GÜNTHER TULIP™ VENA CAVA FILTER
For Jugular Vein Approach

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

DEVICE DESCRIPTION
The Günther Tulip Vena Cava Filter Set consists of a non-magnetic filter (30 mm diameter, 50 mm long), preloaded in a protection sheath on a jugular filter introducer system with a stainless steel grasping hook, a 7.0 French coaxial introducer system (is compatible with a .035 inch wire guide), and a hydrophilic coated dilator. The introducer dilator has 8 sideports and two radiopaque markers 30 mm apart (end to end).

Set Components

Fig. 1

- **DILATOR**
  - Radiopaque hydrophilic coated
  - 10.0 French 20 cm long

- **THREE-WAY STOPCOCK**
  - Plastic

- **Protection Sheath**

- **Preloaded filter introducer**

- **COAXIAL INTRODUCTER SHEATH SYSTEM**
  - Radiopaque introducer sheath, 7.0 French 65 cm long with radiopaque band and radiopaque introducer dilator with 2 radiopaque markers at the distal end

- **Check-Flo Valve**

- **Hook**
  - Secondary Legs
  - Anchors

- **Günther Tulip Vena Cava Filter**
  - supplied preloaded

- **Primary Legs**
INTENDED USE

The Günther Tulip Vena Cava Filter Set 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 pulmonary embolism where anticipated benefits of conventional therapy are reduced; and
• Chronic, recurrent pulmonary embolism where anticoagulant therapy has failed or is contraindicated.

The Günther Tulip Vena Cava Filter may be retrieved according to the instructions supplied in the section labeled “Optional Retrieval Procedure”.

The product is intended for percutaneous placement via a jugular vein for filtration of inferior vena cava (IVC) blood to prevent pulmonary embolism.

The product is intended for use by physicians trained and experienced in diagnostic and interventional techniques.

Standard techniques for placement of vascular access sheaths, angiographic catheters and wire guides should be employed.

CONTRAINDICATIONS

Filter Placement

• Megacava (diameter of the IVC > 30 mm).
• Vena cava filters should not be implanted in patients with risk of septic embolism.

Optional Filter Retrieval

• Retrieval of the filter with significant amounts of trapped thrombus (greater than 25% of the volume of the cone).
• Retrieval of the filter for patients with an on-going high risk for pulmonary embolism.

WARNINGS

Filter Placement

• This Günther Tulip Vena Cava Filter Set is intended for jugular vein approach only and cannot be used for femoral vein approach.
• Manipulation of products requires fluoroscopic control.
• When injecting contrast medium, do not exceed maximum pressure rating of 1000 psi and flow rate of 20 ml/sec. Hand injection is also possible.
• Excessive force should not be used to place filter.

Optional Filter Retrieval

• Excessive force should not be used to retrieve the filter.
• An inferior vena cavagram evaluation for residual captured thrombus should be performed prior to attempted retrieval.
• Available data from retrievals in multicenter and single center studies demonstrate that the device can be safely retrieved. Please refer to the “Clinical Studies” section of this booklet for clinical study references to the retrieval of this filter.
PRECAUTIONS
Possible allergic reactions should be considered.

Filter Placement
• For placement of the filter, the right jugular vein is preferable. An approach via the left jugular vein is possible.
• The filter may be repositioned prior to final deployment by carefully advancing the introducer sheath over the filter only to the anchors. Reposition the system as desired, and again withdraw the introducer sheath by reattaching it to the protection sheath hub, completely exposing the filter.

Optional Filter Retrieval
• For filter retrieval, a right jugular vein approach is preferable. An approach via the left jugular vein is possible; however, there are no available data which demonstrate the safety or effectiveness of filter retrieval via the left jugular vein.
• The filter has been designed to be retrieved with the Günther Tulip Vena Cava Filter Retrieval Set, GTRS- (not included). Cook has not performed testing to evaluate the safety or effectiveness of filter retrieval using other retrieval systems.
• Never re-deploy a retrieved filter.

MR COMPATIBILITY
Non-clinical testing has demonstrated that the Günther Tulip Vena Cava Filter is MR Conditional. It can be scanned safely immediately after placement under the following conditions:
• Static magnetic field of 3.0 Tesla or less
• Spatial gradient field of 525 Gauss/cm or less
• Maximum whole body averaged specific absorption rate (SAR) of 3.0 W/kg for 15 minutes of scanning.

