









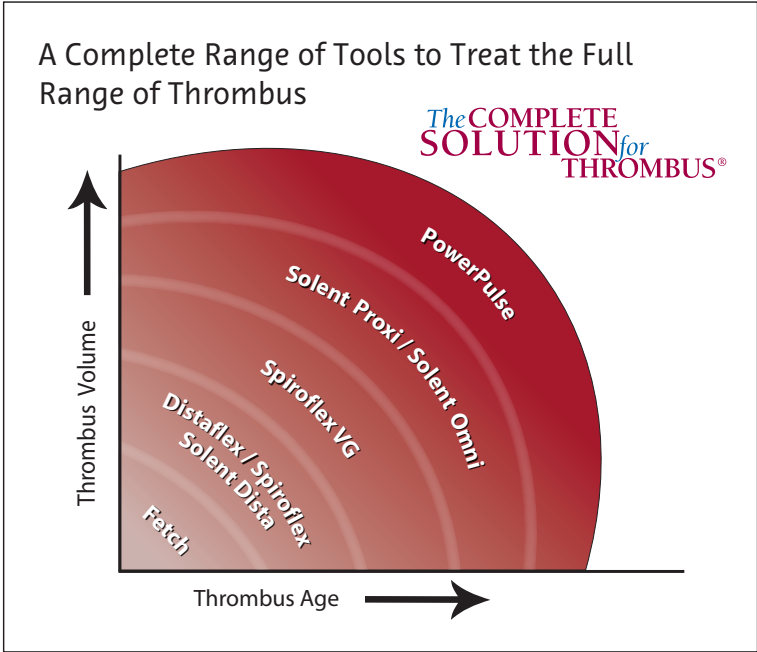




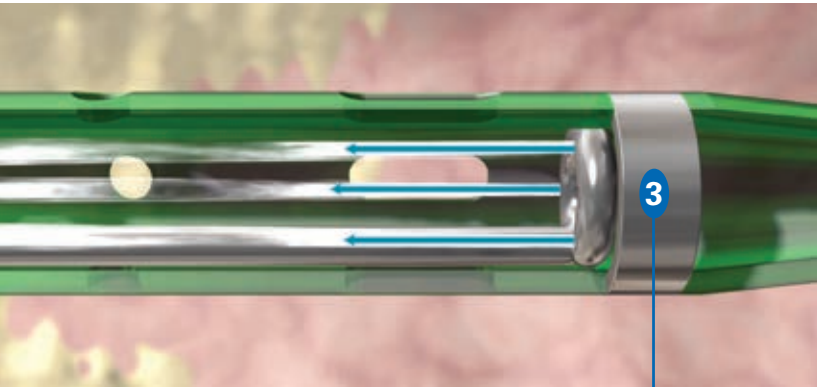
Model	Indication	Delivery Platform	Minimum Vessel Diameter	Catheter Length	Catheter Diameter	Guide Wire	Guide Catheter	Introducer Sheath	Power Pulse® Delivery Enabled	Guidewire Swappable	Contrast Injection Port	Flow Rate	Maximum Run Times		<i>Order Information</i>
													Total Run Time	Run Time with Blood Flow	
 Distaflex®	Coronary Arteries & SVGs	OTW	2mm	145cm	4F/3F	0.014"	6F > .070"	4F (6F*)				23mL/min	600 sec	300 sec	111304-001 (Ultra)
 XMI®	Coronary Arteries, SVGs, & Peripheral Arterial	OTW	2mm	135cm	4F	0.014"	6F > .068"	4F (6F*)				40mL/min	600 sec	300 sec	105041-001 (Ultra)
 Spiroflex®	Coronary Arteries, SVGs, & Peripheral Arterial	RX	2mm	135cm	4F	0.014"	6F > .070"	5F (6F*)				40mL/min	600 sec	300 sec	106553-001 (Ultra)
 XVG®	Peripheral Arterial	OTW	3mm	140cm	5F	0.014"	7F > .076"	5F (6F*)				60mL/min	600 sec	300 sec	105042-001 (Ultra)
 Spiroflex® VG	Coronary Arteries, SVGs, & Peripheral Arterial	RX	3mm	135cm	5F	0.014"	7F > .076"	6F (7F*)				60mL/min	600 sec	300 sec	106608-001 (Ultra)
 Solent® Proxi	Peripheral Arterial and Venous, AV Access	OTW	3mm	90cm	6F	0.035"	—	6F	Yes	Yes	Yes	60mL/min	480 sec	240 sec	109676-001 (Ultra)
 Solent® Omni	Peripheral Arterial and Venous, AV Access	OTW	3mm	120cm	6F	0.035"	—	6F	Yes	Yes	Yes	60mL/min	480 sec	240 sec	109681-001 (Ultra)
 Solent® Dista	Peripheral Arterial	OTW	1.5mm	145cm	4F/3F	0.014"	—	4F	Yes			23mL/min	600 sec	300 sec	111303-001 (Ultra)
 AVX®	AV Access Grafts & Fistula	OTW	3mm	50cm	6F	0.035"	—	6F			Yes	60mL/min	600 sec	300 sec	105039-001 (Ultra)
	Power Pulse Kit for Ultra 104834-002														
 Aspiration Catheter Fetch® 2	Peripheral Arterial & Coronary Vasculature	RX	2mm	135cm	4F	0.014"	6F > .070"	5F	—	—	—	—			109400-001 (Fetch® 2)

