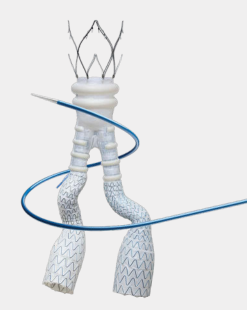


Quick Reference Guide





Indications for Use

The ALTO Abdominal Stent Graft System is indicated for treatment of patients with infrarenal abdominal aortic aneurysms having the vascular morphology suitable for endovascular repair with the device, which includes:

- Adequate iliac/femoral access compatible with vascular access techniques (femoral cutdown or percutaneous), devices, and/or accessories
- A proximal aortic landing zone for the sealing ring 7 mm below the inferior renal artery (Figures A and B)



Figure A: Proximal Landing Zone and Conicity

Figure B: Proximal Landing Zone

Indications

Indications

Indications for Use

- · An aortic sealing zone comprised of healthy aorta defined as:
 - Lack of significant thrombus > 8 mm in thickness; at any point along the aortic circumference at the level of 7 mm below the inferior renal (IR) artery (Figure C)
 - Lack of significant calcification at the level of 7 mm below the inferior renal artery
 - Conicity <10% as measured from the inferior renal artery to the aorta 7 mm below the inferior renal artery (Figure A)
 - An inner wall diameter of no less than 16 mm and no greater than 30 mm at 7 mm below the inferior renal artery
 - An aortic angle of ≤ 60 degrees (Figure D)
- · A distal iliac landing zone:
 - With a length of at least 10 mm
 - With an inner wall diameter of no less than 8 mm and no greater than 25 mm

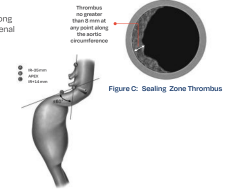


Figure D: Aortic Angle

ALTO System Components



Ovation iX[™] Ovation iX CustomSeal™ Iliac Limbs Iliac Extension Autoinjector 2 **Polymer Fill Kit**

Components

ALTO Procedure: Materials Required

Products*

- ALTO Abdominal Stent Graft Aortic Body preloaded in delivery system
- Ovation iX Iliac Limbs preloaded in delivery systems size one for each: ipsilateral and contralateral limbs (may require additional limbs if extending to external iliac or if long distance from IR to the internal iliac)
- Ovation iX Iliac Extension (as required)
- CustomSeal Polymer Fill Kit (14-minute cure time)
- Autoinjector 2
- □ 0.035" Compatible guidewire (260 cm)

Equipment

- Dever injector
- □ Ultrasound (optional)
- □ IVUS (optional)

 $^{^{\}star}$ All medical devices and tools are to be selected and utilized at the discretion of the medical professional.

ALTO Procedure: Materials Required

Additional Wires

0.035" Guidewires (2) 180 or 260 cm lengths*

□ 0.014" or 0.018", 300 cm Non-hydrophillic, snare compatible guidewires

Angiographic Catheters

- □ 0.035" Compatible radiopaque tip 5 F angiographic pigtail catheter of adequate length**
- Exchange catheters and shaped catheters for contralateral gate cannulation

Balloons

Non-compliant balloons (for leg/limb overlap)

Compliant aortic balloon

Other

- Steerable catheter
- Endovascular snare
- Micro puncture kit
- 30 ml syringe for integrated balloon inflation
- Contrast
- Heparinized saline and flushing syringes
- Time-keeping device
- Balloon expandable stent with appropriately sized balloon
- □ 14 F sheath (e.g. 45 cm)
- Embolization devices (coils)
- Percutaneous closure device (if percutaneous access)

* 180 cm guidewire for standard sheath/260 cm for integrated ALTO sheath ** Suggest using a 100 cm long pigtail when using the ALTO system

Materials Required

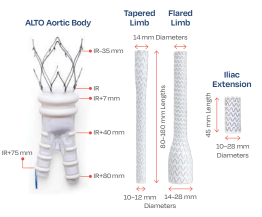
Pre-Case Sizing

- · Select sizes appropriate to the patient's anatomy.
- Confirm that devices on hand match pre-case plan implant sizes.

