TYPE B DISSECTION MEASUREMENT/DEVICE SELECTION FORM



Confidential patient information — **Do not disclose legally protected data** The following information is required to ensure that the appropriate devices and backups are available for the procedure.

Patient ID:

Physician:

Case planning

Device selection (Refer to "Device selection form" on reverse side.)

1. Determine device diameter by identifying the aortic diameter at the proximal extent of the proximal landing zone (D). Distal diameter is not used for device selection.

2. Determine device length by selecting the device length closest to, yet longer than, the total treatment length (TTL) for the selected device diameter. If two or more devices are needed, the proximal end of the distal device must be the same diameter as the distal end of the proximal device. The proximal device must be deployed first.

3. Select the introducer sheath that corresponds to the stent graft diameter. If the outer diameter of the sheath exceeds the minimum right access diameter (A1) or the minimum left access diameter (A2), or there is excessive calcium, thrombus, tortuosity or dissection, a conduit should be used.

A1

Institution: Imaging date:

TT

A2

A2

Diameter

(D) mm

table position Diameter at proximal extent of proximal landing zone Proximal extent of device must be in non-dissected aorta

Length

(L) mm

table position

table position

Proximal neck length from proximal end of primary entry tear to left subclavian artery (LSA) or left common carotid artery (LCCA) along outer curve – L must be ≥ 2 cm

 May include dissected and non-dissected aorta

(TTL) mm

Total treatment length from LSA or LCCA along outer curve

- Device must extend at least 10 cm distal to the primary entry tear.
- Device must terminate in straight segment of descending thoracic aorta.
- Consider long segment coverage in patients with false lumen rupture.

(A1) mm

table position Minimum right access diameter (femoral, external and common iliac) Dissection, tortuosity, calcium and thrombus.

(A2) mm

table position

Minimum left access diameter (femoral, external, and common iliac) Dissection, tortuosity, calcium and thrombus.

Notes:

Gore/Patient confidential information

The following information is required to ensure that the appropriate devices and any additional devices are available for the procedure.

Patient ID:					Institution:		
Physician:					Imaging date:		
Intended device in	troduction site:	□ Right □ Left	_	lliac Femoral	🗌 Infrarenal a	aorta	Conduit

TREATMENT OPTION 1

Devices listed as implanted (proximal to distal)	Order of implantation (#1, #2, etc.)
TREATMENT OPTION 2	
Devices listed as implanted (proximal to distal)	Order of implantation (#1, #2, etc.)

Intended aortic diameters (mm)	Labeled diameter (mm)	Device length (cm)
16-19.5	21	10
19.5-24	26	10
22–26	28	10, 15
24-29	31	10, 15, 20
27-32	34	10, 15, 20
29-34	37	10, 15, 20
31-37	40	10, 15, 20
34-42	45	10, 15, 20

Intended GORE® TAG® Conformable Thoracic Stent Graft size: (Check all device sizes and indicate number of each size to be ordered.)

Device size (mm x cm)	QTY.	Catalogue number*	Device size (mm x cm)	QTY.	Catalogue number*	Device size (mm x cm)	QTY.	Catalogue number*
□ 21 x 10		TGM212110	 34 (proximal), 28 (distal) x 15 		TGM342815			
26 (proximal), 21 (distal) x 10		TGM262110	 37 (proximal), 31 (distal) x 15 		TGMR373115			
□ 26 × 10		TGM262610	□ 40 (proximal), 34 (distal) x 15		TGMR403415			
□ 31 (proximal), 26 (distal) x 10		TGMR312610	□ 45 (proximal), 37 (distal) x 15		TGM453715			
□ 28 x 10		TGM282810	28 x 15		TGM282815			
□ 31 x 10		TGMR313110	□ 31 x 15		TGMR313115	□ 31 x 20		TGMR313120
□ 34 x 10		TGM343410	□ 34 x 15		TGM343415	□ 34 x 20		TGM343420
□ 37 x 10		TGMR373710	□ 37 x 15		TGMR373715	□ 37 x 20		TGMR373720
□ 40 x 10		TGMR404010	□ 40 x 15		TGMR404015	□ 40 x 20		TGMR404020
□ 45 x 10		TGM454510	□ 45 x 15		TGM454515	□ 45 x 20		TGM454520

* Some sizes not available in all markets.

GORE® DRYSEAL Flex Introducer Sheath: (outer diameter)

Sheath size (Fr)	Device diameter (mm)	Sheath working length (cm)	QTY.	Catalogue number*
18 (6.7 mm)	21	33		GDSF1833
18 (6.7 mm)	21	65		GDSF1865
20 (7.5 mm)	26-31	33		GDSF2033
20 (7.5 mm)	26-31	65		GDSF2065
22 (8.2 mm)	34-40	33		GDSF2233
22 (8.2 mm)	34-40	65		GDSF2265
24 (8.8 mm)	45	33		GDSF2433
24 (8.8 mm)	45	65		GDSF2465
26 (9.5 mm)		33		GDSF2633
26 (9.5 mm)		65		GDSF2665

*GDSF catalogue numbers are not available in all regions; use DSF catalogue numbers instead where appropriate.

GORE® Tri-Lobe Balloon Catheter:

Device size	QTY.	Catalogue number
Aortic diameters 16–32 mm		BCM1634
Aortic diameters 26–42 mm		BCL2645

If the physician's proposed use of the device is outside the scope of the *Instructions for Use*, Gore is providing this information in response to the physician inquiry. Physician, as a medical professional, is responsible for evaluating the information provided and deciding on the use of the device.

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Consult Instructions Refer to Instructions for Use at eitu goremedical com for a com eifu.goremedical.com for a complete description of all applicable indications, warnings, precautions and contraindications for the markets where this product is available. $R_{\!X\,\text{Only}}$

