ISOLATED LESION MEASUREMENT/ DEVICE SELECTION FORM



Confidential patient information — Do not disclose legally protected data.

The following information is require	d to ensure that the appropriate	devices and backups are avai	lable for the procedure.

Patier	nt ID:					Institution:	
Physic	cian:					Imaging date:	
Гуре с	of Aneur	ysm,	/lesion:				
			Location	Measurement List single value used to select devices		CT table position/angio Specify CT frame number or specify angio	NOTES
			DIAMETER)) (
ΙΛ.	$\Lambda \Lambda \Lambda$	— А	Proximal implantation site	mm	n mm		1
		В	1 cm from proximal implantation site	mm	n mm		
		С	2 cm from proximal implantation site	mm	n mm		
		D	Maximum aneurysm/lesior	n mm	n mm		
		E	2 cm from distal implantation site	mm	n mm		
W		F	1 cm from distal implantation site	mm	ı mm		
VV	<u> </u>	−G	Distal implantation site	mm	n mm		
		Н	Right common iliac	mm	n mm		
		I	Left common iliac	mm	n mm		
		J	Right extension iliac/ femoral	mm	n mm		
LENG		К	Left extension iliac/femora	l mr	n mm	position/angio Specify CT frame number or specify angio n n n n sum sum sum sum sum sum sum sum sum su	
L ¹	to lef Proxi	nce fr t subo mal r	om aneurysm/lesion clavian	cm	cm		
			mon carotid artery	cm	cm		_//
М		h of a	/lesion aneurysm/lesion	cm	cm		7/1
N	Dista		k rom aneurysm/lesion				CUCCECTED C ADM ANCLE
	to cel		′ '	cm	cm		SUGGESTED CARM ANGLE
0	Total	treat	ment length	cm	cm		RAO
ANG							LAO
Р	Proxi			0			
Q ls th			le (if applicable) It calcium/thrombus at the p	ļ			
imp	lantation	site?			YES	□ №	NOTES
	lantation			istai	YES	□ №	
ls tr			th 10 cm or less?		☐ YES	□ №	
If yes, will both necks (proximal and distal) accommodate a single device?			ai) 	YES	□ №		
Is there a plan for coverage of the left subclavian?			ian?	☐ YES	□ NO		
If yes, is transposition or bypass clinically indicate			indicated?	YES			
Is angle less than 60°?				☐ YES			
If yes, is neck length greater than 2 cm?			k length greater than 2 cm?		☐ YES	□ №	(See reverse for device selection form.

The following information	al information n is required to ensure that the appro	priate devices a	and any additional dev	rices are available	for the procedure.
Patient ID:			Institution:		
Physician:			lmaging date:		
Intended device introd	uction site: Right Left	☐ Iliac ☐ Femo		renal aorta	☐ Conduit
TREATMENT OPTION	1				
Devices listed as impla (proximal to distal)	onted Order of implantation (#1, #2, etc.)		Intended aortic diameters (mm)	Labeled diameter (n	Device nm) length (cm)
			16–19.5	21	10
			19.5-24	26	10
			22-26	28	10, 15
			24-29	31	10, 15, 20
TREATMENT OPTION			27–32	34	10, 15, 20
Devices listed as impla (proximal to distal)	nted Order of implantation (#1, #2, etc.)		29-34	37	10, 15, 20
(proximar to distai)	(117, 112, etc.)		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 x 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 regions.

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		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 eith governedical comfor 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. $R_{\!\! X \, \text{Only}}$

