

Celect Platinum®

VENA CAVA FILTER

Clinically successful design

The four radiopaque platinum markers can verify placement accuracy.³ Anchors and platinum markers on the filter's feet allow secure caval fixation.⁴

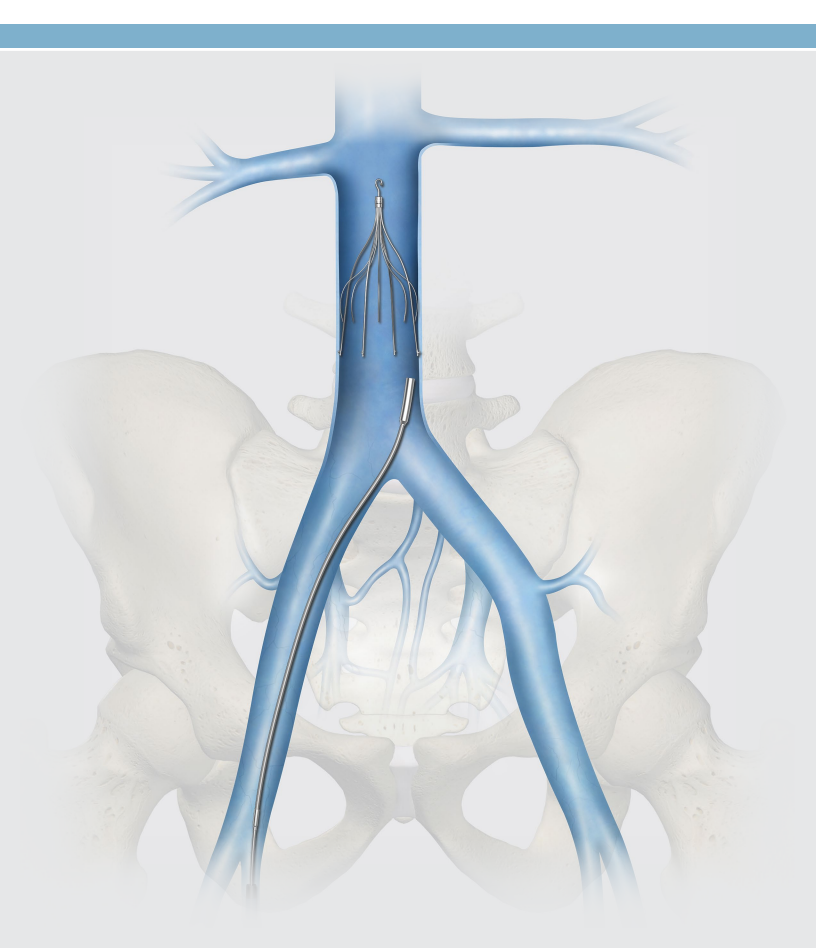
Accurate, controlled deployment

The NavAlign[™] delivery system helps to deploy the filter precisely, easily providing control with accurate deployment.⁵

Constructed of unique material

Cook filters are constructed of a durable Conichrome® alloy.





The **Cook Celect Platinum** filter implant is intended for the prevention of recurrent pulmonary embolism (PE) via placement in the vena cava in the following situations:

- Pulmonary thromboembolism when anticoagulant therapy is contraindicated
- Failure of anticoagulant therapy in thromboembolic diseases
- Emergency treatment following massive PE where anticipated benefits of conventional therapy are reduced
- Chronic, recurrent PE where anticoagulant therapy has failed or is contraindicated

The Cook Celect Platinum filter implant may be retrieved if clinically indicated; please refer to the Optional Filter Retrieval section of the Instructions for Use for more information. The product is intended for percutaneous placement via a femoral vein for filtration of inferior vena cava (IVC) blood to prevent PE.

