



Indicated for  
**Aneurysm**  
**Type B Dissection**  
**Transection**

**PERFORMANCE** by design

*Pioneering TEVAR Therapy,  
Time and Time Again.*



THORACIC  
ENDOPROSTHESIS

## Time-Tested Success

For more than **16 years**, the GORE® TAG® Device has demonstrated impressive success in both clinical studies and real-world commercial use.

### More than **83,000 Devices** Distributed Worldwide

For more than a decade, we have worked alongside physicians in the evolution of the GORE® TAG® Device. Our collaboration has resulted in the distribution of more than 83,000 devices, for the treatment of more than **48,000 patients** worldwide<sup>1</sup>.

### Proven Clinical Results

The GORE® TAG® Device is supported by more than **16 years** of clinical experience.

### Most Studied Thoracic Endograft Available

With the first clinical implant occurring in 1998, the GORE® TAG® Device has been studied in **ten FDA approved clinical studies**, one European clinical trial (ADSORB), and one worldwide registry (GREAT).

### Thirty-Six Years of Experience with ePTFE Graft Material

Having pioneered ePTFE graft technology **36 years** ago, Gore continues to collaborate with physicians and scientists to create a robust and reliable design platform based on proven clinical performance.

<sup>1</sup>Data on file

**1998**

**FIRST** thoracic stent-graft  
to receive CE Mark in Europe

**2005**

**FIRST** thoracic stent-graft  
approved in US

**2008**

**FIRST** thoracic stent-graft approved in Japan

**2012**

**FIRST** thoracic stent-graft approved  
in US for isolated lesions including  
traumatic transections\*

**2013**

**FIRST** thoracic stent-graft  
approved in US for aneurysms,  
transections, *and* acute and  
chronic Type B Dissections



\* Conformable GORE® TAG® Device was approved in US for DTA aneurysms in 2011.



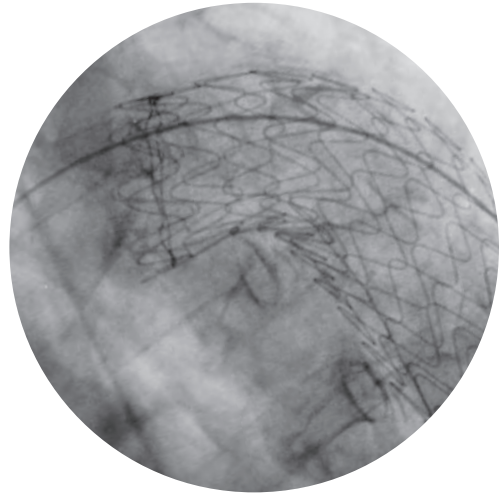
# Conformable GORE® TAG® Device is *Conformability Without Compromise*

## Designed to treat compromised aortas

- 1 No bare springs or barbs
- 2 Designed with optimal radial force to decrease the risk of intimal damage

## Highly conformable to accommodate natural anatomy

- 3 Optimized graft construct to maximize device durability and conformability
- 4 Partially uncovered stents maximize circumferential wall apposition to aid in sealing of the primary entry tear and depressurization of the false lumen while not compromising aortic blood flow
- 5 Fully covered distal end provides a transition between the stent frame and the septum, decreasing the risk of septum perforation



## Able to treat more patients

- Small diameter and tapered devices offer a large treatment range
- Broad 6 – 33% oversizing windows allow physicians to choose device with the optimal radial fit for patient anatomy
- Larger device oversizing windows engineered, tested, and proven to accommodate differences in proximal and distal landing zone diameters



## Flexible delivery system tracks in challenging anatomy

- Soft leading catheter tip for navigation through tortuous and fragile dissection anatomy
- Easy one-step deployment



## Proven compression resistance

- No reports of compression with more than 28,000 devices distributed worldwide\*
- Increased wire diameter optimizes radial force to resist compression in high flow aortas
- Nine apex stent pattern further distributes point load and contributes to long-term durability in maximum oversizing conditions
- Unique sutureless design and stent-graft construction facilitates consistent conformability throughout the device for uniform arch support

\* Through January 17, 2014. GORE® TAG® Device, *Annual Clinical Update*, 2014 not published.

**With the FDA approved indication for acute and chronic Type B dissections, Conformable GORE® TAG® Device is the *first thoracic stent-graft* approved to treat aneurysms, transections, and Type B dissections.**

For more than a decade, we have worked closely with physicians to evolve TEVAR therapies and improve patient outcomes. That's why today, Conformable GORE® TAG® Device is still a leading less-invasive treatment option.

Engineered for flexibility in tortuous anatomy, the Conformable GORE® TAG® Device provides enhanced conformability to treat the challenges associated with dissected aortas.





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**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,  $\geq 20$  mm non-aneurysmal aorta proximal and distal to the lesion; Type B dissections in patients who have appropriate anatomy, including: adequate iliac / femoral access,  $\geq 20$  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. **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](http://goremedical.com) for a complete description of all warnings, precautions, and adverse events.  $\text{K}_014$

Products listed may not be available in all markets.

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