



Penumbra System®

ELEVATING PERFORMANCE WITH

JET **3D**TM
System

FOR COMBINATION TECHNIQUE

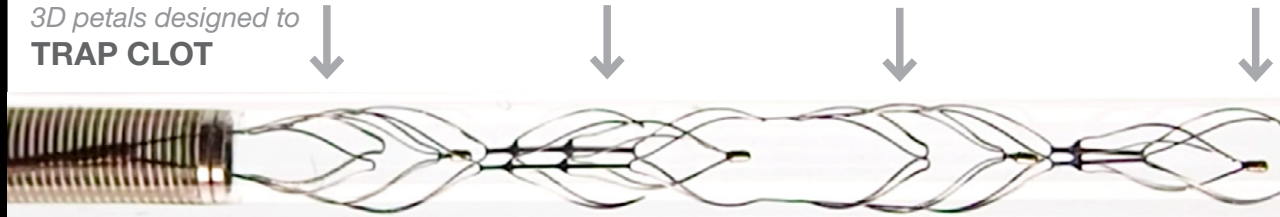
Penumbra 

3D™ LOCKS AND TRAPS CLOT

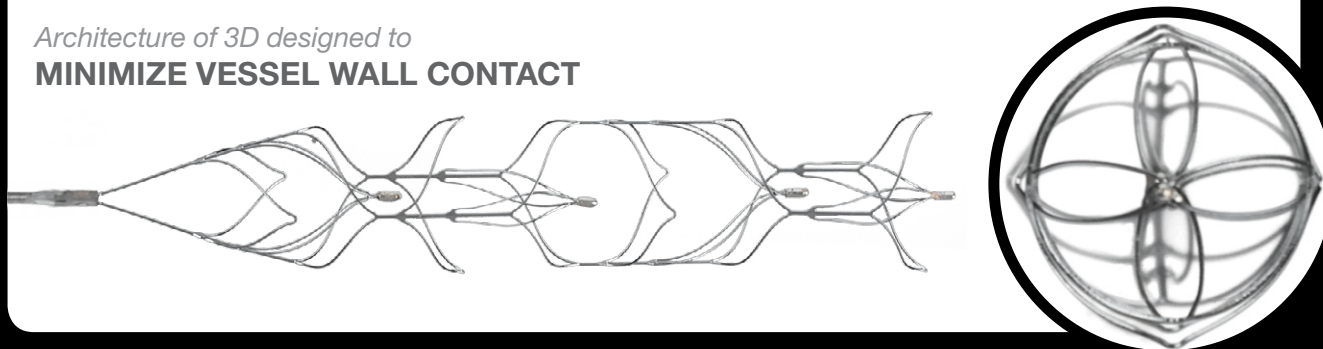
Intraluminal chambers designed to
LOCK CLOT



3D petals designed to
TRAP CLOT



Architecture of 3D designed to
MINIMIZE VESSEL WALL CONTACT



Photographs taken by and on file at Penumbra, Inc.

