

Designed For Conformability

Building Confidence in the Treatment of Type B Dissection

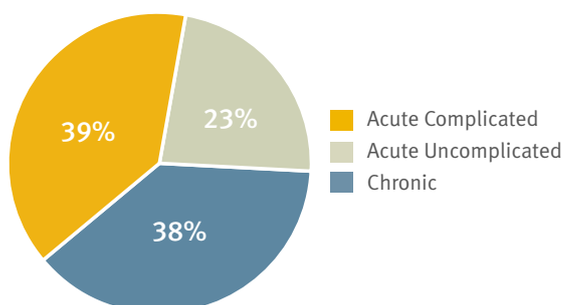


Worldwide data supports performance in the treatment of all Type B dissections

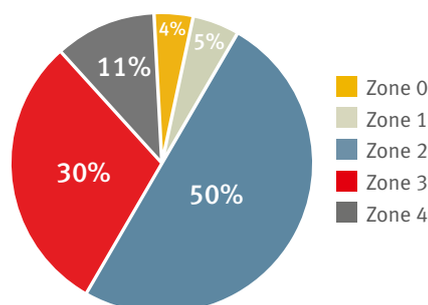
Gore GREAT Registry¹ provides worldwide experience and outcomes in real-world dissection use.

GREAT Dissection Experience

Dissection Sub-Types Treated



Proximal Landing Zone²



Adverse Event

GREAT Data³

Dissection-Related Survival	96.4 %
Procedural Survival	100 %
RTAD	0.6 %
Stroke	1.2 %
Type IA Endoleak	0 %
Device Compressions	0 %
Paraparesis / Paralysis	1.2 %
Conversion	0.6 %



1. The GREAT Registry is a prospective, observational, multicenter registry to actively track Gore commercial aortic endovascular device performance and associated patient outcomes in global markets with 10 years of follow-up.
2. The GORE® TAG® Thoracic Endoprosthesis is not indicated for the treatment of Zone 0 and Zone 1.
3. Includes all Type B dissections, events occurring within 12 months post-procedure.

Images courtesy of Nimesh Desai, MD

INDICATIONS FOR USE IN THE US: The GORE® TAG® Thoracic Endoprosthesis is intended for endovascular repair of all lesions of the descending thoracic aorta, including: Isolated lesions in patients who have appropriate anatomy, including: adequate iliac / femoral access, aortic inner diameter in the range of 16–42 mm, ≥ 20 mm non-aneurysmal aorta proximal and distal to the lesion; Type B dissections in patients who have appropriate anatomy, including: adequate iliac / femoral access, ≥ 20 mm landing zone proximal to the primary entry tear; proximal extent of the landing zone must not be dissected, diameter at proximal extent of proximal landing zone in the range of 16–42 mm. **CONTRAINDICATIONS:** Patients with known sensitivities or allergies to the device materials; patients who have a condition that threatens to infect the graft. Refer to *Instructions for Use* at goremedical.com for a complete description of all warnings, precautions, and adverse events. **INDICATIONS FOR USE UNDER CE MARK:** The GORE® TAG® Thoracic Endoprosthesis is indicated for endovascular repair of the descending thoracic aorta. **CONTRAINDICATIONS:** 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. Refer to *Instructions for Use* at goremedical.com for a complete description of all warnings, precautions, and adverse events. Rx Only

Products listed may not be available in all markets.

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To view further dissection information, visit goremedical.com/aortic/tevar



PERFORMANCE through experience