Amplatzer™ Vascular Plug II

Instructions for Use

Device Description

The Amplatzer™ Vascular Plug II is a self-expandable nitinol mesh occlusion device (see Figure 1).

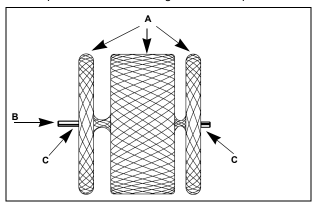


Figure 1. Amplatzer™ Vascular Plug II

- A. Nitinol mesh
- B. Screw attachment
- C. Marker bands

The device has a screw attachment for a delivery wire and radiopaque marker bands at both ends. The Amplatzer™ Vascular Plug II is attached to a 135 cm delivery wire with a stainless steel screw. The Amplatzer™ Vascular Plug II is packaged within a loader, pre-connected to the delivery wire in a hoop dispenser. A plastic vise is included with each device (Figure 2).

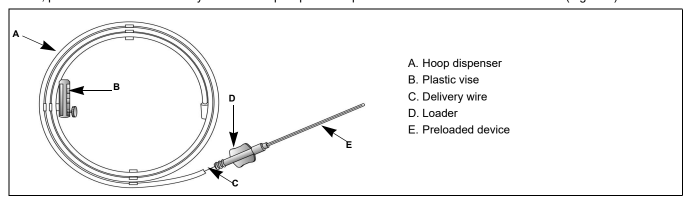


Figure 2. Amplatzer™ Vascular Plug II preloaded device.

Plugs are available in the sizes referenced in Table 1.











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Manufacturing facility: Abbott Medical 5050 Nathan Lane North Plymouth, MN 55442 USA

+1 855 478 5833

+1 651 756 5833

[™] Indicates a trademark of the Abbott group of companies. ‡ Indicates a third party trademark, which is property of its respective owner.

Table 1. Recommended delivery catheter or delivery sheath size

Order Number ^a	Vascular Plug II Diameter Fully Expanded (mm)	Vascular Plug II Length Fully Expanded (mm)	Minimum Delivery Catheter or Delivery Sheath ID ^b (inch)	Maximum Delivery Catheter or Delivery Sheath ID ^b (inch)	Maximum Delivery Catheter or Delivery Sheath Length (cm)
9-AVP2-003	3	6	0.056	0.067	100
9-AVP2-004	4	6	0.056	0.098	100
9-AVP2-006	6	6	0.056	0.098	100
9-AVP2-008	8	7	0.056	0.106	100
9-AVP2-010	10	7	0.070	0.106	100
9-AVP2-012	12	9	0.070	0.106	100
9-AVP2-014	14	10	0.086	0.106	100
9-AVP2-016	16	12	0.086	0.106	100
9-AVP2-018	18	14	0.098	0.106	100
9-AVP2-020	20	16	0.098	0.106	100
9-AVP2-022	22	18	0.098	0.106	100

Intended Use

The Amplatzer™ Vascular Plug II is indicated for arterial and venous embolizations in the peripheral vasculature.

Contraindications

None known.

Warnings

 The safety and effectiveness of this device for cardiac uses (for example, patent ductus arteriosus or paravalvular leak closures) and neurological uses have not been established.

Potential Adverse Events

Potential complications include, but are not limited to: death, migration of the device, stroke, or vessel perforation.

Precautions

- · This device was sterilized with ethylene oxide and is for single use only. Do not reuse or resterilize this device. Attempts to resterilize this device can cause a malfunction, insufficient sterilization, or harm to the patient.
- · Use on or before the last day of the expiration month that is printed on the product packaging label.
- Patients with nickel allergy can experience an allergic reaction to this device.
- Do not use this device if the sterile package is open or damaged.
- This device should be used only by physicians who are trained in standard endovascular techniques. The physician should determine which patients are candidates for procedures that use this device.
- · The physician should exercise clinical judgment in situations that involve the use of anticoagulants or antiplatelet drugs before, during, and/or after the use of this device.
- · Store in a dry place.

a. Some sizes not available in all areas.b. Refer to delivery catheter or delivery sheath manufacturer's product labeling for inner diameter (ID).

MRI Safety Information

MR	A patient with the Amplatzer™ Vascular Plug II may be safely scanned under the following conditions. Failure to follow these conditions may result in injury to the patient.
Device Name	Amplatzer™ Vascular Plug II
Static Magnetic Field Strength (B ₀)	1.5 T or 3.0 T
Maximum Field Spatial Gradient	19T/m (1900 gauss/cm)
RF Excitation	Circularly Polarized (CP)
RF Transmit Coil Type	Body Coil
Operating Mode	Normal Operating Mode
Maximum Whole Body SAR	2.0W/kg (Normal Operating Mode)
Maximum Head SAR	N/A
Scan Duration	2.0W/kg whole-body-averaged SAR for 15 minutes of continuous scanning
MR Image Artifact	The presence of this implant may produce an image artifact.