	Aortic	lliac Limb /	Extension	
Stent Graft Diameter (mm)	Aortic ID* (mm)	Maximum Aortic Vessel Diameter at Anchors (mm) 35 mm above the lowest renal	Labeled Diameter (mm)	Native Iliac Vessel ID Range (mm)
20	16-17	≤24	10	8-9
23	18-20	≤26	12	10-11
26	21-23	≤29	14	12-13
29	24-26	≤32	16	14-15
34	27-30 ≤35		18	16-17
* At the inter	ided proximal	22	18-20	

28

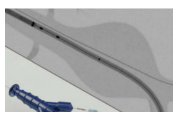
21-25



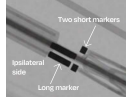
System Set-Up



• Orient the graft outside the patient in the planned orientation.



 Using fluoroscopic guidance, advance the delivery system until the radiopaque markers are about 1 cm proximal to the intended landing site.



 Orient the aortic body by rotating the entire delivery system as a unit to align the delivery system markers with the long marker so the ipsilateral (or deployment knobs) are in desired orientation.

System Set-Up

Aortic Body Deployment



Mid Crown Deployment

- To uncover the graft, stabilize the handle and retract the delivery system outer sheath until the sheath retraction knob meets the handle.
- Verify that aortic body radiopaque markers are approximately 1 cm proximal to the intended landing site.
- Stabilize the delivery system and deploy first segment of proximal stent (mid crown): rotate first deployment knob ¼ turn counterclockwise and then steadily pull knob and attached wire from handle.

Aortic Body Deployment



Integrated Balloon

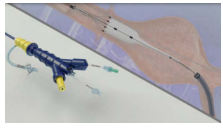
- Remove white cap from balloon injection port on handle.
- Inflate balloon with standard 4:1 saline and contrast mixture. Adhere to recommended inflation volumes (see table).
- Completely deflate the integrated balloon by pulling vacuum on the inflation syringe.

Aortic Body Stent Graft Diameter (mm)	Recommended Integrated Balloon Inflation Volume (mL)	
20		
23	5	
26		
29	10	
34	10	

Aortic Body Deployment Aortic Body Deployment

Aortic Body Deployment

Proximal Crown Deployment



- Correct for parallax and repeat angiogram (if adjustments are made).
- Position bottom of radiopaque markers at floor of inferior renal ostium.



- Stabilize the delivery catheter while retracting the angiographic catheter away from proximal stent.
- Deploy proximal crown. Stabilize the delivery system. Rotate second deployment knob ¼ turn counterclockwise, then pull knob and attached wire from handle.

Adaptive Sealing Technology

Polymer Mixing



- Open the fill polymer kit stopcocks and mix the polymer using a minimum of twenty full syringe strokes.
 - Confirm each syringe has bottomed out prior to beginning the 20 full uninterrupted strokes.
- Start a timer for 14 minutes when mixing is complete.
 - After mixing, polymer should be discarded if not used within 2 minutes.
- Transfer contents to syringe with green band, then expel to the minimum fill syringe volume appropriate to the aortic body size. Close stopcocks. Remove green tear tab and disconnect fill syringe.

Adaptive Seal

Adaptive Seal

Adaptive Sealing Technology

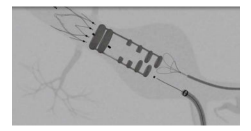
Polymer Fill



- Remove green fill cap from the polymer injection port on the catheter handle and attach fill syringe.
- Push Autoinjector 2 over the syringe plunger and lock into place by rotating 90 degrees, until you hear an audible click. Fill polymer will begin filling aortic body in ~1 minute.
- Retract aortic body guidewire until the stiff to floppy transition is in the ipsilateral leg of the aortic body. Under fluoroscopy, intermittently observe filling of graft with radiopaque fill polymer.

NOTE: Alert anesthesia that polymer fill will begin.

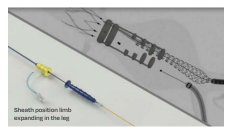
Contralateral Limb Deployment



- Cannulate the contralateral gate with guidewire. The integrated crossover lumen may be used to facilitate the process using a maximum 0.018" guidewire through the crossover lumen port on the handle.
- Advance wire until the wire exits the crossover lumen at the second ring of the contralateral leg.
- Snare the guidewire and externalize on the contralateral side.
 - Insert a 5 F sheath over the contralateral crossover guidewire. Insert a buddy guidewire into the aortic body leg and advance proximally.
 - Retract wire within crossover lumen through the ipsilateral side and remove sheath on the contralateral side.

