Obtryx™ II
Transobturator Mid-urethral Sling System
Obtryx™ II
Transvaginal Mid-urethral Sling System

How can we continue to innovate our family of mid-urethral slings that have already been the products of choice for nearly 800,000 patients? By creating the same reliable mesh in an easy-to-see, optical blue color. So whatever your preferred surgical approach, Advantage™ Blue mesh provides improved visibility so you can treat your patients with confidence.

Improved visibility. Evidence based.
- The same mesh properties as our patented Advantage mesh, which is documented in more than 35 publications to date
- The easy-to-see, optical blue color helps to improve your visibility for more accurate intra-operative sling tensioning and makes it easier to locate post-operatively

Trusted polypropylene mesh¹
- Mesh thickness: 0.66 mm
- Pore size: 1182 μm
- Fiber size (diameter): 0.15 mm
- Weight: 100 g/m²

Transobturator approach

Needle
- Designed to facilitate transobturator device passage
- Two needle configurations (curved and halo) allow physicians to choose their preferred approach (curved and halo sold separately)

Association loop
- Facilitates needle and mesh engagement and removal

Blue dilator leg
- Designed to create a smooth transition from the needle to the mesh assembly
- Improves intra-operative visibility in the event of vaginal perforation
- Minimizes the force needed to deliver the mesh assembly through the patient’s anatomy

Mesh assembly
- No sleeve coverage under the suburethral segment to allow for mesh visibility and to aid in precise placement
- Reduced plastic sleeve minimizes the force needed to remove mesh assembly

Blue centering tab
- Allows for proper alignment of the center of the mesh under the urethra. It also allows the physician to apply counter tension to the sling while preserving the mesh integrity.
Obtryx™ II Transobturator Mid-urethral Sling System Procedure

1. Prepare the skin lateral to the inferior pubic ramus and vaginal operative sites.
2a. Make a 1.0 cm to 1.5 cm vertical midline incision on the anterior vaginal wall at the level of the mid-urethra. Dissect bilaterally to the interior portion of the inferior pubic ramus at the 45 degree angle off the midline creating a vertical pathway for delivery device placement.
2b. Create a vertical skin incision large enough to insert tip of needle just lateral to the edge of the inferior pubic ramus at the junction where the inferior pubic ramus and adductor longus muscle meet. Repeat on the contralateral side.
3. Grasp the device handle and insert one needle through one skin incision, piercing through the obturator muscle and obturator membrane.
4. Turn the handle at a 45 degree angle medial towards the midline. Place the opposite hand’s forefinger into the lateral dissection of the vaginal incision, placing the fingertip on the distal end of the needle. Guide the distal end of the needle around the inferior pubic ramus through the vaginal incision, maintaining contact with the finger.
5. Appropriately tension the mesh/sleeve assembly according to physician preference.
6. Once proper tension is achieved, cut the leader loop that is on the outside of the sleeve that is connecting the dilator leg and sleeve to the mesh. Pull outward on the dilator to remove the sleeve leaving the mesh in place. Repeat on the other side.
7. Grasp the blue centering tab and cut the exposed white suture located to the distal end of the needle.

Ordering Information

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<th>Product code</th>
<th>Description</th>
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<td>Obtryx™ II Transobturator Sling System – Curved</td>
<td>2 Delivery Devices and 1 Mesh Assembly</td>
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<td>M0068505110</td>
<td>Obtryx™ II Transobturator Sling System – Halo</td>
<td>2 Delivery Devices and 1 Mesh Assembly</td>
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Caution: For Female Mid-Urethral Slings: Federal (US) law restricts this device to sale by or on the order of a physician trained in use of surgical mesh for repair of stress urinary incontinence.

CAUTION: The law restricts these devices to sale by or on the order of a physician. Indications, contraindications, warnings and instructions for use can be found in the product labeling supplied with each device. Information for use only in countries with applicable health authority registrations. Material not intended for use in France. Products shown for INFORMATION purposes only and may not be approved or for sale in certain countries. Please check availability with your local sales representative or customer service for availability in other markets.

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