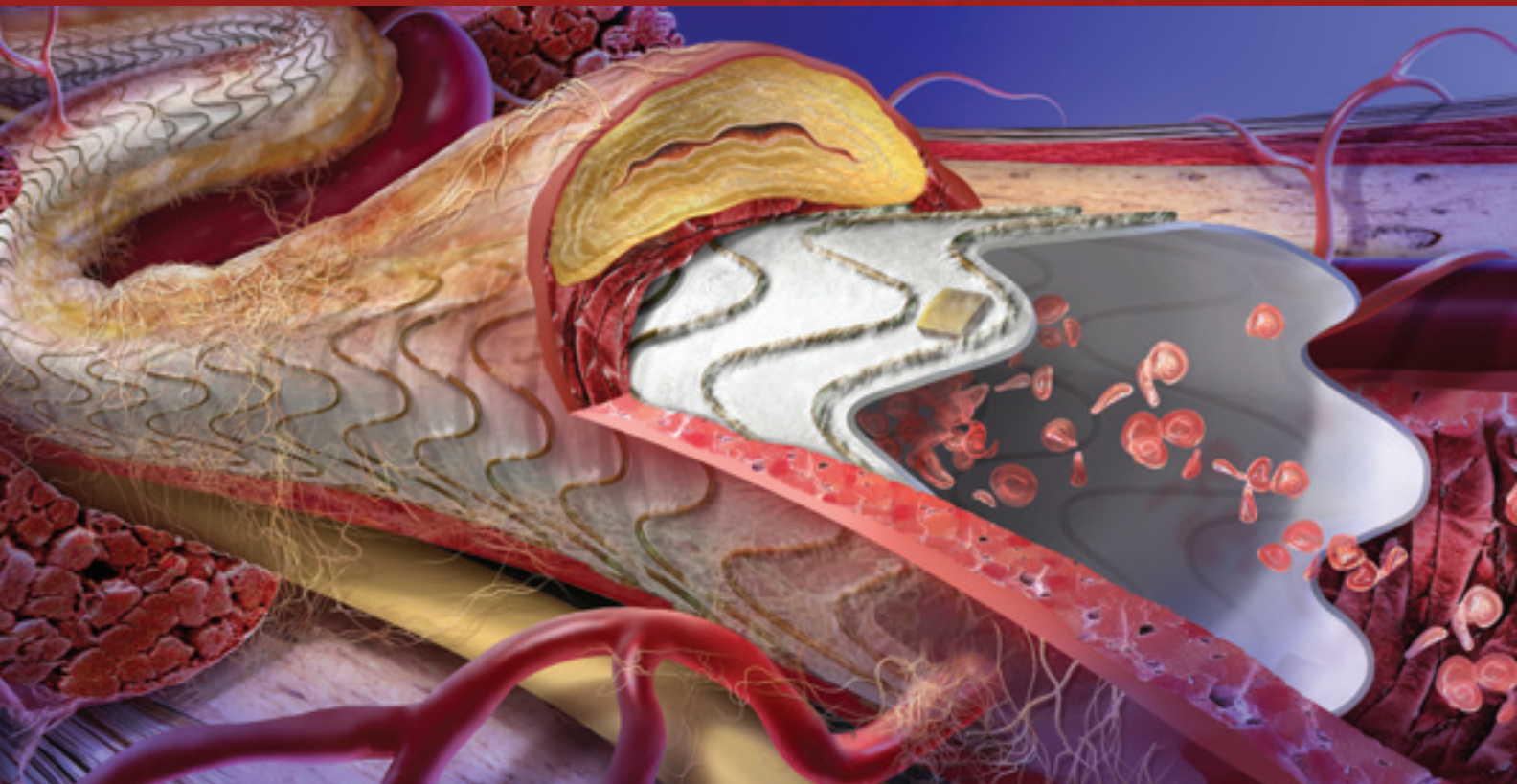


*In-stent restenosis stops here.
RELINE with confidence.*



THE CONTINUING EVOLUTION OF A REVOLUTIONARY DEVICE



PERFORMANCE
through innovation

1996

*Original GORE®
HEMOBAHN®
Endoprosthesis
introduced in Europe*

2008

*GORE® VIABAHN®
Endoprosthesis with
Heparin Bioactive Surface
introduced in Europe*

*5–8 mm devices decreased
in profile by one French size*

2009

*Laser technology
enables the new
contoured edge
at proximal end*

*9–13 mm devices
introduced with
0.035" guidewire
compatibility*

2011

*GORE® V
Endopro
Heparin
5–8 mm
decreas
by one F*

*TIP to HUB
deployment
introduced on
6–8 mm devices*

2003

*25 cm Length:
Longest stent-graft
introduced in EUROPE*

2010



*GORE® VIABAHN®
Endoprosthesis with
Bioactive Surface
introduced in profile
3 French size*



