rates²
 - Reduced vascular access complications^{3,4}

150 CM CATHETER LENGTH

Long catheter allows for an extended reach

TRIAXIAL DESIGN

Gold isolation sheath reduces friction from the system for increased accuracy and more predictable outcomes

0.035 IN GUIDEWIRE COMPATIBLE

Guidewire provides greater support for SFA procedures

EVERFLEX STENT

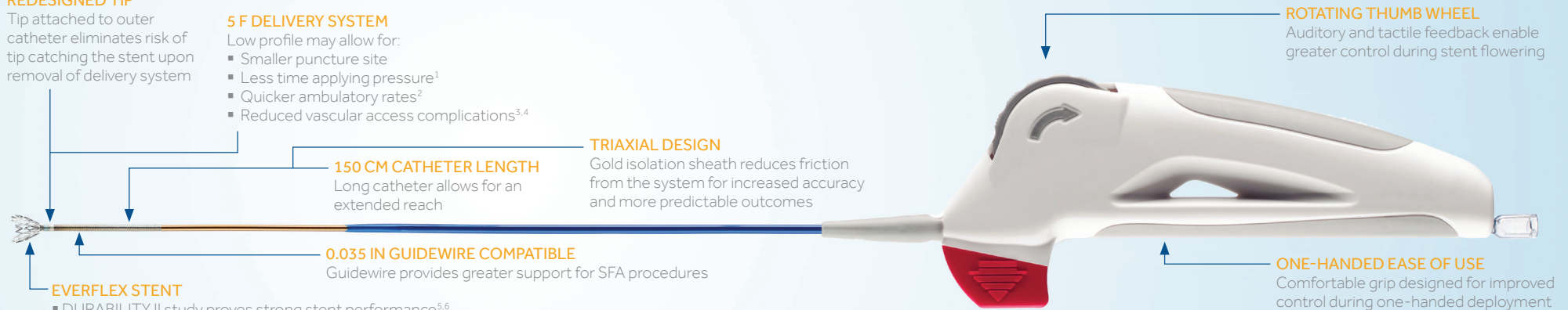
- DURABILITY II study proves strong stent performance^{5,6}
- Broad stent matrix minimizes need to place multiple stents
- Second-generation design for flexibility and durability in the SFA

ROTATING THUMB WHEEL

Auditory and tactile feedback enable greater control during stent flowering

ONE-HANDED EASE OF USE

Comfortable grip designed for improved control during one-handed deployment



Medtronic

EverFlex™ Self-expanding Peripheral Stent with Entrust™ Delivery System

Catheter			Stent dimensions		Size compatibility		
80 cm Product catalog	120 cm Product catalog	150 cm Product catalog	Unconstrained stent diameter (mm)	Unconstrained stent length (mm)	Sheath/guide compatibility (F)	Guidewire acceptance (in)	Recommended vessel size (mm)
EVD35-06-020-080	EVD35-06-020-120	EVD35-06-020-150	6	20	5	0.035	4.5–5.5
EVD35-06-040-080	EVD35-06-040-120	EVD35-06-040-150	6	40	5	0.035	4.5–5.5
EVD35-06-060-080	EVD35-06-060-120	EVD35-06-060-150	6	60	5	0.035	4.5–5.5
EVD35-06-080-080	EVD35-06-080-120	EVD35-06-080-150	6	80	5	0.035	4.5–5.5
EVD35-06-100-080	EVD35-06-100-120	EVD35-06-100-150	6	100	5	0.035	4.5–5.5
EVD35-06-120-080	EVD35-06-120-120	EVD35-06-120-150	6	120	5	0.035	4.5–5.5
EVD35-06-150-080	EVD35-06-150-120	EVD35-06-150-150	6	150	5	0.035	4.5–5.5
EVD35-07-020-080	EVD35-07-020-120	EVD35-07-020-150	7	20	5	0.035	5.5–6.5
EVD35-07-040-080	EVD35-07-040-120	EVD35-07-040-150	7	40	5	0.035	5.5–6.5
EVD35-07-060-080	EVD35-07-060-120	EVD35-07-060-150	7	60	5	0.035	5.5–6.5
EVD35-07-080-080	EVD35-07-080-120	EVD35-07-080-150	7	80	5	0.035	5.5–6.5
EVD35-07-100-080	EVD35-07-100-120	EVD35-07-100-150	7	100	5	0.035	5.5–6.5
EVD35-07-120-080	EVD35-07-120-120	EVD35-07-120-150	7	120	5	0.035	5.5–6.5
EVD35-07-150-080	EVD35-07-150-120	EVD35-07-150-150	7	150	5	0.035	5.5–6.5
EVD35-08-020-080	EVD35-08-020-120	EVD35-08-020-150	8	20	5	0.035	6.5–7.5
EVD35-08-040-080	EVD35-08-040-120	EVD35-08-040-150	8	40	5	0.035	6.5–7.5
EVD35-08-060-080	EVD35-08-060-120	EVD35-08-060-150	8	60	5	0.035	6.5–7.5
EVD35-08-080-080	EVD35-08-080-120	EVD35-08-080-150	8	80	5	0.035	6.5–7.5
EVD35-08-100-080	EVD35-08-100-120	EVD35-08-100-150	8	100	5	0.035	6.5–7.5
EVD35-08-120-080	EVD35-08-120-120	EVD35-08-120-150	8	120	5	0.035	6.5–7.5
EVD35-08-150-080	EVD35-08-150-120	EVD35-08-150-150	8	150	5	0.035	6.5–7.5

