

The Next Evolution for TIPS



Control
DIAMETER
Target
PRESSURE

***GORE® VIATORR® TIPS Endoprosthesis
with Controlled Expansion***





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Control Diameter Target Pressure


- **Lasting Diameter Control**
 - Control the diameter to reach a targeted portal pressure gradient.
 - Size and set the diameter to stay.¹
- **Single Device**
 - Select from a range of diameters² during implantation.
- **Performance**
 - Count on advanced performance that combines a legacy of proven patency³ with diameter control.

1. Based on benchtop data on file. Less than 0.25 mm increase in diameter (diameter expansion) demonstrated by a simulated 10 year period at physiologic portal pressures.

2. Labeled diameter range is between 8–10 mm.

3. Based on GORE® VIATORR® TIPS Endoprosthesis: Bureau C, Pagan JCG, Layrargues GP, et al. Patency of stents covered with polytetrafluoroethylene in patients treated by transjugular intrahepatic portosystemic shunts: long term results of a randomized multicentre study. *Liver International* 2007;27(6):742-747.

 Consult Instructions for Use

INDICATIONS FOR USE IN THE US: The GORE® VIATORR® TIPS Endoprosthesis is indicated for use in the *de novo* and revision treatment of portal hypertension and its complications such as variceal bleeding, gastropathy, refractory ascites, and / or hepatic hydrothorax. **INDICATIONS FOR USE UNDER CE MARK:** The GORE® VIATORR® TIPS Endoprosthesis is indicated for use in the treatment of portal hypertension and its complications such as: variceal bleeding refractory to, or intolerant of, conventional therapies, inaccessible varices, gastropathy, refractory ascites, and / or hepatic hydrothorax. Refer to *Instructions for Use* at goremedical.com for a complete description of all contraindications, warnings, precautions and adverse events. 



W. L. GORE & ASSOCIATES, INC.
Flagstaff, AZ 86004

+65.67332882 (Asia Pacific)
00800.6334.4673 (Europe)
800.437.8181 (United States)
928.779.2771 (United States)

goremedical.com

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