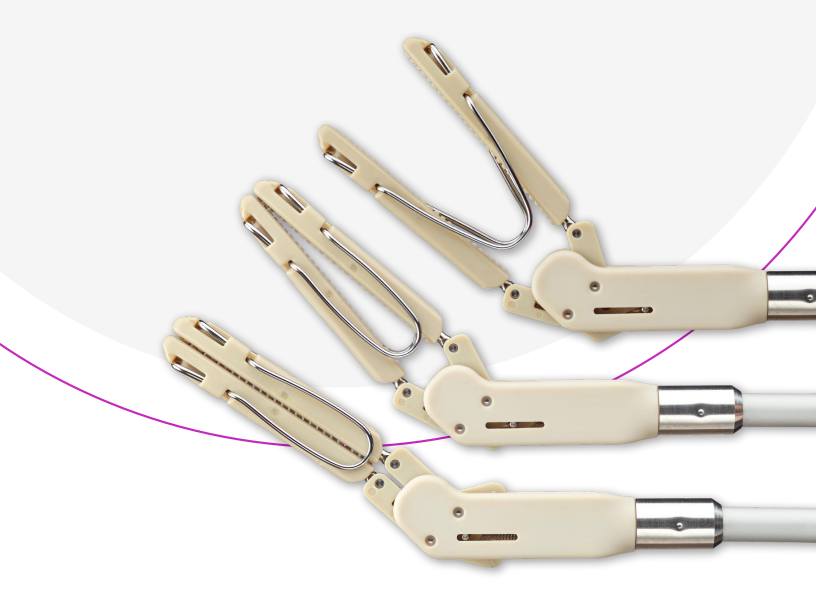
Medtronic

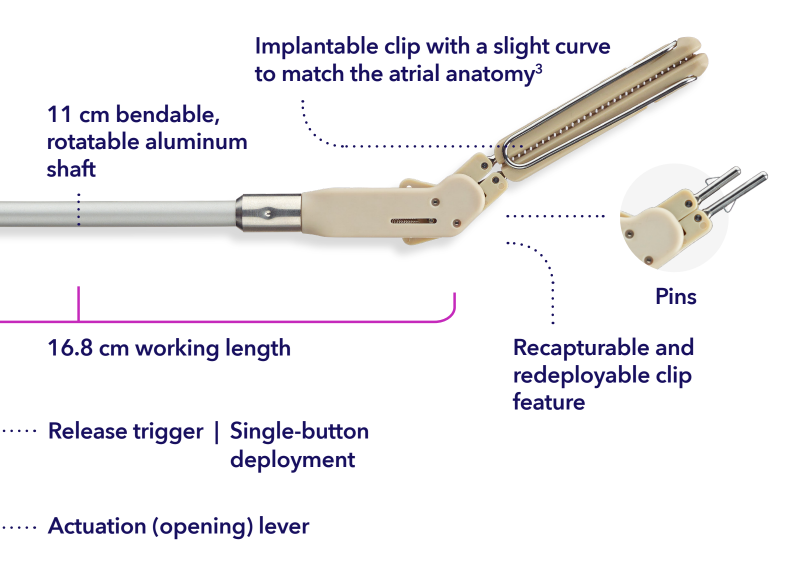
Penditure[™] LAA Exclusion System



The Medtronic Penditure left atrial appendage (LAA) clip

- Fabric-free design for atraumatic closure and reduced risk of inflammation¹
- Tip-first closure helps provide complete and secure exclusion of the appendage²
- Curved clip to better match the atrial anatomy³
- Recapturable, redeployable, and repositionable after deployment to ensure precise placement²
- Wide opening at the base offers good visibility, which may assist in proper clip placement⁴







Handle

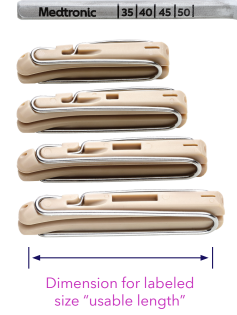
Possible complications related to the use of Penditure[™] LAA exclusion system in combination with open heart surgery are: bleeding, tissue damage, thromboembolism, and pericardial effusion. For a complete listing of all indications, contraindications, precautions, and warnings, please refer to the Instructions for Use, which accompany each product.

Ordering information | Penditure LAA exclusion system

| CFN | Length (A) | Width (B) | Height (C) | Opening (D) [†] |
|-----------------|----------------------------|-----------|------------|-----------------------------|
| LAAC35 35 mm | 39.6 mm | 6.9 mm | 11 mm | 19.0 mm |
| LAAC40 40 mm | 44.6 mm | 7.4 mm | 11 mm | 19.9 mm |
| LAAC45 45 mm | 50.0 mm | 8.1 mm | 11 mm | 23.2 mm |
| LAAC50 50 mm | 55.0 mm | 8.8 mm | 11 mm | 20.0 mm |
| LAACSIZER | Packaged separately (3/pk) | | | |

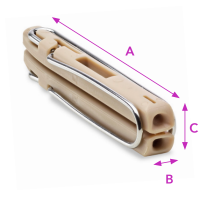
†Implant tip opening is a function of the moment arm and force of the spring. The location where spring force acts is further from the delivery system with longer clips. The result is that tip openings vary, and may not be consistent, with clip size.

Penditure $^{\text{m}}$ LAA Exclusion System is 510(k) cleared, and only available for sale in the United States.



Approximate size shown

Penditure clips are textured PEEK material with a subtle curve (radius). Nitinol springs provide the constant closing force.

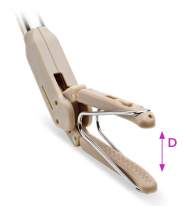


35 mm

40 mm

45 mm

50 mm



¹Medtronic data on file: GLP Animal Study Report TR-00023. These tests may not be indicative of clinical performance.

²Medtronic data on file: Product Specification Matrix PRS-00001. These tests may not be indicative of clinical performance.

³Medtronic data on file: Product Specification Matrix PRS-00001 and Curvature report EGS-00006. These tests may not be indicative of clinical performance.

⁴Medtronic Data on File: VOC August 29, 2023, may not be indicative of clinical performance

LifeLine

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medtronic.com

Important Safety Information

Indications for use

The Penditure LAA Exclusion System is indicated for the exclusion of the left atrial appendage of the heart, performed under direct visualization and in conjunction with other cardiac surgical procedures. Direct visualization, in this context, requires that the physician can see the heart directly, with or without assistance from a camera, endoscope, and so forth, or any other appropriate viewing technologies.

Only physicians who are trained in standard cardiac surgical procedures can use this device.

Contraindications

- Do not use this device if the patient has a known allergy to nitinol (nickel titanium alloy).
- Do not use this device as a contraceptive tubal occlusion device

Potential Adverse Effects

Possible complications related to the use of Penditure™ LAA exclusion system in combination with open heart surgery are: bleeding, tissue damage, thromboembolism, and pericardial effusion. For a complete listing of all indications, contraindications, precautions, and warnings, please refer to the Instructions for Use, which accompany each product.

For more information, contact your local Medtronic cardiac surgery representative. U.S. Customer Service: 1-800-328-1357

Caution: Federal law (USA) restricts this device to sale by or on the order of a physician. For a listing of indications, contraindications, precautions, and warnings, please refer to the Instructions for Use.

