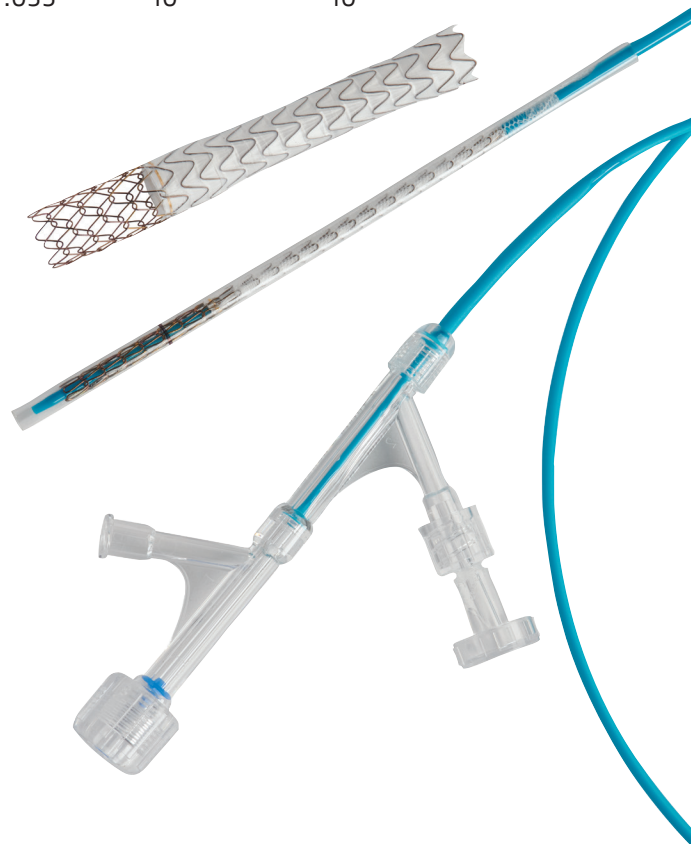


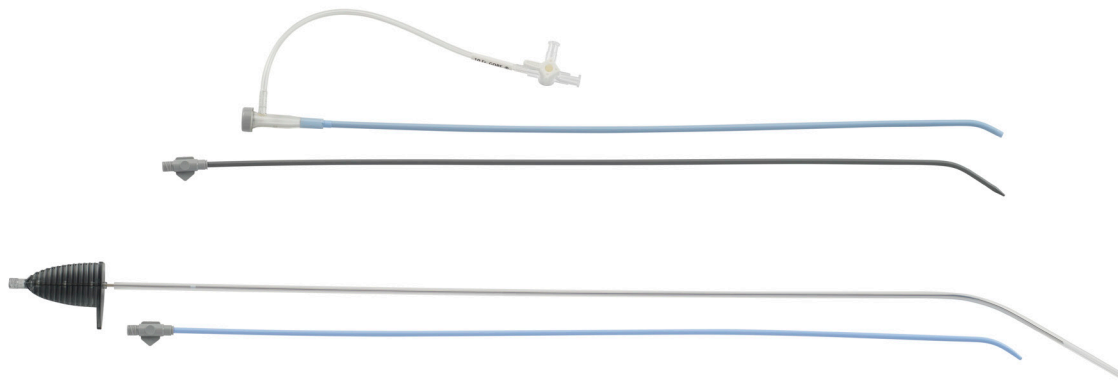
# CONFIGURATIONS

## GORE® VIATORR® TIPS Endoprosthesis with Controlled Expansion

Catalogue number	Endoprosthesis internal diameter (mm)*	Graft-lined length (cm)	Unlined length (cm)	Maximum guidewire diameter (in)	Introducer sheath size (Fr)	Maximum dilatation balloon diameter (mm)*
PTB8104275	8–10	4	2	≤ .035	10	10
PTB8105275	8–10	5	2	≤ .035	10	10
PTB8106275	8–10	6	2	≤ .035	10	10
PTB8107275	8–10	7	2	≤ .035	10	10
PTB8108275	8–10	8	2	≤ .035	10	10

\*The selected balloon diameter may be 8, 9 or 10 mm and should not exceed 10 mm. A balloon which can reach an inflation pressure of 10 ATM must be selected and should be inflated to a minimum of 10 ATM.





# CONFIGURATIONS

## GORE TIPS Set

Catalogue number	GORE TIPS Set component	Effective length (cm)	Accepts guidewire diameter (in)
TSET1016	16 Gauge needle	56	≤ .035
	10 Fr Introducer sheath	40	≤ .035
	10 Fr Dilator	47	≤ .035
	10 Fr Guiding catheter	49	≤ .035

 Consult Instructions for Use  
[eifu.goremedical.com](http://eifu.goremedical.com)

**INDICATIONS FOR USE IN THE U.S.:** 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.

**CONTRAINDICATIONS:** There are no known contraindications for this device. Refer to *Instructions for Use* at [eifu.goremedical.com](http://eifu.goremedical.com) for a complete description of all applicable indications, warnings, precautions and contraindications for the markets where this product is available.  $\text{Rx}$  Only

**INTENDED USE:** The GORE TIPS Set, GORE® TIPS Sheath and GORE TIPS Needle, are intended to be used together for percutaneous transjugular liver access during diagnostic and interventional procedures in patients undergoing a Transjugular Intrahepatic Portosystemic Shunt (TIPS) procedure. Refer to *Instructions for Use* at [eifu.goremedical.com](http://eifu.goremedical.com) for a complete description of all applicable indications, warnings, precautions and contraindications for the markets where this product is available.  $\text{Rx}$  Only

Products listed may not be available in all markets.

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