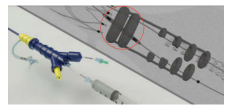
Contralateral Limb

Contralateral Limb Deployment



- Advance marker catheter over the wire to measure required length for the contralateral limb.
- Remove marker catheter and advance limb delivery system over guidewire and into the aortic body leg.
- Position iliac limb radiopaque markers between the third and fourth half ring of the aortic body leg to ensure appropriate overlap.
- Retract sheath to deploy iliac limb while maintaining catheter handle position.
- Maintain position of sheath and retract catheter handle to position nosecone in end of delivery system outer sheath.
- If maintaining the sheath, reposition sheath tip to desired location and continue to retract the blue handle until entire inner catheter is removed.
 - Alternatively, remove entire delivery system from vasculature and replace with access sheath of choice.

Ballooning the Proximal Sealing Ring



Aortic Body Stent Graft Diameter (mm)	Recommended Integrated Balloon Inflation Volume (mL)	Maximum Integrated Balloon Inflation Volume (mL)
20		7
23	5	8
26		12
29	10	15
34	10	19

Balloon Proximal Seal

- · Verify 14-minute polymer cure time has elapsed.
- Remove the green autoinjector from fill syringe.
- Readvance aortic body guidewire through aortic body delivery system.
- Using fluoroscopy, position balloon radiopaque markers proximal to the primary sealing ring and distal to secondary ring.
- Manually inflate balloon with 4:1 saline and contrast mixture to recommended volume (see table).
- Completely deflate balloon by pulling vacuum and confirm seal angiographically.

NOTE: In order to maximize polymer moldability, it may be necessary to balloon the proximal sealing ring prior to the contralateral limb placement.

Ballooning

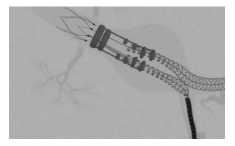
Delivery System Removal

Aortic Body Delivery System Removal



- Rotate the third deployment knob ¼ turn counterclockwise and pull knob and wire from handle.
- Advance the delivery system sheath to the first ring of the aortic body. Stabilize the delivery system and retract inner catheter.
- When the nosecone is within the aortic body, stop retracting.
- Slightly retract the sheath 2-4 cm, then continue to retract the inner catheter handle to reseat the nosecone into the outer sheath.
- To use the integrated sheath, while maintaining guidewire position, move entire delivery system to desired position. Retract handle to remove the inner catheter from the outer sheath.
 - Alternatively, remove entire delivery system from vasculature.

Ipsilateral Limb Deployment



- Advance marker catheter over the wire to measure required length for the ipsilateral limb.
- Remove marker catheter and advance limb delivery system over guidewire and into the aortic body leg.
- Position iliac limb radiopaque markers between the third and fourth half ring of the aortic body leg to ensure appropriate overlap.
- Retract sheath to deploy iliac limb while maintaining catheter handle position.
- Maintain position of sheath and retract catheter handle to
 position nosecone in end of delivery system outer sheath.
- If maintaining the sheath, reposition sheath tip to desired location and continue to retract the blue handle until entire inner catheter is removed.
 - Alternatively, remove entire delivery system from vasculature and replace with access sheath of choice.

Ipsilateral Limb

System Removal

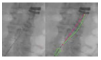
Confirm Aneurysm Exclusion and System Removal



- Advance angiographic catheter through the integrated sheath, through the aortic body and into the suprarenal space.
- Perform angiogram to confirm aneurysm exclusion and graft patency.
- · If iliac extensions are required, proceed with deployment.
- Remove angiographic catheter.
- Remove delivery system sheath from the aortic guidewire.

Best Practices to Optimize Proximal Seal







1. Accuracy

- a. Adjust C-Arm for parallax, magnify at renals, and ensure radiopaque markers are in a line.
- b. Repeat angiogram if imaging orientation is changed in any way.
- c. Position bottom of radiopaque markers at the floor of the renal ostium.

2. Apposition (prior to and during the polymer fill)

- Retract angiographic catheter before deploying second yellow deployment knob for proximal crown deployment.
- b. Retract aortic body guidewire until the stiff-to-floppy transition of the wire is in the ipsilateral leg of the aortic body.
- c. Ensure no tension on delivery system.

3. Confirm Seal

- a. From 1 to 14 minutes post polymer mix, visually check that the edges of the sealing ring are flat vs rounded and are of uniform density (no apparent infolding).
- b. Change orientation to see circular sealing ring.

Best Practices

Best Practices

Best Practices to Optimize Proximal Seal



4. Balloon

- a. Integrated balloon can be used if the aortic body system is in place.
- b. Non-compliant balloon use in the proximal sealing zone can only occur after aortic body delivery system has been removed.
- c. If unable to achieve seal post balloon deployment, follow next steps.