Directions for Use

- 1. Access the vessel and perform an angiogram using standard technique to measure the diameter of the vessel at the desired occlusion site.
- Select an Amplatzer™ Vascular Plug II with a diameter approximately 30%–50% larger than the vessel diameter at the
 occlusion site. Ensure that the occlusion site has sufficient length to accommodate the deployed device length so that
 the device will not obstruct other vessels or anatomical structures.
- 3. Flush the hoop dispenser and loader with sterile saline until fluid exits the distal tip to purge air from the loader.
- 4. Remove the device (in the loader) and the delivery wire from the hoop dispenser.
- 5. Select a delivery catheter (see Table 1 for delivery catheter size).

NOTE: If the inner diameter of the original access catheter is sufficient for the device size selected, that catheter may be used for delivery.

NOTE: The delivery catheter length should be no more than 100 cm.

- 6. Advance the delivery catheter over the guidewire until the distal tip of the catheter is at the distal edge of the occlusion site.
- 7. Remove the guidewire.
- 8. Insert the loader into the delivery catheter through the Y-connector or hemostasis valve.
- 9. A single Y-Connector kit is recommended for use with the Amplatzer™ Vascular Plug II. B. Braun Medical Inc., Order No. 610400.
- 10. Allow blood backflow or aspirate the system to ensure air is purged from the catheter and loader.

CAUTION: Do not overtighten Y-connector screw to avoid damaging loader.

- 11. Push on the delivery wire to advance the device into the delivery catheter. Remove the loader from the wire if desired.
- 12. Advance the delivery wire and device to the distal end of delivery catheter. Do not twist or rotate the delivery wire during advancement to ensure device does not prematurely detach.
- 13. Hold the delivery wire in place and slowly withdraw the delivery catheter to deploy the device at the occlusion site.
- 14. Verify position of the device.
- 15. If device position is unsatisfactory:
 - Stabilize the wire and re-advance the delivery catheter until the device is completely within the catheter.
 - Reposition and deploy, or remove the device from the patient.
- 16. If the device position is satisfactory:
 - Attach the plastic vise to the wire, and release the device, by rotating the delivery wire counterclockwise until it separates from the device.
 - Remove the delivery catheter and wire from the patient.

Post-procedure Instructions

Temporary patient ID card – Go to www.amplatzer.com/tempIDcard to print the temporary patient identification card.
 Complete this card and give it to the patient.

Disposal

• The carton and Instructions for Use are recyclable. Dispose of all packaging materials as appropriate.

- Devices can be returned to Abbott Medical for disposal. Contact an Abbott Medical representative or returns@amplatzer.com for instructions.
- · Use solid biohazard waste procedures to discard devices.

Warranty

Abbott Medical warrants to buyer that, for a period equal to the validated shelf life of the product, this product shall meet the product specifications established by the manufacturer when used in accordance with the manufacturer's instructions for use and shall be free from defects in materials and workmanship. Abbott Medical's obligation under this warranty is limited to replacing or repairing at its option, at its factory, this product if returned within the warranty period to Abbott Medical and after confirmed to be defective by the manufacturer.

EXCEPT AS EXPRESSLY PROVIDED IN THIS WARRANTY, ABBOTT MEDICAL DISCLAIMS ANY REPRESENTATION OR WARRANTY OF ANY KIND, EXPRESS OR IMPLIED, INCLUDING ANY WARRANTY AS TO MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

See the Terms and Conditions of Sale for further information.

State of California (USA) Only:

WARNING: This product can expose you to chemicals including ethylene oxide, which is known to the State of California to cause cancer and birth defects or other reproductive harm. For more information, go to www.P65Warnings.ca.gov.

Symbol Definitions

The following symbols may appear on the device packaging:

Symbol	Definition		
\triangle	Caution, consult accompanying documents		
•••	Manufacturer		
PHT	Contains Phthalate		
REF	Reference number		
SN	Product serial number		
LOT	Batch Code		
	Use-by date		
2	Do not re-use		
UDI	Unique device identification		
STERILE EO	Sterilized using ethylene oxide		
[]i	Consult instructions for use		
medical.abbott/manuals	Follow instructions for use on this website		

Symbol	Definition		
Ť	Keep dry; keep away from rain		
	Do not use if package is damaged		
Not made with natural rubber latex	Not made with natural rubber latex		
MR	MR Conditional		
	Inner diameter		
Q	Outer diameter		
\longleftrightarrow	Length		
←	Usable length		
<u></u>	Recommended delivery sheath/catheter dimensions		
Ronly	Federal law restricts this device to sale by or on the order of a physician.		
	Quantity		
~~~	Date of manufacture		
Vascular Plug	Vascular Plug		