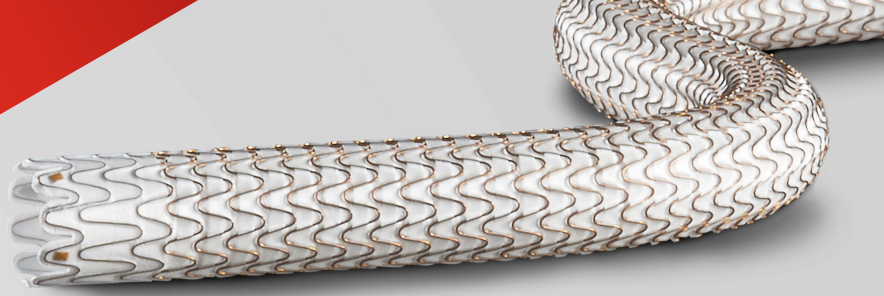


GORE® VIABAHN®

Endoprosthesis with
PROPATEN Bioactive Surface*



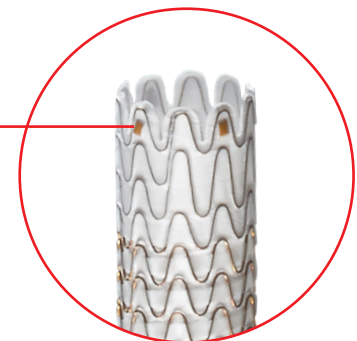
PREDICTABLE PERFORMANCE WITH ENHANCEMENTS IN ACCESS AND DELIVERY

Large diameter configurations with lower delivery profile

- Expanded accessibility through smaller sheaths.
- Smaller sheath sizes result in a lower risk of vascular access complications in select patient populations.¹

Endoprosthesis labeled diameter (mm) [†]	Recommended vessel diameter (mm) [‡]	Legacy profile (Fr)	New lower profile (Fr)
9	7.6–8.5	○ 9	○ 8
10	8.6–9.5	○ 11	○ 8
11	9.6–10.5	○ 11	○ 10
13	10.6–12.0	○ 12	○ 10 [§]

- Enhanced visualization under fluoroscopy
 - Four gold radiopaque markers on the distal and proximal ends for more confident delivery in complex cases



* As used by Gore, PROPATEN Bioactive Surface refers to Gore's proprietary CBAS Heparin Surface

[†] Labeled device diameters and lengths are nominal.

[‡] Recommended endoprosthesis compression within the vessel is approximately 5–20%.

[§] The 13 mm diameter device is not compatible with the 10 Fr COOK® FLEXOR® CHECK-FLO® Introducer.

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Together, improving life



Reference

1. Applegate RJ, Sacrinty MT, Kutcher MA, et al. Trends in vascular complications after diagnostic cardiac catheterization and percutaneous coronary intervention via the femoral artery, 1998 to 2007. *JACC: Cardiovascular Interventions* 2008;1(3):317-326.

 Consult Instructions
for Use
eifu.goremedical.com

Refer to *Instructions for Use* at eifu.goremedical.com for a complete description of all applicable indications, warnings, precautions and contraindications for the markets where this product is available. Rx_{only}

Products listed may not be available in all markets.

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W. L. Gore & Associates, Inc.
goremedical.com

Asia Pacific +65 67332882 **Australia/New Zealand** 1800 680 424 **Europe** 00800 6334 4673
United States Flagstaff, AZ 86004 800 437 8181 928 779 2771