NOTE: Balloon has greatest effect on molding customized seal immediately following 14 minutes post-polymer mix (with reduced efficiency after 25 minutes). By 30 minutes post-polymer fill, the ability of the balloon to mold the seal is minimized. Prioritize ballooning over limb deployment.



Ballooning Best Practices—Aortic Body

Timing (post polymer mix)	Type of Balloon
0 to 14 minutes	No ballooning allowed
≥14 minutes	Integrated balloon (or compliant if integrated balloon has been removed or damaged)
>30 minutes	Ballooning much less effective

• Keep the lower margin of the balloon proximal to the aortic body bifurcation

Non-compliant balloons only used after aortic body delivery system has been removed

Best Practices

Graft Selection

ALTO Aortic Body Graft Selection

Proximal Graft Nominal Size (mm)	Min–Max Vessel Diameter at Sealing Ring (mm)	Max Aortic Vessel Diameter at Anchors (mm)	Inner "Bore" Diameter (mm)	Labeled Length (mm)	ALTO Delivery Catheter ID/OD (F)
20	16-17	≤24	15		
23	18-20	≤26	17		
26	21-23	≤29	20	80	13/15
29	24-26	≤32	23		
34	27-30	≤35	26		

Note: Standard rounding rules apply to vessel diameter measurements. Indications are for proximal aortic neck inner wall diameter of no less than 16 mm and no greater than 30 mm.



Limb Graft Selection

Iliac Limb: Diameter Sizing

Distal Graft Nominal Size (mm)	Native Iliac Vessel ID Range (mm)	Min – Max Vessel Diameter (mm)	Ovation iX Delivery Catheter ID/OD (F)
10	8–9	8–9	
12	10-11	10-11	10/12
14	12-13	12-13	
16	14–15	14–15	11/10
18	16-17	16-17	11/13
22	18-20	18-20	12/14
281	21-25	21-25	13/15

Note: Standard rounding rules apply to vessel diameter measurements. Indications are for inner wall diameter of no less than 8 mm and no greater than 25 mm².

- 1. 28 mm limb diameter available with the Ovation iX Iliac Stent Graft.
- 2. Ovation iX Iliac Stent Graft allows treatment up to 25 mm iliac vessel.

Graft Selection



US Model Numbers

ALTO Aortic Body

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Length (mm)
TV-AB2080-N			20	
TV-AB2380-N			23	
TV-AB2680-N	13	15	26	80
TV-AB2980-N			29	
TV-AB3480-N			34]

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141080-J					80	125
TV-IL1410100-J					100	145
TV-IL1410120-J	10	12	14	10	120	165
TV-IL1410140-J					140	185
TV-IL1410160-J					160	205

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141280-J					80	125
TV-IL1412100-J					100	145
TV-IL1412120-J	10	12	14	12	120	165
TV-IL1412140-J					140	185
TV-IL1412160-J					160	205
TV-IL141480-J					80	125
TV-IL1414100-J					100	145
TV-IL1414120-J	10	0 12	14	14	120	165
TV-IL1414140-J					140	185
TV-IL1414160-J					160	205
TV-IL141680-J					80	125
TV-IL1416100-J					100	145
TV-IL1416120-J	11	13	14	16	120	165
TV-IL1416140-J					140	185
TV-IL1416160-J					160	205

US Model Numbers

US Model Numbers

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141880-J					80	125
TV-IL1418100-J					100	145
TV-IL1418120-J	11	13	14	18	120	165
TV-IL1418140-J					140	185
TV-IL1418160-J					160	205
TV-IL142280-J					80	125
TV-IL1422100-J					100	145
TV-IL1422120-J	12	14	14	22	120	165
TV-IL1422140-J					140	185
TV-IL1422160-J					160	205
TV-IL142880-J					80	125
TV-IL1428100-J					100	145
TV-IL1428120-J	13	15	14	28	120	165
TV-IL1428140-J]				140	185
TV-IL1428160-J					160	205

Ovation iX Iliac Extension

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)
TV-EX101045-J	10		10	10	
TV-EX121245-J	10	12	12	12	
TV-EX141445-J	10		14	14	
TV-EX161645-J	11	13	16	16	45
TV-EX181845-J	11	13	18	18	
TV-EX222245-J	12	14	22	22	
TV-EX282845-J	13	15	28	28	

