

GORE® TAG®

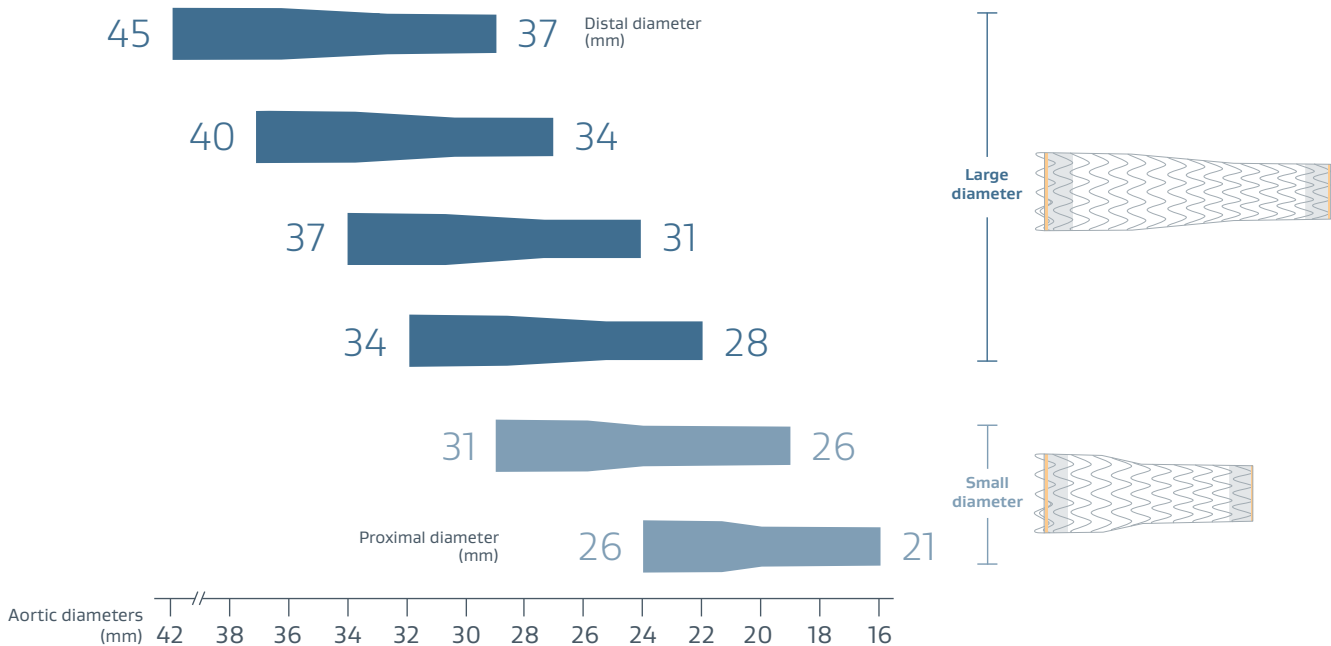
Conformable Thoracic Stent Graft
with ACTIVE CONTROL System



Expanded sizes, proven performance

Offer more patients the treatment you trust
with our large-diameter tapers.

Optimize oversizing for a broad range of tapered aortas



Together, improving life



A design against dSINE

Tapered designs may reduce distal stent graft-induced new entry (dSINE) tears associated with oversizing in a narrow true lumen.¹⁻³

dSINE is linked to increased reinterventions and rupture risk¹⁻³

Tapered devices

Catalogue number	Labeled diameter (mm)	Intended aortic diameter (mm)	Device length (cm)	Device profile (Fr)	Oversizing range (%)	Partially uncovered stent length (mm)
TGM262110	26 × 21	19.5-24/16-19.5	10	20	8-33	4
TGMR312610	31 × 26	24-29/19.5-24	10	20	7-33	4
TGM342815	34 × 28	27-32/22-26	15	22	6-27	5
TGMR373115	37 × 31	29-34/24-29	15	22	7-29	5
TGMR403415	40 × 34	31-37/27-32	15	22	6-29	6
TGM453715	45 × 37	34-42/29-34	15	24	7-32	6.5

For further information, reach out to our customer service teams.

Canada: 800 245 3416

US: 800 528 8763

References

1. Canaud L, Gandet T, Sfeir J, Ozdemir BA, Solovei L, Alric P. Risk factors for distal stent graft-induced new entry tear after endovascular repair of thoracic aortic dissection. *J Vasc Surg.* 2019;69(5):1610-1614. doi:10.1016/j.jvs.2018.07.086. OPEN ACCESS <https://www.sciencedirect.com/science/article/pii/S0741521418322560>
2. Jánosi RA, Tsagakis K, Bettin M, et al. Thoracic aortic aneurysm expansion due to late distal stent graft-induced new entry. *Catheter Cardiovasc Interv.* 2015;85(2):E43-53. doi:10.1002/ccd.25614
3. Lortz J, Leinburger F, Tsagakis K, et al. Distal stent graft induced new entry: risk factors in acute and chronic type B aortic dissections. *Eur J Vasc Endovasc Surg.* 2019;58(6):822-830. doi:10.1016/j.ejvs.2019.04.015. OPEN ACCESS <https://www.sciencedirect.com/science/article/pii/S1078588419302928>

 Consult Instructions for Use at eifu.goremedical.com 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. 

INDICATIONS FOR USE: The GORE® TAG® Conformable Thoracic Stent Graft 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.

Products listed may not be available in all markets.

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