GORE® EXCLUDER®

AAA Endoprosthesis

GORE® EXCLUDER® Iliac Branch Endoprosthesis

GORE® EXCLUDER® Conformable AAA Endoprosthesis

The GORE® EXCLUDER® Device family

AN EVAR PORTFOLIO YOU CAN COUNT ON



Together, improving life



GORE® EXCLUDER® AAA Endoprosthesis

Repositionable to obtain optimal seal.

Unique ability to reconstrain the proximal end and reposition for ideal placement.

More opportunities to maximize infrarenal seal.

3,274 patients through 5 years of follow-up⁺

94.7% Freedom from device-related reintervention

0.1% Migration

1.6% Type I endoleak

* Based on company-sponsored trials and registries shown on clinicaltrials.gov for currently available stent grafts

+ Results from the real-world patient population enrolled in the Global Registry for Endovascular Aortic Treatment (GREAT) (n = 3,274). To calculate the overall rates from procedure through end of study period, all subjects who could have had events, regardless of length of follow-up, were included. For outcome data, GREAT only collects site-reported serious adverse events (SAE).

+ Inclusion for renal complication rate: Subjects with renal complication were identified with MedDRA code. Of those identified with MedDRA code as having a renal complication, only those who showed the SAE occurring within 75 days of the procedure AND were reported by the site/physician as being related to the device or procedure were included in the renal complication rate.

DURABILITY you can count on

The most-studied EVAR device,* designed to provide an optimal infrarenal seal and reliable results.

Click to see device animation.





0.2% Type III endoleak

0.7% Limb occlusion

0.4% Renal complications[†]

GORE® EXCLUDER® Conformable AAA Endoprosthesis

The only on-label solution for highly angulated aortic necks up to 90° and neck lengths of at least 10 mm.

The conformability to adapt to challenging anatomies.

The capability to reconstrain, reposition and refine angulation in stages.



Patients with neck angles between 60° and 90° outcomes: 1-year follow-up*

97.9% Technical success

serious adverse events

REPORTED

98.9% Freedom from device-related

4 Type I endoleaks⁺ open repair[‡]

* For these data points, 91 patients were eligible for 1-year outcome analysis.

1-year follow-up.

+ Subject underwent open surgical repair without explant of the EXCLUDER® Conformable Device, was determined the event to be unrelated to the device or the procedure.

§ Same patient that had to convert to open repair.

II Change in AAA diameter as identified from baseline imaging taken closest to post-operative day 30 and no later than post-operative day 90.

CONTROL you can count on

The only EVAR device with angulation control, offering controlled conformability when you need it most.



Click to see device animation.





- Type III endoleaks Migrations
- Ruptures
- Stent fractures



occlusion⁵

Change in AAA diameter^{II}





48 (64.0%) NO CHANGE



+ Core Lab identified endoleaks. 3 patients with Type 1 endoleaks were resolved at 1-year follow-up, 1 patient died of unrelated causes before

GORE® EXCLUDER®

lliac Branch Endoprosthesis

The broadest treatment range for iliac preservation.

The flexibility needed in tortuous iliac anatomy, with proven kink resistance.⁴

The lowest profile (16 Fr) for enhanced vessel access and trackability.

ADVANCES you can count on

The first FDA-approved iliac branch device and the only all-in-one system. Preservation is the recommended treatment to sustain quality of life.¹⁻³

Click to see device animation.

5-year data from the U.S. IDE clinical trial.*

100% patency[†]

95.2%

External iliac artery

95.1% patency[†] Internal iliac artery

Freedom from IBE-related reintervention

98.3%

Freedom from CIAA⁺ enlargement[†]

* U.S. IDE Clinical Trial. 63 subjects with device implanted in initial cohort. 36 patients have completed 5-year follow-up. + Core Lab reported assessment for patency, endoleak, migration and CIAA enlargement (> 5 mm). Denominator is number of subjects

evaluated for primary effectiveness endpoint result with an evaluable result. ‡ On the side treated with the IBE.





ZERO REPORTED

Buttock claudication⁺ New onset erectile dysfunction Type I/III endoleaks[†] Migrations⁺

DESIGN and MATERIALS you can count on

Every device in the portfolio is meticulously designed, made from durable materials and exhaustively tested.



25 years of aortic device experience



525,000 patients treated worldwide*



Most studied⁺ EVAR device family



* Based on the number of Trunk-Ipsilateral Legs distributed for GORE® EXCLUDER® Device family as of May 2022. † Based on company-sponsored trials and registries shown on clinicaltrials.gov for currently available stent grafts.