Autoinjector 2

CustomSeal Polymer Fill Kit

Catalog Number TV-AI01-N

Catalog Number TV-CS14-G

US Model Numbers

ALTO Aortic Body

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Length (mm)
TV-AB2080-L			20	
TV-AB2380-L			23	
TV-AB2680-L	13	15	26	80
TV-AB2980-L			29	
TV-AB3480-L			34	

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141080-I	10		14	10	80	125
TV-IL1410100-I					100	145
TV-IL1410120-I		12			120	165
TV-IL1410140-I					140	185
TV-IL1410160-I					160	205

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141280-I		12	14	12	80	125
TV-IL1412100-I					100	145
TV-IL1412120-I	10				120	165
TV-IL1412140-I					140	185
TV-IL1412160-I					160	205
TV-IL141480-I		12	14	14	80	125
TV-IL1414100-I					100	145
TV-IL1414120-I	10				120	165
TV-IL1414140-I					140	185
TV-IL1414160-I					160	205
TV-IL141680-I		13	14	16	80	125
TV-IL1416100-I	11				100	145
TV-IL1416120-I					120	165
TV-IL1416140-I					140	185
TV-IL1416160-I					160	205

OUS Model Numbers

Ovation iX Iliac Limb

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)	Stent Graft Length with Aortic Body (mm)
TV-IL141880-I		13	14	18	80	125
TV-IL1418100-I					100	145
TV-IL1418120-I	11				120	165
TV-IL1418140-I	-				140	185
TV-IL1418160-I					160	205
TV-IL142280-I	12	14	14	22	80	125
TV-IL1422100-I					100	145
TV-IL1422120-I					120	165
TV-IL1422140-I					140	185
TV-IL1422160-I					160	205
TV-IL142880-I	13	15	14	28	80	125
TV-IL1428100-I					100	145
TV-IL1428120-I					120	165
TV-IL1428140-I					140	185
TV-IL1428160-I					160	205

Ovation iX Iliac Extension

Catalog Number	Delivery System Inner Diameter (F)	Delivery System Outer Diameter (F)	Stent Graft Proximal Diameter (mm)	Stent Graft Distal Diameter (mm)	Stent Graft Length (mm)
TV-EX101045-I	10		10	10	
TV-EX121245-I	10	12	12	12	
TV-EX141445-I	10		14	14	
TV-EX161645-I	11	13	16	16	45
TV-EX181845-I	11	13	18	18	
TV-EX222245-I	12	14	22	22	
TV-EX282845-I	13	15	28	28	

Autoinjector 2

CustomSeal Polymer Fill Kit

Catalog Number TV-AI01-L

Catalog Number TV-CS14-F

OUS Model Numbers

INDICATIONS FOR USE: The ALTO Abdominal Stent Graft System is indicated for treatment of patients with infrarenal abdominal aortic aneurysms having the vascular morphology suitable for endovascular repair with the device, which includes the following:

- Adequate iliac/femoral access compatible with vascular access techniques (femoral cutdown or percutaneous), devices, and/or accessories,
- · A proximal aortic landing zone for the sealing ring 7 mm below the inferior renal artery.
- · An aortic sealing zone comprised of healthy aorta defined as:
 - Lack of significant thrombus >8 mm in thickness; at any point along the aortic circumference at the level of 7 mm below the inferior renal artery,
 - Lack of significant calcification at the level of 7 mm below the inferior renal artery,
 - Conicity < 10% as measured from the inferior renal artery to the aorta 7 mm below the inferior renal artery,
 - An inner wall diameter of no less than 16 mm and no greater than 30 mm at 7 mm below the inferior renal artery, and
 - An aortic angle of ≤ 60 degrees
- · A distal iliac landing zone:
 - With a length of at least 10 mm, and
 - With an inner wall diameter of no less than 8 mm and no greater than 25 mm.

CONTRAINDICATIONS: The system is contraindicated for use in patients who have a condition that threatens to infect the graft and in patients with known sensitivities or allergies to the device materials including polytetrafluoroethylene [PTFE], polyethylene glycol [PEG]-based polymers, contrast agents, fluorinated ethylene propylene [FEP], titanium, nickel, platinum, or iridium.

Refer to Instructions for Use for more information concerning Indications, Contraindications, Specific Anatomic Considerations, Warnings, Precautions, and Adverse Events.

CAUTION: Federal (USA) law restricts this device to sale by or on the order of a physician.

NOTE: Not all product components are available in every country. Please consult with your Endologix representative to confirm product availability.

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