GORE® VIABAHN® VBX

Balloon Expandable Endoprosthesis

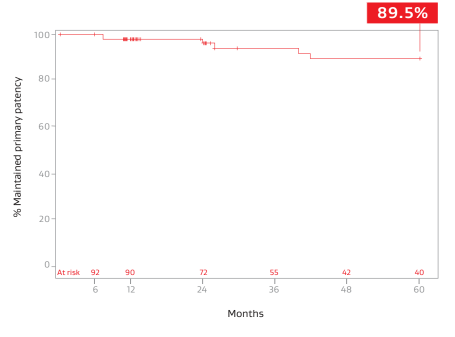
FLEXIBLE STRENGTH. PROVEN SUCCESS.

Demonstrated long-term durable clinical outcomes in complex aortoiliac occlusive disease (AIOD) treatment through **5 years.**

Sustained clinical effectiveness through 5 years:

- 89.5% primary patency and 96.1% primary assisted patency per lesion¹
- 89.1% freedom from target lesion revascularization (fTLR) per subject¹

This physician-initiated study enrolled 59 patients from 3 participating centers with patients followed out to **5 years** and beyond.



Kaplan-Meier graph of primary patency with number of lesions at risk



DURABLE PATIENT BENEFIT VS. BASELINE BEYOND 5 YEARS¹

100% of patients improved ≥ 1 Rutherford category from baseline¹

improvement in mean resting ankle-brachial index (ABI) (*P* < .001, .95 mean ABI)¹

3x improvement in median WIQ measures sustained beyond 5 years in long-term follow-up cohort¹

	Preprocedure (N = 59)	3 years (N = 39)	5 years [*] (N = 27)
Walking distance	7	25	21
Walking speed	3	10	9
Stair climbing	3	11	9

* Median follow-up of 6.6 years.

Reference

1. Holden A, Takele E, Hill A, et al. Long-term follow-up of subjects with iliac occlusive disease treated with the Viabahn VBX Balloon-Expandable Endoprosthesis. *Journal of Endovascular Therapy*. In press.



eifu.goremedical.com

INDICATIONS FOR USE IN THE U.S.: The GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis is indicated for the treatment of de novo or restenotic lesions found in Iliac arteries with reference vessel diameters ranging from 5 mm–13 mm and lesion lengths up to 110 mm, including lesions at the aortic bifurcation. The GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis is also indicated for use with thoracoabdominal and pararenal branched devices indicated with the GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis as a branch component.[†] **CONTRAINDICATIONS**: Do not use the GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis as a branch component.[†] **CONTRAINDICATIONS**: Do not use the GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis in patients with known hypersensitivity to heparin, including those patients who have had a previous incident of Heparin-Induced Thrombocytopenia (HIT) type II. Refer to *Instructions for Use* at effu.goremedical.com for a complete description of all applicable indications, warnings, precautions and contraindications for the markets where this product is available. R_{X ONV}

+ Not applicable to Reduced Profile GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis. (BXB catalogue numbers.)

Products listed may not be available in all markets.

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