COLLABORATION you can count on

With the GORE[®] EXCLUDER[®] Device family, you get a commitment that goes far beyond products.



Unbiased support

- A non-commissioned sales force Our focus is on outcomes.
- Committed to supporting physicians in choosing the right product for each patient.



Resources for physician education

- Gore MEDICAL MASTERY Courses Clinical knowledge through peer-to-peer collaboration.
- Gore Medical Fellows Program One-stop access to people, training and registration for key events.



Deep device expertise

- Aortic representatives support hundreds of cases per year.
- Pre-case planning and procedural consultation.



A forward-looking legacy

- Multiple clinical and investigational trials in process supporting a robust aortic pipeline.
- Continually advancing aortic treatment capabilities with innovations in device design, deployment and materials.

References

- 1. Chaikof EL, Dalman RL, Eskandari MK, et al. The Society for Vascular Surgery practice guidelines on the care of patients with an abdominal aortic aneurysm. Journal of Vascular Surgery 2018;67(1):2-77.e2.
- 2. Moll FL, Powell JT, Fraedrich G, et al. European Society for Vascular Surgery. Management of abdominal aortic aneurysms clinical practice guidelines of the European Society for Vascular Surgery. European Journal of Vascular & Endovascular Surgery 2011;41(Supplement 1):S1-S58.
- 3. Schneider DB. One-year U.S. results of GORE[®] EXCLUDER[®] Iliac Branched Endograft: Advantages and limitations. Presented at the 42nd Annual Symposium on Vascular and Endovascular Issues, Techniques, Horizons, (VEITHsymposium); November 17-21, 2015; New York, NY.
- 4. Simonte G, Parlani G, Farchioni L, et al. Lesson learned with the use of iliac branch devices: single centre 10 year experience in 157 consecutive procedures. European Journal of Vascular & Endovascular Surgery 2017;54(1):95-103.



GORE® EXCLUDER® AAA Endoprosthesis

INDICATIONS FOR USE: Trunk-Ipsilateral Leg and Contralateral Leg Endoprosthesis. The GORE® EXCLUDER® AAA Endoprosthesis is intended to exclude the aneurysm from the blood circulation in patients diagnosed with infrarenal abdominal aortic aneurysm (AAA) disease and who have appropriate anatomy as described below: Adequate iliac/femoral access; Infrarenal aortic neck treatment diameter range of 19–32 mm and a minimum aortic neck length of 15 mm; Proximal aortic neck angulation \leq 60°; Iliac artery treatment diameter range of 8–25 mm and iliac distal vessel seal zone length of at least 10 mm. Aortic Extender Endoprosthesis. The Aortic and Iliac Extender Endoprostheses are intended to be used after deployment of the GORE® EXCLUDER® AAA Endoprosthesis. These extensions are intended to be used when additional length and/or sealing for aneurysmal exclusion is desired. **CONTRAINDICATIONS:** The GORE® EXCLUDER® AAA Endoprosthesis is contraindicated in: patients with known sensitivities or allergies to the device materials: patients with a systemic infection who may be at increased risk of endovascular graft infection.

GORE® EXCLUDER® Conformable AAA Endoprosthesis

INDICATIONS FOR USE: The GORE® EXCLUDER® Conformable AAA Endoprosthesis is intended to exclude the aneurysm from the blood circulation in patients diagnosed with infrarenal abdominal aortic aneurysm (AAA) disease and who have appropriate anatomy as described below: Adequate iliac/femoral access; Infrarenal aortic neck treatment diameter range of 8-25 mm and a minimum aortic neck length of 10 mm; Proximal aortic neck angulation < 90°; Iliac artery treatment diameter range of 8-25 mm and iliac distal vessel seal zone length of at least 10 mm. Aortic Extender Endoprosthesis and Iliac Extender Endoprosthesis Components are intended to be used after deployment of the GORE® EXCLUDER® Conformable AAA Endoprosthesis. These extensions are intended to be used when additional length and/or sealing for aneurysmal exclusion is desired. CONTRAINDICATIONS: The GORE® EXCLUDER® Conformable AAA Endoprosthesis is contraindicated in: patients with known sensitivities or allergies to the device materials. All components of the GORE® EXCLUDER® Conformable Endoprosthesis contain expanded polytetrafluoroethylene (ePTFE), fluorinated ethylene propylene (FEP), nitinol (nickel-titanium alloy) and gold. Patients with systemic infection who may be at increased risk of endovascular graft infection.