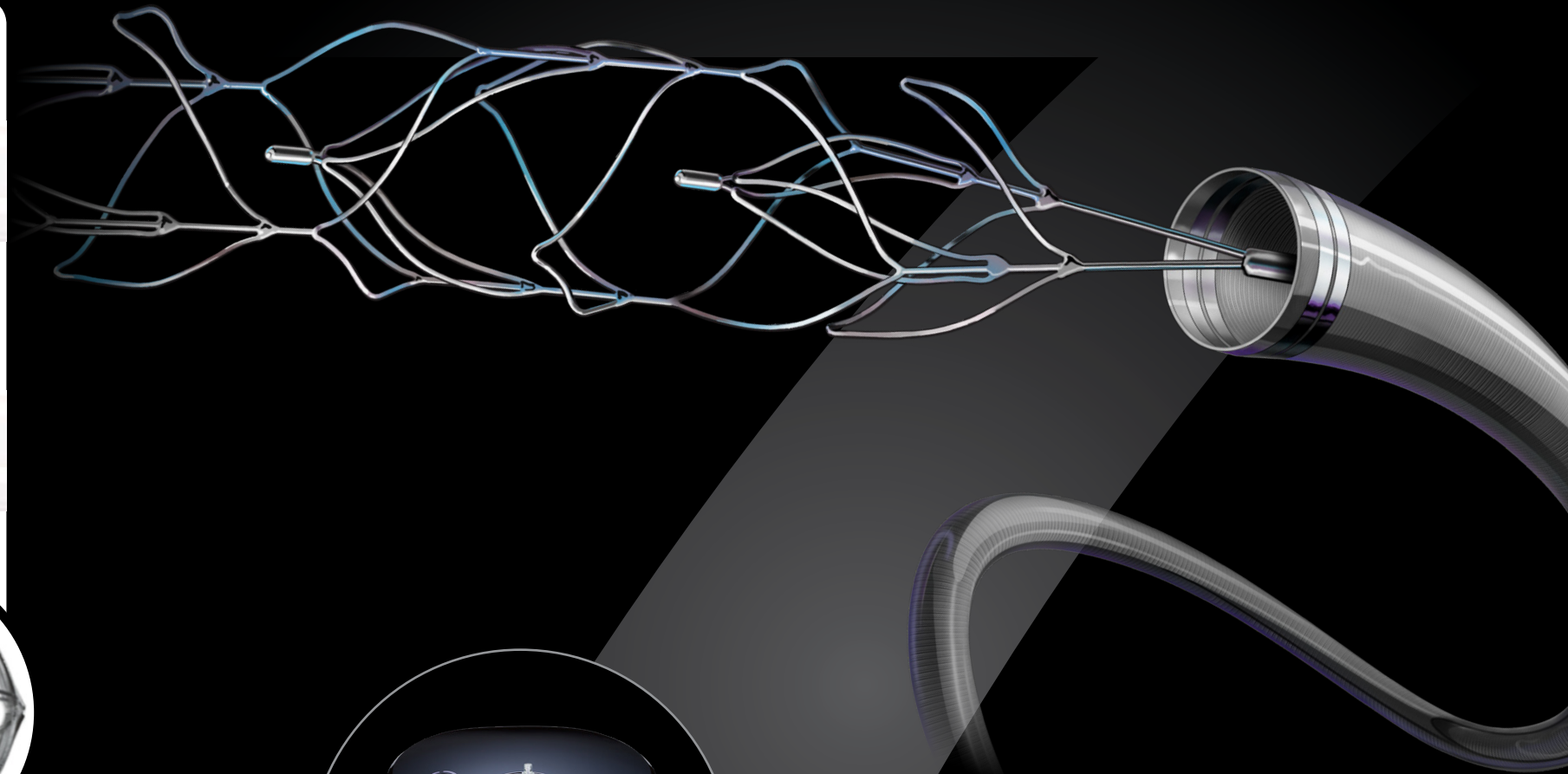
Compatible with
**NEURON™
MAX 088**



JET 3D SYSTEM WHAT'S INSIDE:

Penumbra JET™ 7 Reperfusion Catheter
Penumbra Hi-Flow Aspiration Tubing
3D Revascularization Device™
Velocity® Microcatheter

ELEVATING PERFORMANCE WITH JET AND 3D FOR COMBINATION TECHNIQUE



.072" LUMEN

20 TRANSITIONS
for trackability and navigation

ARTICULATING MARKER BAND
designed to improve tip softness

SUPERIOR FLEXIBILITY
enabled by progressive distal coil wind

ENHANCED PUSHABILITY
featuring Quad-Wire technology

FULL LENGTH PTFE LINER
designed for durability with adjunctive devices

POWERED BY
PENUMBRA ENGINE™

-29.2 inHg^a
Deep Vacuum

a. Data on file at Penumbra, Inc.