Brief Statement

Indication: The EverFlex™ Self-Expanding Peripheral Stent with Entrust™ Delivery System is intended to improve luminal diameter in the treatment of symptomatic *de novo* or restenotic lesions up to 140 mm in length in the native Superficial Femoral Artery (SFA) and/or proximal popliteal arteries with reference vessel diameters ranging from 4.5–7.5 mm.

Contraindications: Use of the EverFlex™ Self-Expanding Peripheral Stent with Entrust™ Delivery System is contraindicated in patients with known hypersensitivity to nickel titanium; patients contraindicated for anticoagulant and/or antiplatelet therapy; patients who have a lesion that prevents complete inflation of an angioplasty balloon or proper placement of the stent or stent delivery system. The EverFlex™ Self-Expanding Peripheral Stent with Entrust™ Delivery System is contraindicated for use in the carotid artery.

Potential Adverse Events: Potential adverse events that may be associated with the use of a stent in the SFA and proximal popliteal arteries include, but are not limited to: Allergic reaction, Amputation, Arterial dissection/perforation, Bleeding disorders (including GI, lymphatic), Infection (local or systemic including bacteremia or septicemia), Pseudoaneurysm, Restenosis, Stent/Vessel Thrombosis, and Surgical or endovascular intervention.

See the Instructions for Use provided with the product for a complete list of warnings, precautions, adverse events, and device information.

*EverFlex stent, U.S. only. DRG market share data for bare metal stents, November 2020.

¹ Büchler JR, Ribeiro EE, Falcão JL, et al. A randomized trial of 5 versus 7 French guiding catheters for transfemoral percutaneous coronary stent implantation. *J Interv Cardiol.* February 2008;21(1):50-55.

² Rodriguez A, Katz S. The use of the StarClose device for obtaining femoral artery hemostasis. *Vasc Endovascular Surg.* October 2011;45(7):627-630.

³ Meis A, Osada N, Schlegel PM, Fischbach R, Heindel W, Kloska SP. Sonographic follow-up of the access site after arterial angiography: Impact on the detected complication rate. *J Ultrasound Med.* September 2009;28(9):1151-1157.

⁴ Zahn R, Thoma S, Fromm E, et al. Do 5-F Catheters reduce the incidence of a pseudoaneurysm? *Int Angiol.* September 1996;15(5):257-260.

⁵ Rocha-Singh KJ, Bosiers M, Schultz G, et al. A single stent strategy in patients with lifestyle limiting claudication: 3-year results from the Durability II trial. *Catheter Cardiovasc Interv.* July 2015;86(1):164-170.

⁶ Matsumura J. DURABILITY II 12-month data. Presented at ISET 2012; Miami, FL.