Order Number	Reference Part Number	Sheath Diameter Fr	Max. indicated Cava Diameter for Filter Placement mm	Filter Length mm	Sheath Working Length cm		
Celect Platinum	t Platinum® Vena Cava Filter Set for Femoral Vein Approach						
G34502	IGTCFS-65-1-FEM-CELECT-PT	7	30	49	65		
Celect Platinum	Vena Cava Filter Set for Jugular Veir	Approach		er for Filter Sheath nent Length Working Length mm cm			
G34309	IGTCFS-65-1-JUG-CELECT-PT	7	30	49	65		
Celect Platinum ^e	Vena Cava Filter Set for Femoral and	Sheath Diameter Filter Placement mm Sheath Working Length mm cm for Femoral Vein Approach -CELECT-PT 7 30 49 65 for Jugular Vein Approach -CELECT-PT 7 30 49 65 for Femoral and Jugular Vein Approach					
G34505	IGTCFS-65-1-UNI-CELECT-PT	7	30	49	65		

Some products or part numbers may not be available in all markets. Contact your local Cook representative or Customer Support & Distribution for details.

- 1. Lyon SM, Riojas GE, Uberoi R, et al. Short- and long-term retrievability of the Celect vena cava filter: results from a multi-institutional registry. *J Vasc Interv Radiol*. 2009;20(11):1441–1448.
- 2. Prospective study of the Cook Celect Filter, including permanent and retrievable use (Study Protocol #05-507) final study report; September 2009.
- 3. Demonstrated in simulated use tests: D00302649 and D00006045.
- 4. Demonstrated in testing: D00006050 (TR-3789) and D00006018 (TR-3787).
- 5. Celect Platinum® Vena Cava Filter: View the Instructions for Use for a thorough examination of the procedural instructions, intended use, contraindications, warnings and precautions, and potential adverse events.

Cook Celect® Platinum Vena Cava Filter Set for Femoral Vein Approach

CAUTION: U.S. federal law restricts this device to sale by or on the order of a physician (or a properly licensed practitioner).

INTENDED USE: The Cook Celect Platinum Filter implant is intended for the prevention of recurrent pulmonary embolsm (PE) via placement in the vena cava in the following situations: Pulmonary thromboembolism when anticoagulant therapy is contraindicated: Failure of anticoagulant therapy in thromboembolis diseases; Emergency treatment following massive PE where anticipated benefits of conventional therapy are reduced; and • Chronic, recurrent PE where anticoagulant therapy has failed or is contraindicated. The Cook Celect Platinum Filter implant may be retrieved if clinically indicated; please refer to the "Optional Filter Retrieval" section in the complete Instructions for Use for more information. The product is intended for percutaneous placement via a femoral vein for filtration of inferior vena cava (VCV) blood to prevent PE.

CONTRAINDICATIONS: Filter Placement • Megacava (diameter of the IVC > 30 mm). • Diameter of the IVC < 15 mm.
• Extensive thrombus in the vein chosen for approach. • Patients with risk of septic embolism. • Use in pregnant womer
• Use in minors/pediatric patients. Optional Filter Retrieval - Filters with significant amounts of trapped thrombus
(greater than 25% of the volume of the cone). • Patients with an ongoing high risk for PE.

WARNINGS: Filter Placement - If severe resistance is met when advancing the wire guide or the introducer system, then critact and choose a different approach. Excessive force should not be exerted: -When power injecting contrast media, do not exceed the maximum pressure rating of 68 bar/1000 psi and flow rate of 20 mL/sec. Hand injection is also possible. -When inserting the preloaded filter into the Check-Flo* valve of the introducer sheath, hold the introducer system. Do not re-sheath the expanded filter. -Do not attempt to rotate the preloaded filter inside the introducer system. Do not re-sheath the expanded filter during femoral approach. -Do not attempt to rotate, advance or retract the expanded filter inside the vena cava. -Excessive force should not be exerted in placement of the fleter ident of the filter inside the vena cava. -Excessive force should not be exerted in placement of the fleter inside the rotation of the filter inside the introducer system on the special context of the filter inside the value of the filter inside value of the filter ins