*Receives CE mark for the
treatment of symptomatic
venous obstruction*

2014

2016

*Radiopaque markers
introduced on 5–8 mm
devices in Europe*

We continue

to evolve the

GORE® VIABAHN®

Endoprosthesis,

demonstrating our

commitment to

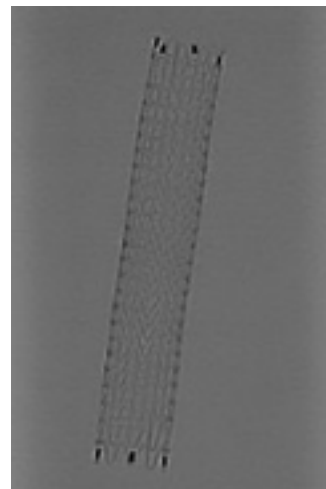
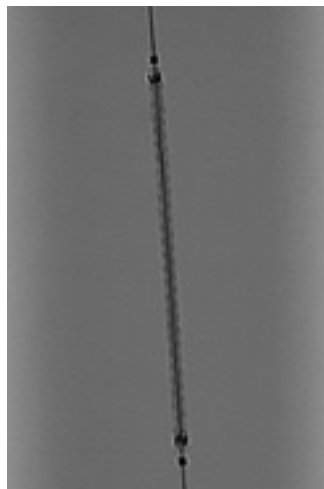
providing our

customers with

innovative products.

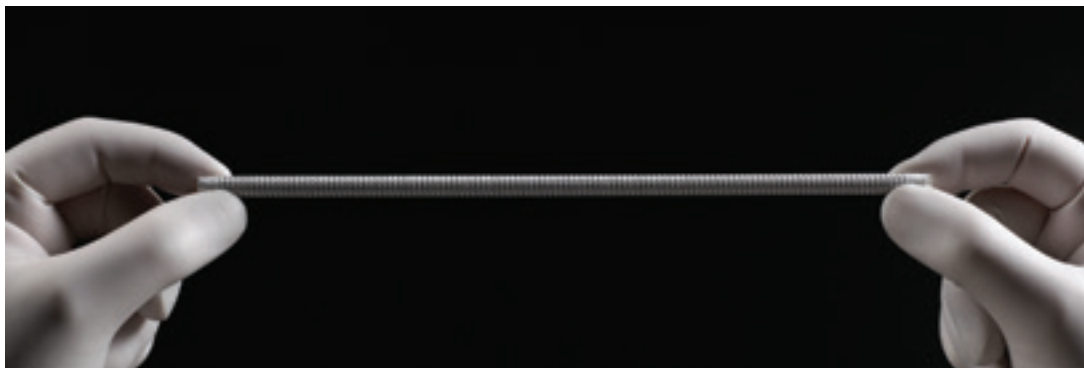
▶ **Now with Radiopaque Markers for Enhanced Visibility**

- Addition of four gold radiopaque markers bonded to the graft at each end of endoprosthesis
- 5–8 mm diameter devices incorporate this change
- Delivery system and profile unchanged



▶ **The Longest Stent-Graft for Endoluminal Bypass**

- 25 cm longest length available
- Covers more lesion with one device
- May reduce the need for overlapping devices