*When a guide catheter is utilized

AngioJet® / Fetch® Catheter Reference Guide

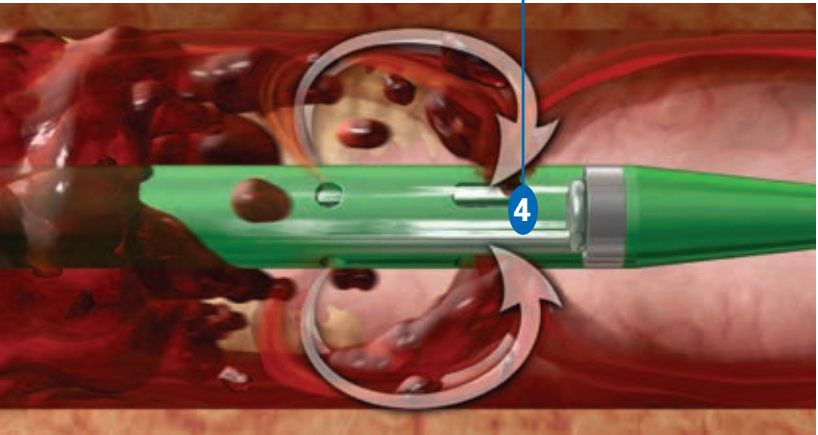


AngioJet® Ultra Thrombectomy System Mechanism of Action



Saline jets travel backwards to create a low pressure zone causing a vacuum effect.

Thrombus is drawn into the in-flow windows and the jets push the thrombus back down the catheter.



AngioJet® Thrombectomy Systems

General Indications/Contraindications - AngioJet System peripheral indications include: breaking up and removing thrombus from infra-inguinal peripheral arteries, upper and lower extremity peripheral arteries, upper extremity peripheral veins, iliofemoral, infra-iliac and lower extremity veins, A-V access conduits, and for use with the AngioJet Power Pulse Kit for the control and selective infusion of physician specified fluids, including thrombolytic agents, into the peripheral vascular system. AngioJet System coronary indications include: removing thrombus in the treatment of patients with symptomatic coronary artery or saphenous vein graft lesions prior to balloon angioplasty or stent placement. Do not use in patients: who are contraindicated for intracoronary or endovascular procedures, who cannot tolerate contrast media, and in whom the lesion cannot be accessed with the wire guide.

General Warnings and Precautions - The System has not been evaluated for treatment of pulmonary embolism in the US and some other countries or for use in the carotid or cerebral vasculature. Some AngioJet devices have not been evaluated for use in coronary vasculature. Operation of the catheter may cause embolization of some thrombus and/or thrombotic particulate debris. Cardiac arrhythmias may occur and cardiac rhythm should be monitored during catheter use and appropriate management employed, if needed. Systemic heparinization is advisable to avoid pericatheterization thrombus and acute rethrombosis. Operation of the System causes transient hemolysis. Large thrombus burdens may result in significant hemoglobinemia which should be monitored. Consider hydration, as appropriate. Before coronary AngioJet treatment, verify the presence of thrombus because routine use of AngioJet in every STEMI patient, without proper selection for thrombus, has been associated with increased mortality risk. Do not use the system in the coronary vasculature without placing a temporary pacing catheter to support the patient through hemodynamically significant arrhythmias which may occur.

Potential Adverse Events - Potential adverse events (in alphabetical order) which may be associated with use of the system are similar to those associated with other interventional procedures and include but are not limited to the following: abrupt closure of treated vessel, acute myocardial infarction, acute renal failure, arrhythmias (including VF and VT), bleeding from access site, death, dissection, embolization (proximal or distal), emergent CABG, hematoma, hemolysis, hemorrhage requiring transfusion, hypotension/hypertension, infection at access site, myocardial ischemia, pain, pancreatitis, perforation, pseudoaneurysm, reactions to contrast medium, stroke/CVA, thrombosis/occlusion, total occlusion of treated vessel, vascular aneurysm, vascular spasm, vessel wall or valve damage.