PRECAUTIONS: The product is intended for use by physicians trained and experienced in diagnostic and interventional endovascular techniques. • Standard techniques for placement of vascular access sheaths, angiographic catheters and wire guides should be employed. • The Cook Celect Platinum Filter Set should be used in patients with vessel diameters compatible with the associated device components. • Product (filter or introducer system) modification or alteration is not recommended, as the product's safety and effectiveness has not been established following any modifications. • Manipulation of products (e.g., placement and retrieval) requires imaging control. • Before injecting any contrast media (by either power or hand injection) through the introducer dilator, ensure that the introducer sheath hub antiroducer dilator are correctly connected. • Possible allergic reactions (e.g., to cobalt, chromium, nickel and platinum) should be considered. • Ensure that the patient is not with the anesthetic procedure. • Ensure that he patient is not allergic/sensitive to contrast media, since the use of contrast media during the procedure and/or during postoperative imaging may cause an allergic reaction and/or other contrast-induced harms. • Placement in the suprarenal position has been reported. The safety and effectiveness of the filter has not been established in these patients. • Filter tilt has been reported. Potential causes may include filter placement in IVCs with diameters smaller or larger than those specified in the complete Instructions for Use; improper deployment; manipulations near an implanted filter (e.g., a surgical or endovascular procedure in the vicinity of a filter); and in a filter filter (e.g., a surgical or endovascular procedure in the vicinity of a filter); and penetration/perforation has been reported. Potential causes may include improper deployment; and only be either symptomatic or asymptomatic. Potential causes may include improper deployment; and of or excessive force or manipulations nea

be due to repetitive motion on a filter leg in an unusual, stressed position, such as a filter leg penetrating/perforating the IVC; or a filter leg being caught in a side branch (e.g., a renal vein). Other potential causes of filter fracture may include excessive force or manipulations near an implanted filter (e.g., a surgical or endovascular procedure in the vicinity of a filter). Retrieval of a fractured filter or filter fragment implanted filter (e.g., a surgical or endovascular procedure in the vicinity of a filter). Retrieval of a fractured filter or filter fragment migration and (or) embolization (e.g., movement to the heart or lungs) has been reported. Filter of filter fragment movement has occurred in both the cranial and caudal direction and may be either symptomatic or asymptomatic. Potential causes may include filter placement in IVCs with diameters smaller or larger than those specified in the complete Instructions for Use; improper deployment; deployment into thrombus; disbdogment due to large thrombus burdens; and (or) excessive force or manipulations near an in situ filter (e.g., a surgical or endovascular procedure in the vicinity of a filter). Increased friction and/or compression at the access site during the procedure may lead to increased risk of thrombosis at the access site. Follow the instructions throughly to ensure successful deployment, and to avoid any harm to the patient or damage to the device. If the introduction system or parts of the introduction system malfunctions during procedure, the device should be replaced. If the device malfunctions during procedure, perform careful replacement to avoid injuries to the access site and vessel. Failure to store the device correctly may result in material degradation and/or damage to the device and the filter or the access site and vessel. Failure to store the device correctly may result in susually preferred due to its straighter route to the vena cava. The left femoral vein can be used, but is more tortuous. Prior to choosing an approach,

POTENTIAL ADVERSE EVENTS: Potential adverse events that may occur include, but are not limited to, the following:

- Access site thrombosis/occlusion - Air embolism - Arrhythmia - Back or abdominal pain - Blood loss - Branch vessel occlusion - Cardiac damage - Cardiac tamge - Verange to the vene cava - Death - Deep vein thrombosis - Edema - Extravasation of contrast material - Failure of filter expansion/incomplete expansion - Filter or filter fragment embolization - Filter fracture - Filter migration - Filter malpositioning - Hemortmage - Hematoma at vascular access site - Infection at vascular access site - Intimal tear - Obstruction of blood flow - Pneumothorax - Postphlebitic syndrome - Pulmonary embolism - Retrieval failure - Trauma to adjacent structures - Unacceptable filter tilt - Vascular trauma - Vena cava perforation - Vena cava settoriss - Vena cava settoriss - Vena cava stenosis

See Instructions for Use for full product information.

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Customer Service

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