Echo PS™
Positioning System
with Ventralight™ ST Mesh or Composix™ L/P Mesh

Laparoscopic ventral hernia repair

Echo PS™ Positioning System
with Ventralight™ ST Mesh

Echo PS™ Positioning System
with Composix™ L/P Mesh

KEY BENEFITS
• Easy Insertion
• Effortless Placement and Positioning
• Assisted Fixation
• Compatible with Robotic Surgical Systems

SOFT TISSUE REPAIR
Right Procedure. Right Product. Right Outcome.
Compatible with Robotic Surgical Systems

The ECHO PS™ Positioning System provides traditional laparoscopic advantages to robotic surgical systems.
For an efficient, reproducible repair

The Echo PS™ Positioning System is a low profile balloon that comes preattached to either VENTRALIGHT™ ST Mesh or COMPOSIX™ L/P Mesh. When inflated, the Echo PS™ Positioning System facilitates the deployment (including unrolling, positioning and placement) of the mesh during laparoscopic ventral hernia repair. Upon completion of initial perimeter fixation, the balloon is deflated quickly and completely removed from the body.

KEY BENEFITS

Easy Insertion
- The prepackaged Introducer Tool holds the mesh in place, ensuring a tight, uniform roll
- The positioning system and the mesh are low profile, facilitating trocar deployment

Effortless Placement and Positioning
- Mesh easily unrolls and opens during inflation
- No additional trocar site is needed to hold the mesh in place
- Positioning system design and orientation markers allow for ease of orientation (anterior vs. posterior side, long vs. short axis, and the midline of the mesh)
- System designed to facilitate centering of the mesh over the defect
- System designed to eliminate the time and effort involved with placing and pulling up of positioning sutures

Assisted Fixation
- Positioning system keeps the mesh open and up against the abdominal wall with no additional graspers or spreading devices, allowing for complete visibility during fixation
- Following initial perimeter mesh fixation with the OptiFix™ Absorbable Fixation System, the positioning system is deflated and quickly and completely removed from the body
The **Echo PS™** Positioning System comes preattached to either **Ventralight™** ST Mesh or **Composix™** L/P Mesh, requiring no assembly or specialty instruments.

**Anterior Side:** Uncoated monofilament polypropylene mesh

**Posterior Side:** Absorbable hydrogel barrier based on Sepra® Technology (**Ventralight™** ST Mesh only)

- Low profile, thermoplastic polyurethane (TPU) coated nylon balloon
- Logo identifies long axis
- Removal points marked by arrows
- Tabs clearly identify the connector locations
- Marked center of proximal ends represent the midline of the mesh
- Connectors keep the **Echo PS™** Positioning System attached to the mesh, and are simultaneously removed from the abdominal cavity with the **Echo PS™** Positioning System
- Inflation tube
- Anchor allows for a secure connection between the inflation tube and inflation assembly
- Tube cut location
- Retrieval loop
Basic Steps for a Reproducible Repair

1. **Roll**
2. **Insert**
3. **Inflate**
4. **Position**
5. **Initially Fixate**
6. **Deflate**
7. **Remove Echo PS™ Positioning System**
8. **Complete Fixation**

Also included:
- Introducer Tool facilitates rolling and insertion of mesh
- Prepackaged inflation assembly inflates balloon
**Echo PS™ Positioning System with VENTRALIGHT™ ST Mesh or Composix™ L/P Mesh**

**VENTRALIGHT™ ST Mesh with Echo PS™ Positioning System**

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<th>Mesh Size</th>
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**Also Available:**

**Composix™ L/P Mesh with Echo PS™ Positioning System**

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<td>Rectangle</td>
<td>10.2” x 14.2” (26.1 cm x 36.2 cm)</td>
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**Order Form**

- Please add these marked products to my preference card.
- I would like to have these marked products in stock.
- I would like to trial these marked products.

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<tr>
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**ECHO PS™ Positioning System with VENTRALIGHT™ ST Mesh or Composix™ L/P Mesh**

**Indications**

VENTRALIGHT™ ST Mesh is indicated for use in the reconstruction of soft tissue deficiencies, such as the repair of hernias. **Composix™ L/P Mesh** is indicated for use in the reconstruction of soft tissue deficiencies, such as for the repair of hernias and chest wall defects. The **Echo PS™ Positioning System** is intended to be used to facilitate the delivery of soft tissue prostheses through laparoscopic hernia repair.

**Contraindications**

Do not use **VENTRALIGHT™ ST Mesh** in infants or children whereby future growth will be compromised by use of such material.

Do not use **VENTRALIGHT™ ST Mesh** for the reconstruction of cardiovascular defects.

**Warnings**

**VENTRALIGHT™ ST Mesh/Composix™ L/P Mesh** is the only permanent implant component of the device. The inflation adapter and syringe are to be kept external to the patient and discarded after use. The **Echo PS™ Positioning System** (including the balloon, all connectors, and inflation tube) is to be removed from the patient and appropriately discarded as it is not part of the permanent implant.

**Adverse Reactions**

Possible complications include seroma, adhesions, inflammation, extrusion, fistula formation, infection, allergic reaction, and recurrence of the hernia or soft tissue defect.

**OptiFix™ Absorbable Fixation System**

**Indications**

The **OptiFix™ Absorbable Fixation System** is indicated for the approximation of soft tissue and fixation of surgical mesh to tissues during open or laparoscopic surgical procedures, such as hernia repair.

**Contraindications**

Contraindications associated with laparoscopic and open surgical procedures relative to mesh fixation apply, including but not limited to:

- Fixation of vascular or neural structures
- Fixation of bone and cartilage
- Situations with insufficient ingrowth of tissue into the mesh over time, which could result in inadequate fixation once the fastener is absorbed

Carefully inspect the area in the vicinity of the tissue being fastened to avoid inadvertent penetration of underlying structures such as nerves, vessels, viscera or bone. Use of the **OptiFix™ Absorbable Fixation System** in the close vicinity of such underlying structures is contraindicated. For reference, the length of the fastener below the fastener head is 6.1 mm, the fastener head is another 0.6 mm (total 6.7 mm).

**Warnings**

The device may not fixate through prosthetics derived from biologic material such as xenografts and allografts. Prosthetic should be evaluated for compatibility prior to use.

After use, the **OptiFix™ Absorbable Fixation System** may be a potential biohazard. Handle and dispose of in accordance with any local and federal laws regarding medical waste.

**Adverse Reactions**

Adverse reactions and potential complications associated with fixation devices such as the **OptiFix™ Absorbable Fixation System** may include, but are not limited to the following:

- Hemorrhage; pain, edema and erythema at wound site; allergic reaction to Poly(D, L)-lactide; infection/septicemia; hernia recurrence/wound dehiscence.

To learn more, contact your local **Bard** Representative or call 1.800.556.6275.

Please consult package insert for more detailed safety information and instructions for use.

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