ORDERING INFORMATION

Penumbra System®


Catalog Number	Description	Proximal OD (F) (in.)	Distal OD (mm)	Proximal ID (in.)	Distal ID (in.)	Working Length (cm)
Aspiration Kits						
5MAXJET7KIT	JET 7 Reperfusion Catheter + Penumbra Hi-Flow Tubing	6.0 (.085)	2.16	.072	.072	132
5MAXJETDKIT	JET D Reperfusion Catheter + Penumbra Hi-Flow Tubing	6.0 (.080)	1.65	.064	.054	138
5MAXACE068KIT	ACE™68 Reperfusion Catheter + Penumbra Hi-Flow Tubing	6.0 (.080)	2.03	.068	.068	132
5MAXACE132KIT	ACE60 Reperfusion Catheter + Penumbra Hi-Flow Tubing	6.0 (.080)	1.80	.068	.060	132
3MAXCKIT	3MAX™ Reperfusion Catheter + Penumbra Hi-Flow Tubing	4.7 (.062)	1.27	.043	.035	160

Systems		<p>be suspended until 24 hours post-treatment. Medical management and acute post stroke care should follow the ASA guidelines. Any neurological deterioration should be evaluated by urgent CT scan and other evaluations as indicated according to investigator/hospital best practice.</p> <ul style="list-style-type: none">As in all surgical interventions, monitoring of intra-procedural blood loss is recommended so that appropriate management may be instituted.Confirm vessel diameter, and select an appropriate size Penumbra Reperfusion Catheter. Limit the usage of Reperfusion Catheters to arteries larger than the catheter's outer diameter. Refer to the Reperfusion Catheter labeling for dimensional information. <p>Potential Adverse Events</p> <p>Possible complications include, but are not limited to, the following:</p> <p>allergic reaction and anaphylaxis from contrast media; acute occlusion; air embolism; arteriovenous fistula; death; device malfunction; distal embolization; emboli; false aneurysm formation; hematoma or hemorrhage at access site; inability to completely remove thrombus; infection; intracranial hemorrhage; ischemia; kidney damage from contrast media; neurological deficits including stroke; vessel spasm, thrombosis, dissection, or perforation; radiation exposure that may lead to cataracts, skin reddening or burns from x-ray exposure.</p> <p>1. Adams, et al., Guidelines for the Early Management of Adults with Ischemic Stroke: A Guideline from the AHA/ASA Stroke Council, Clinical Cardiology Council, Cardiovascular Radiology and Intervention Council, and the Atherosclerotic Peripheral Vascular Disease and Quality of Care Outcomes in Research Interdisciplinary Working Groups: The American Academy of</p>
JET73DVELSYS	JET 7 Reperfusion Catheter + Penumbra Hi-Flow Tubing + 3D Revascularization Device™ + Velocity® Microcatheter	
ACE603DVELSYS	ACE60 Reperfusion Catheter + Penumbra Hi-Flow Tubing + 3D Revascularization Device + Velocity Microcatheter	
Reperfusion Catheters		
5MAXJETD	JET D Reperfusion Catheter	
5MAXACE068	ACE68 Reperfusion Catheter	
5MAXACE132	ACE60 Reperfusion Catheter	
3MAXC	3MAX Reperfusion Catheter	
Revascularization Device		
PSR3D	3D Revascularization Device	
Delivery Microcatheter		
VEL160STR	Velocity Microcatheter	
Separator™ Devices		
PSF054	5MAX Separator	
3MAXS	3MAX Separator	
Aspiration Accessories		
PMXENGN	Penumbra ENGINE™	
PAPS3	ENGINE Canister	

Neuron™ MAX 6 F 088 Lumen Long Sheath

Catalog Number	Description	Tip Shape	Working Length (cm)
(Crosscut Valve, RHV, and Dilator Included)			
PNML6F088904	6 F 088 Neuron MAX Long Sheath, Straight		90

6 F Select™ Catheters

Catalog Number	Description	Tip Shape	Working Length (cm)
PNS6F105BER	6 F Select Catheter, BER		105
PNS6F125SIM	6 F Select Catheter, SIM		125
PNS6F125SIMV	6 F Select Catheter, SIMV		125
PNS6F125BER	6 F Select Catheter, BER		125

Indication for Use

Penumbra Reperfusion Catheters and Separators

As part of the Penumbra System, the Reperfusion Catheters and Separators are indicated for use in the revascularization of patients with acute ischemic stroke secondary to intracranial large vessel occlusive disease (within the internal carotid, middle cerebral – M1 and M2 segments, basilar, and vertebral arteries) within 8 hours of symptom onset. Patients who are ineligible for intravenous tissue plasminogen activator (IV t-PA) or who fail IV t-PA therapy are candidates for treatment.

