

EsophyX® Z+ Device



Helical Retractor

- Engages and retracts tissue
- Anchors gastroesophageal junction during fundoplication
- Stows/locks safely inside tissue mold during insertion and removal

Tissue Mold and Chassis

- Plicates and compresses tissue
- Rotates fundus around esophagus to create partial wrap

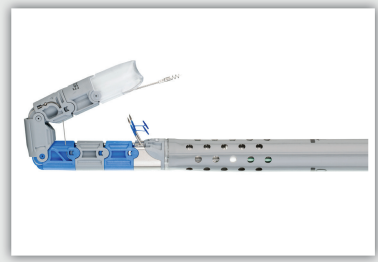
Invaginator

- Suctions esophageal tissue circumferentially
- Reduces small hiatal hernia
- Facilitates proper position of fundoplication caudal to diaphragm

Stylets and SerosaFuse® Fasteners*

- Provides durable tissue apposition
- Fasteners maintain tissue compression throughout healing process

- Increased compatibility with a wide range of high definition endoscopes (8.6 to 11.4mm diameter).
- Simultaneous dual fastener deployment using new trigger handle design
- Tissue mold designed to shroud stylets during fastener delivery
- Compatible with 7.5 mm SerosaFuse® Implantable Fastener cartridges



*"On average, 21 ± 4 fasteners were used to secure the newly created valves."

Trad et al: TEMPO Trial at 5 years: Surgical Innovation Feb 2018



EsophyX Device	Z+ Model
Catalog No	R2007
Endoscope Compatibility	Minimum size OD > 8.6 mm Maximum size OD < 11.4 mm Ensure endoscope compatibility with endo- scope sizing tool R4007
Sterility	Sterile, single-use device
Shelf Life	2 year shelf life
SerosaFuse® Fasteners	7.5 mm
Catalog No.	R2175
Fastener web length	7.5 mm
Fasteners Per Cartridge	20
Material	Non-resorbable polypropylene
Sterility	Sterile, single-use device
Shelf Life	2 year shelf life

SerosaFuse® Implantable Fastener Kit	
Catalog No.	R2275
	Includes (2) 7.5mm cartridges Ships with (1) EsophyX Z+ device
Sterility	Sterile, single-use device
Shelf Life	2-year shelf life
Z+ Scope Compatibility Tool	
Catalog No.	R4007
	Ensure endoscope compatibility with endoscope sizing tool





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INDICATIONS

The EsophyX device with SerosaFuse® fasteners and accessories are indicated for use in transoral tissue approximation, full thickness plication and ligation in the gastrointestinal tract. They are indicated for the treatment of symptomatic chronic GERD in patients who require and respond to pharmacological therapy. The device is also indicated to narrow the gastroesophageal junction, and reduce hiatal hernia < 2cm in size in patients with symptomatic chronic GERD. Patients with hiatal hernias larger than 2cm may be included, when a laparoscopic hiatal hernia repair reduces the hernia to 2cm or less.