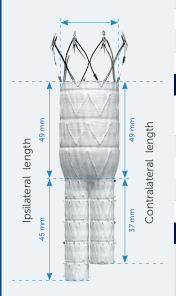




PRODUCT PORTFOLIO Proven with standard and challenging anatomies, treat more with less.

BIFURCATES

Outer diameter 22 mm, 26 mm, 30 mm, 34 mm



Caudal diameter for all aortic bifurcates 11 mm

Aortic Bifurcate Prosthesis Dimensions/Sizing Guides Treatment Delivery Delivery Ipsi Contra **Product AB Size** Range System System Length Length Code (mm) (mm) ID (F) OD (F) (mm) (mm) AB2298 22 17.0 - 19.913 94 86 14 13 94 AB2698 26 20.0-22.9 14 86

13

15

14

16

94

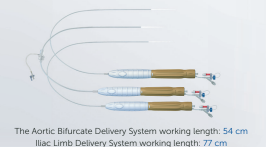
94

86

86

Aortic Extension									
Product Code	AE Size (mm)	Treatment Range (mm)	Delivery System ID (F)	Delivery System OD (F)	lpsi Length (mm)				
AE2204	22	17.0-19.9	13	14	42				
AE2604	26	20.0-22.9	13	14	42				
AE3004	30	23.0-26.9	13	14	42				
AE3404	34	27.0-31.0	15	16	42				

Bifurcate and Limb Delivery Systems



LIMBS





Iliac Limb	o/Limb	Extension Pr	osthesis	S Dimen	sions/Sizii	ng Guide
Product Code	IL Size (mm)	Treatment Range (mm)	IL Length (mm)	Delivery System OD (F)	lpsi Length (mm)	Contra Length (mm)
IL1008	10	7.0-8.9	82	12	128-156	128-147
IL1010	10	7.0-8.9	101	12	147-175	147-166
IL1012	10	7.0-8.9	120	12	166-194	166-185
IL1014	10	7.0-8.9	138	12	184-212	184-203
IL1308	13	9.0-10.9	82	12	128-156	128-147
IL1310	13	9.0-10.9	101	12	147-175	147-166
IL1312	13	9.0-10.9	120	12	166-194	166-185
IL1314	13	9.0-10.9	138	12	184-212	184-203
IL1608	16	11.0-13.9	82	12	128-156	128-147
IL1610	16	11.0-13.9	101	12	147-175	147-166
IL1612	16	11.0-13.9	120	12	166-194	166-185
IL1614	16	11.0-13.9	138	12	184-212	184-203
IL2008	20	14.0-17.9	82	12	128-156	128-147
IL2010	20	14.0-17.9	101	12	147-175	147-166
IL2012	20	14.0-17.9	120	12	166-194	166-185
IL2014	20	14.0-17.9	138	12	184-212	184-203
IL2410	24	18.0-22.0	101	13	147-175	147-166
IL2412	24	18.0-22.0	120	13	166-194	166-185
IL2414	24	18.0-22.0	138	13	184-212	184-203

For additional information and availability, please contact your Cordis representative.

AB3098

AB3498

30

34

23.0-26.9

27.0 - 31.0

For healthcare professionals only. Important information: Prior to use, refer to the "Instructions for Use" supplied with these devices for indications, contraindications, side effects, suggested procedure, warnings, and precautions.

As part of the Cordis policy of continuous product development, we reserve the right to change product specifications without prior notification. The use of INCRAFT[™] AAA Stent-Graft System requires that physicians be specifically trained in endovascular abdominal aortic aneurysm repair techniques, including experience with high resolution fluoroscopy and radiation safety. Cordis Corporation will provide training specific to the INCRAFT* AAA Stent-Graft System. Please contact your Cordis representative for additional product availability information. CORDIS, Cordis LOGO, INCRAFT are trademarks of Cordis and may be registered in the US and/or in other countries. © 2023 Cordis. All Rights Reserved. 100574919-2 01/2023