In non-clinical testing, the Günther Tulip Vena Cava Filter produced a temperature rise of less than 0.6°C at a maximum whole body averaged specific absorption rate (SAR) of 3.0 W/kg for 15 minutes of MR scanning in a 3.0 Tesla MR system using a transmit/receive body coil (Excite, Software G3.0-052B, General Electric Healthcare, Milwaukee, WI) MR scanner.

MR image quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the Günther Tulip Vena Cava Filter. Therefore, it may be necessary to optimize MR imaging parameters for the presence of this metallic implant.

POTENTIAL ADVERSE EVENTS
• Damage to the vena cava
• Pulmonary embolism
• Filter embolization
• Vena cava perforation
• Vena cava occlusion or thrombosis
• Hemorrhage
• Extravasation of contrast material at time of vena cavagram
• Hematoma at vascular access site
• Infection at vascular access site
• Thrombosis or stenosis at implant site
• Death
**CLINICAL STUDIES**

To evaluate the safety of retrieving the Günther Tulip Vena Filter, a clinical study was conducted in which 41 patients (female (n=19); male (n=22)) were enrolled for possible retrieval of the filter. The results of this and other published and presented sources listed below demonstrate that the Günther Tulip Vena Cava Filter may be safely retrieved:

<table>
<thead>
<tr>
<th>Reference</th>
<th>Filters Inserted</th>
<th>Retrieval Attempts</th>
<th>Successful Retrievals</th>
<th>Range (Days)</th>
<th>Mean (Days)</th>
<th>Adverse Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kachura JR “Inferior Vena Cava Filter Removal After 475-day Implantation”. JVIR 2005; 16: 1156-1158.</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>475</td>
<td>475</td>
<td>None</td>
</tr>
<tr>
<td>Binkert CA, Bansal A, Gates JD, “Inferior Vena Cava Filter Removal After 317 day Implantation.” JVIR 2005; 16:1395-1398.</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>317</td>
<td>317</td>
<td>Mild caval stenosis following 317-day retrieval; follow-up OK</td>
</tr>
<tr>
<td>Piano G, et al. “Safety, Feasibility, and Outcome of retrievable Vena Cava Filters in High-risk Surgical Patients.” J Vasc Surg 2007; 45:784-788</td>
<td>60</td>
<td>54</td>
<td>52</td>
<td>32-162</td>
<td>63</td>
<td>One patient had non-fatal PE with filter in place. Three patients required second retrieval attempt. All three were successful.</td>
</tr>
<tr>
<td>Morris CS, Rogers FB, Najarian KE, Bhave AD, Shachford SR. “Current Trends In Vena Cava Filtra-tion with the Introduction of a retrievable Filter at a Level 1 Trauma Center.” J Trauma, 2004; 57(1):32-36.</td>
<td>58</td>
<td>14</td>
<td>13</td>
<td>11-41</td>
<td>19</td>
<td>One patient had a non-fatal PE after filter retrieval.</td>
</tr>
</tbody>
</table>

1 The unsuccessful retrievals did not result in adverse events; the device was left in place in place as a permanent implant.
INSTRUCTIONS FOR USE

Preparation
1. Flush the introducer sheath and dilator.

2. Advance the introducer dilator through the middle of the silicone valve on the introducer sheath. Secure the introducer dilator to the introducer sheath by twisting the dilator hub clockwise (Fig. 2).

Filter Placement
3. Puncture the jugular vein using the Seldinger technique.
4. Over the wire guide, dilate the puncture site with the dilator. Remove the dilator.
5. Advance the coaxial introducer sheath system over the wire guide.
6. Remove the wire guide.
7. Using either power or hand injection, perform cavography to verify position below (caudal to) the renal veins.
   **WARNING:** Do not exceed maximum pressure rating of 1000 psi and flow rate of 20 ml/sec.

8. Remove the introducer dilator by twisting the dilator hub counter clockwise (Fig. 3).

9. Place the filter introducer with the protection sheath containing the preloaded filter into the hub of the introducer sheath. Advance the filter introducer with the protection sheath into the sheath (Fig. 4).
10. Advance the introducer with the protection sheath to the introducer sheath hub and connect the sheath hub and protection sheath by twisting clockwise. The filter is now at the radiopaque band of the introducer sheath. The hook of the filter should be below the renal veins (Fig. 5).