▶ Total Endoluminal Bypass

Cover with Confidence

Covers and excludes the diseased irregular tissue of the arterial wall

ePTFE Lining

Provides barrier to in-stent restenosis

Nitinol Stent

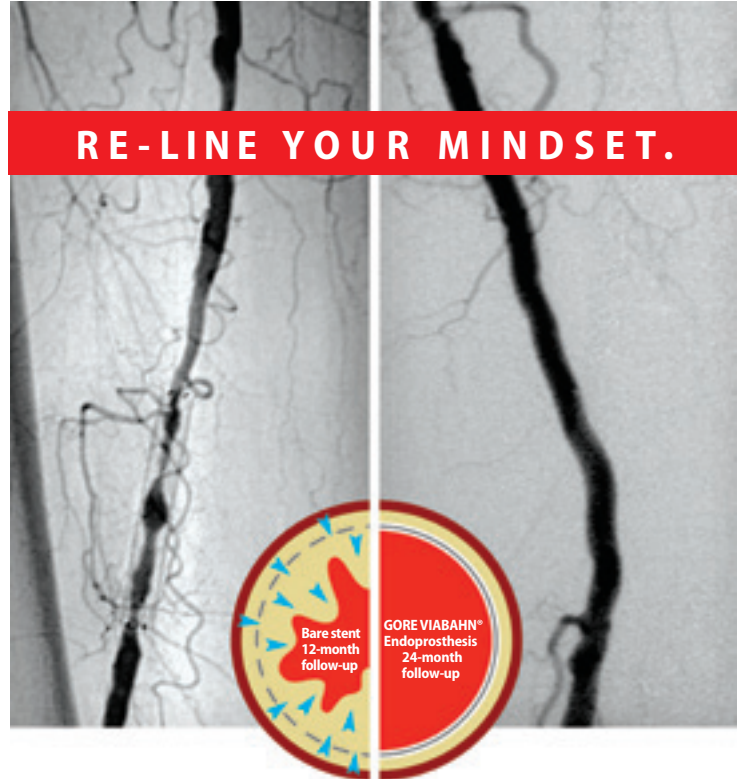
Conformable yet durable

Heparin-Bonded Surface

Intended to provide sustained thromboresistance

Lowest Profile Stent-Graft

Reduced profile delivery system makes it even easier to reach and treat challenging SFA lesions

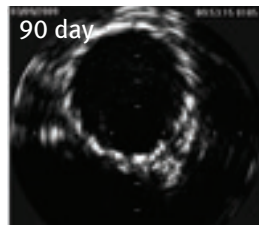
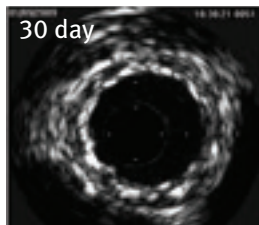
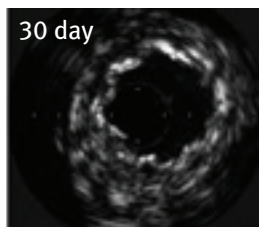


Individual Results May Vary

▶ Contoured Proximal Edge

- Precision laser trimming technology enables manufacturing change
- Excess material at the proximal edge removed
- Contoured edge may improve flow dynamics at proximal end

Canine In Vivo IVUS Examples

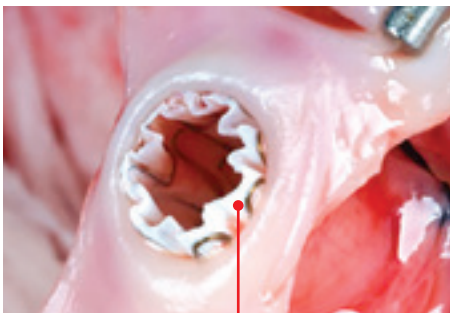


Excess material removed at the device margin of the contoured edge compared to a non-contoured edge

Contoured edge

Animal Acute Examples

Non-contoured edge



Contoured edge



Excess material removed at the device margin of the contoured edge compared to a non-contoured edge

CBAS Heparin Surface

- Intended to provide a thromboresistant surface
- Sustained bioactivity*
- Proprietary end-point covalent bonding

GORE® VIABAHN® Endoprosthesis with PROPATEN Bioactive Surface



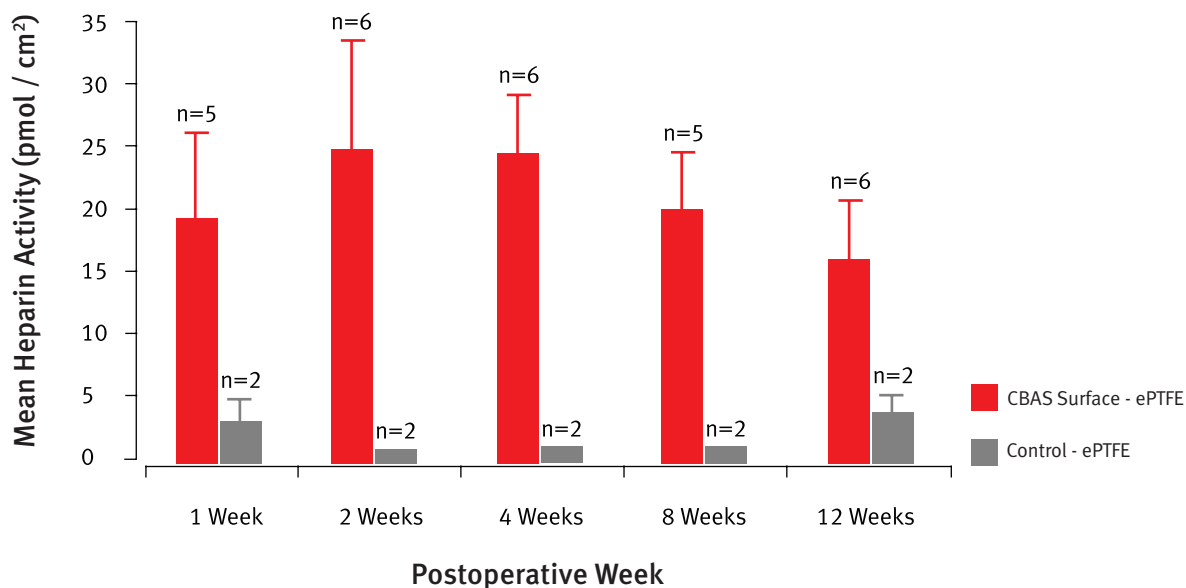
The bioactive luminal surface of a 5 mm diameter GORE® VIABAHN® Endoprosthesis with PROPATEN Bioactive Surface appears free of thrombus after two hours in an in vitro blood loop model.

