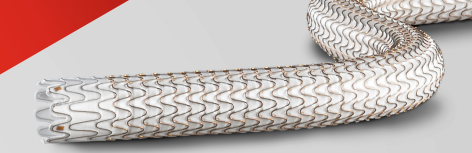
GORE® VIABAHN®
Endoprosthesis with
PROPATEN Bioactive Surface*



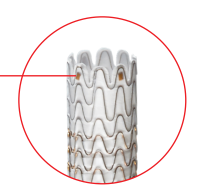
PREDICTABLE PERFORMANCE WITH ENHANCEMENTS IN ACCESS AND DELIVERY

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	O 9	0 8
10	8.6-9.5	<u> </u>	○ 8
11	9.6-10.5	O 11	O 10
13	10.6–12.0	<u> </u>	○ 10⁵

- 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%.
- $^{\rm 5}$ The 13 mm diameter device is not compatible with the 10 Fr COOK $^{\rm 6}$ FLEXOR $^{\rm 6}$ CHECK-FLO $^{\rm 6}$ Introducer.

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Reference
 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.
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