11. Stabilize the introducer, and withdraw the introducer sheath and protection sheath until the protection sheath and handle are connected. At this point the filter is expanded, still connected to the filter introducer (Fig. 6).

12. If the filter is not in the desired position, carefully advance the introducer sheath over the filter only to the anchors. Reposition the system as desired, and again withdraw the introducer sheath by reattaching it to the protection sheath hub, completely exposing the filter.

**WARNING:** Do not advance the sheath over the anchors to avoid scratching particles off the sheath.

13. When the filter position is correct, push the red safety button to prepare filter release (Fig. 7).

14. While keeping slight back tension on the introducer push the release button completely to ensure proper release of the filter. Repositioning of the filter is no longer possible. The filter is now released (Fig. 8).
15. Remove the introducer and perform cavography to verify filter position and then withdraw the introducer sheath.

**Optional Retrieval Procedure**

**NOTE:** If a filter retrieval is going to be performed, please refer to Instructions for Use provided with the Günther Tulip Vena Cava Filter Retrieval Set (not included) for device description and caution statement.

1. Hold the clear Y-fitting and pull back the plastic pin vise on the wire loop retriever to cover the loop. Tighten the screw of the clear Y-fitting to keep the loop inside the catheter (Fig. 9).
2. Puncture the right jugular vein using the Seldinger technique.
3. Position a flush catheter inferior to the filter and perform a diagnostic vena cavagram.
4. Exchange the flush catheter for the coaxial retrieval sheath system, advancing it over the wire guide.

5. Remove the red inner dilator and the wire guide. Verify the position by injection of contrast medium (Fig. 10).

6. Introduce the retrieval loop system through the coaxial retrieval sheath system, advance and connect the white Tuohy-Borst side-arm adapter of the loop system to the sheath system. The Tuohy-Borst adapter can be tightened around the catheter to prevent loss of blood (Fig. 11).

7. Loosen the screw of the clear Y-fitting to enable advancement of the loop inside the catheter. Hold the clear Y-fitting and push the pin vise forward. Advance until the loop has fully expanded inside the vena cava and surrounds the filter (Fig. 12).

8. Pull back the loop until it engages the hook of the filter (Fig. 13).

**CAUTION:** Do not pull on the filter beyond what is required to keep tension on the loop. Doing so may cause damage to the caval wall.
9. Hold the wire loop steady with the pin vise, then push the clear Y-fitting with the catheter forward until it touches the hook. To snare the filter in this position make sure to firmly lock the screw of the clear Y-fitting on the wire loop (Fig. 14).

**NOTE:** If the retrieval wire loop is losing its shape during the attempt to engage the hook of the filter, it can be removed and gently reshaped. After reshaping, clean loop and proceed from step 6.

10. While holding steady the retrieval loop system with the clear Y-fitting, advance the white Tuohy-Borst side-arm adapter and the black inner sheath with the coaxial retrieval system. The filter collapses and the hooks disengage from the caval wall (Fig. 15).

**CAUTION:** Advance the inner sheath over the filter to collapse it. Do not retract the loop snare. This may cause damage to the caval wall.

11. When the tip of the coaxial retrieval system is at the anchors, loosen the hub of the outer sheath, advance the outer sheath forward to cover the whole filter, and retrieve the complete assembly (Fig. 16).

**CAUTION:** If the outer sheath is not advanced over the inner sheath to cover the anchors, the anchors may scratch or damage the caval wall.

**POST-RETRIEVAL CARE**

After retrieval of filter, hospital standard of care should be followed for removing the sheath and providing hemostasis to prevent bleeding at the vascular access site.

**HOW SUPPLIED**

Supplied sterilized by ethylene oxide gas in peel-open packages. Intended for one-time use. Sterile if package is unopened or undamaged. Do not use the product if there is doubt as to whether the product is sterile. Store in a dark, dry, cool place. Avoid extended exposure to light. Upon removal from package, inspect the product to ensure no damage has occurred.

**REFERENCES**

These instructions for use are based on experience from physicians and (or) their published literature. Refer to your local Cook sales representative for information on available literature.