Penumbra 3D Revascularization Device

As part of the Penumbra System, the Penumbra 3D Revascularization Device is indicated for use in the revascularization of patients with acute ischemic stroke secondary to intracranial large vessel occlusive disease (within the internal carotid, middle cerebral – M1 and M2 segments) within 8 hours of symptom onset. Patients who are ineligible for intravenous tissue plasminogen activator (IV t-PA) or who fail IV t-PA therapy are candidates for treatment.

Penumbra Aspiration Tubing

As part of the Penumbra System, the Penumbra Sterile Aspiration Tubing is indicated to connect the Penumbra Reperfusion Catheters to the Penumbra Aspiration Pump.

Penumbra Aspiration Pump

The Penumbra Aspiration Pump is indicated as a vacuum source for Penumbra Aspiration Systems.

Contraindications

There are no known contraindications.

Warnings

- The Penumbra System should only be used by physicians who have received appropriate training in interventional neuro-endovascular techniques and treatment of acute ischemic stroke.
- Do not advance, retract or use any component of the Penumbra System against resistance without careful assessment of the cause using fluoroscopy. If the cause cannot be determined, withdraw the device or system as a unit. Unrestrained torquing or forced insertion of the catheter, revascularization device, or separator against resistance may result in damage to the device or vessel.
- Do not use the Penumbra System with a pump other than the

Penumbra Aspiration Pump.

- The Penumbra 3D Revascularization Device has not been evaluated in patients with angiographic evidence of pre-existing arterial injury.
- The use of fluoroscopy may present potential risks from radiation exposure. The probability of adverse events due to radiation exposure increases with the total amount of radiation observed, the number of procedures and total procedure time.

Precautions

- The device is intended for single use only. Do not resterilize or reuse. Resterilization and/or Reuse may result in ineffective catheter coating lubrication, which may result in high friction and the inability to access the target neuro vasculature location.
- Do not use kinked or damaged devices. Do not use open or damaged packages. Return all damaged devices and packaging to the manufacturer/distributor.
- Use prior to the "Use By" date.
- Use the Penumbra System in conjunction with fluoroscopic visualization.
- As in all fluoroscopy procedures, consider all necessary precautions to limit patient radiation exposure by using sufficient shielding, reducing fluoroscopy times and modifying radiation technical factors whenever possible.
- Maintain a constant infusion of appropriate flush solution.
- When performing aspiration, ensure that the Penumbra Aspiration Tubing valve is open for only the minimum time needed to remove thrombus. Excessive aspiration or failure to close the Penumbra Aspiration Tubing valve when aspiration is complete is not recommended.
- The Penumbra Separator is not intended for use as a neurovascular guidewire. If repositioning of the Penumbra Reperfusion Catheter is necessary during the revascularization procedure, such repositioning should be performed over an appropriate neurovascular guidewire using standard microcatheter and guidewire techniques.
- Do not use automated high-pressure contrast injection equipment with the Penumbra Reperfusion Catheter because it may damage the device.
- Administration of anticoagulants and antiplatelets should

Penumbra 

www.penumbrainc.com

Penumbra, Inc. USA

One Penumbra Place
Alameda, CA 94502
USA
1.888.272.4606
T 1.510.748.3200
F 1.510.748.3232
order@penumbrainc.com
info@penumbrainc.com

Caution: Federal (USA) law restricts these devices to sale by or on the order of a physician. Prior to use, please refer to the Instructions for Use for Penumbra System, Penumbra System with 3D Revascularization Device, Penumbra ENGINE, Penumbra Delivery Microcatheters, and Neuron MAX System for complete product indications, contraindications, warnings, precautions, potential adverse events, and detailed instructions for use. Please contact your local Penumbra representative for more information.

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