GORE® EXCLUDER® Iliac Branch Endoprosthesis (IBE)

INDICATIONS FOR USE IN THE U.S.: Iliac Branch and Internal Iliac Components. The GORE® EXCLUDER® Iliac Branch Endoprosthesis (IBE) is intended to be used with the GORE® EXCLUDER® AAA Endoprosthesis or the GORE® EXCLUDER® Conformable Endoprosthesis to isolate the common iliac artery from systemic blood flow and preserve blood flow in the external iliac arteries in patients with a common iliac or aortoiliac aneurysm, who have appropriate anatomy, including: Adequate iliac/femoral access; Minimum common iliac diameter of 17 mm at the proximal implantation zone of the IBE; External Iliac artery treatment diameter range of 6.5–25 mm and seal zone length of at least 10 mm; Internal iliac artery treatment diameter range of 6.5–13.5 mm and seal zone length of at least 10 mm; Adequate length from the lowest major renal artery to the internal iliac artery to accommodate the total endoprosthesis length, calculated by adding the minimum lengths of required components, taking into account appropriate overlaps between components. GORE[®] EXCLUDER[®] Iliac Branch Endoprosthesis: Trunk-Ipsilateral Leg Component. The Trunk-Ipsilateral Leg is intended to provide proximal seal and fixation for the endovascular repair of the aneurysm. For more information on the Trunk-Ipsilateral Leg Component indications for use and deployment, see the GORE® EXCLUDER® AAA Endoprosthesis or the GORE® EXCLUDER® Conformable Endoprosthesis Instructions for Use. Contralateral Leg Endoprosthesis Component. The Contralateral Leg Endoprosthesis is intended to bridge the GORE® EXCLUDER® Device Trunk-Ipsilateral Component to the GORE® EXCLUDER® Iliac Branch Endoprosthesis following deployment of the GORE[®] EXCLUDER[®] Iliac Branch Endoprosthesis. Additionally, the Contralateral Leg Endoprosthesis is intended to be used for distal extension of the Iliac Branch Component in the external iliac artery. The Iliac Branch Component can treat external iliac artery diameters up to 13.5 mm. This ability to extend the Iliac Branch Component distally with any Contralateral Leg Endoprosthesis expands the external iliac artery treatment range up to 25 mm. For more information on the Trunk-Ipsilateral Leg and Contralateral Leg Endoprosthesis Component indications for use and deployment, see the GORE® EXCLUDER® AAA Endoprosthesis Instructions for Use. Aortic Extender and Iliac Extender Components. The Aortic and Iliac Extender Components can be used after deployment of the GORE[®] EXCLUDER[®] liac Branch and GORE[®] EXCLUDER[®] AAA Endoprostheses or the GORE[®] EXCLUDER[®] Conformable Endoprosthesis. These extensions are used when additional length and/or sealing for aneurysmal exclusion is desired. For more information on Aortic Extender and Iliac Extender indications for use and deployment, see the GORE® EXCLUDER® AAA Endoprosthesis or the GORE® EXCLUDER® Conformable Endoprosthesis Instructions for Use. CONTRAINDICATIONS: The GORE® EXCLUDER® Iliac Branch Endoprosthesis (IBE) is contraindicated in: patients with known sensitivities or allergies to the device materials. All components of the GORE® EXCLUDER® Iliac Branch Endoprosthesis (IBE), the GORE® EXCLUDER® AAA Endoprosthesis and GORE® EXCLUDER® Conformable Endoprosthesis contain ePTFE, FEP, nitinol (nickel-titanium alloy) and gold. Patients with a systemic infection who may be at increased risk of endovascular graft infection. INDICATIONS FOR USE IN CANADA: The GORE® EXCLUDER® Iliac Branch Endoprosthesis (IBE) is intended to isolate the common iliac artery from systemic blood flow and preserve blood flow in the external iliac and internal iliac arteries in patients with a common iliac or aortoiliac aneurysm, who have appropriate anatomy, including: Adequate iliac/femoral access, Minimum common iliac diameter of 17 mm at the proximal implantation zone of the IBE, External Iliac artery treatment diameter range of 6.5 – 25 mm and seal zone length of at least 10 mm, Internal iliac artery treatment diameter range of 6.5 – 13.5 mm and seal zone length of at least 10 mm.

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. Romy

Products listed may not be available in all markets.

GORE, Together, improving life, EXCLUDER, MEDICAL MASTERY and designs are trademarks of W. L. Gore & Associates, Inc. © 2023, 2024 W. L. Gore & Associates, Inc. 24AR1184-EN01 NOVEMBER 2024

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