Control Endoprosthesis



The non-bioactive luminal surface of a control endoprosthesis (5 mm diameter) appears covered with thrombus after 90 minutes in the same blood loop model (data on file).

Sustained Bioactivity



Long-term Heparin Activity of Explanted Heparin-bonded ePTFE Vascular Grafts in a Canine Model*

* Begovac PC, Thomson RC, Fisher JL, Hughson A, Gällhagen A. Improvements in GORE-TEX® Vascular Graft Performance by Carmeda® BioActive Surface Heparin Immobilization. *European Journal of Vascular and Endovascular Surgery* 2003;25(5):432-437.

➤ Sizing Table

TIP to HUB Device Deployment – 0.014" or 0.018" Guidewire Compatibility (With radiopaque markers)

Device Sizing		Introducer Sheath (Fr)					RECOMMENDED BALLOON DIAMETER FOR DEVICE TOUCH-UP ³ (mm)
ENDOPROSTHESIS LABELED DIAMETER ¹ (mm)	RECOMMENDED VESSEL DIAMETER ² (mm)	2.5 cm DEVICE LENGTH ¹	5 cm DEVICE LENGTH ¹	10 cm DEVICE LENGTH ¹	15 cm DEVICE LENGTH ¹	25 cm DEVICE LENGTH ¹	
5	4.0 – 4.7	6	6	6	6	6	5
6	4.8 – 5.5	6	6	6	6	6	6
7	5.6 – 6.5	7	7	7	7	7	7
8	6.6 – 7.5	7	7	7	7	7 ⁵	8

TIP to HUB Device Deployment – 0.035" Guidewire Compatibility (Radiopaque markers on 5–8 mm devices)

Device Sizing		Introducer Sheath (Fr)					RECOMMENDED BALLOON DIAMETER FOR DEVICE TOUCH-UP ³ (mm)
ENDOPROSTHESIS LABELED DIAMETER ¹ (mm)	RECOMMENDED VESSEL DIAMETER ² (mm)	2.5 cm DEVICE LENGTH ¹	5 cm DEVICE LENGTH ¹	10 cm DEVICE LENGTH ¹	15 cm DEVICE LENGTH ¹	25 cm DEVICE LENGTH ¹	
5	4.0 – 4.7	7	7	7	7	7	5
6	4.8 – 5.5	7	7	7	7	7	6
7	5.6 – 6.5	8	8	8	8	8	7
8	6.6 – 7.5	8	8	8	8	8	8
9	7.6 – 8.5	–	9	9	9	–	9
10	8.6 – 9.5	–	11 ⁴	11 ⁴	11 ⁴	–	10
11	9.6 – 10.5	–	11	11	–	–	12
13	10.6 – 12.0	–	12	12	–	–	14

¹ Labeled device diameters and lengths are nominal.

² Recommended endoprosthesis compression within the vessel is approximately 5 – 20%.

³ For the 11 and 13 mm diameter devices, balloon inflation pressure should not exceed 8 atm.

⁴ The 10 mm diameter device is compatible with the following 10 Fr introducer sheaths: CORDIS® AVANTI® Sheath Introducer, Boston Scientific SUPER SHEATH Introducer Sheath, B. Braun INTRADYN Tear-Away Introducer Sheath.

⁵ The 8 mm x 25 cm device is not compatible with the 7 Fr COOK® CHECK-FLO® FLEXOR® Sheath.



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goremedical.com

The GORE® VIABAHN® Endoprosthesis with PROPATEN Bioactive Surface is sold in some markets as the GORE® VIABAHN® Endoprosthesis with Heparin Bioactive Surface.

Products listed may not be available in all markets.

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