Capio™ SLIM

Suture Capturing Device

Scientific Scientific

Building on the history, reducing the profile

Always there.

Dedicated to Women's Health

Capio™ SLIM Suture Capturing Device

The Capio SLIM Suture Capturing device features a reduced profile to minimize the space required within the surgical field, easy suture loading and a funnel shaped catch, designed for consistent ease of use.



Suture Capturing Device

Since 1995, the Capio Device has provided consistent suture placement in difficult to access pelvic floor locations.

70% Reduction in Shaft Diameter

Easy Suture

Dart Loading

New "funnel" suture dart catch design

36% Reduction in Head Width



Capio SLIM
Suture Capturing Device



Capio OPEN ACCESS
Suture Capturing Device

The evolution continues...



1995

The Capio Suture Capturing Device debuts, revolutionizing the field of urogynecology.



1999

The Capio Device is used with Repliform™ Graft, offering another treatment option.



2006

Ergonomic Handle

Providing

3 Finger Grip

The Capio Device is used with Xenform™ Graft, offering more material options for prolapse repair.



Comparison of Device Heads



Figure 1: Comparison of Device Heads illustrates the difference in carrier diameter amongst these fixation devices. The Capio SLIM Suture Capturing Device carrier diameter is 45% less than the FIXT Suturing Device and 43% less than the Digitex Suture Delivery System.

Building on the history and reducing the profile, the Capio SLIM Suture Capturing Device features one of the smallest device profiles and suture dart carrier diameter while keeping the same bite depth as the current Capio Open Access Suturing Device.

Fixation Device Comparison

	Shaft Diameter (mm)	Head Width (mm)	Device Weight (g)	Carrier Diameter (mm)
Capio OPEN ACCESS Suture Capturing Device**	10.1	9.8	45.4	1.2
Capio SLIM Suture Capturing Device*	3.0	6.3	38.2	1.2
Digitex Suture Delivery System*	9.5	9.6	133.3	2.1
FiXT Suturing Device*	4.8	6.9	121.1	2.2

Figure 2: Fixation Device Comparison illustrates the device profile differences amongst these fixation devices. The Capio SLIM Suture Capturing Device has the smallest device profile based on the following measurements: shaft diameter, head width and device weight.

with the Capio SLIM Suture Capturing Device.



2008

The Capio Device is used with the Uphold™ System, one of the smallest mesh footprints for pelvic floor reconstruction.



2011

The Capio Device remains "the foundation" of Pelvic Floor Reconstruction.



2012

Capio Slim: the evolution continues.

^{*}Measurements were recorded using one (1) of each device. Data on file at Boston Scientific.

^{**}Head width and device weight values are an average of five (5) device measurement recordings. Shaft diameter values are an average of fifteen (15) device measurement recordings. Data on file at Boston Scientific.

Capio™ SLIM Suture Capturing Device

Order Number	Description	Unit
M006 831825 0	Capio SLIM Suture Capturing Device	bx 1
M006 831826 1	Capio SLIM Suture Capturing Device	bx 5

Capio Sutures

Order Number	Description	Size	Unit
M006 833113 1	Non-absorbable, coated braided polyester, double armed with TC tapercut needle (dart), 48"	0	bx 12
M006 833123 1	Non-absorbable, polypropylene monofilament, double armed with TC tapercut needle (dart), 48"	0	bx 12
M006 833213 1	Absorbable, coated braided PGA, double armed with TC tapercut needle (dart), 48"	0	bx 12
M006 833114 1	Non-absorbable, coated braided polyester, double armed with TC tapercut needle (dart) and a T 26mm 1/2 circle taper needle, 36"	0	bx 12
M006 833124 1	Non-absorbable, polypropylene monofilament, double armed with TC tapercut needle (dart) and a T 26mm 1/2 circle taper needle, 36"	0	bx 12
M006 833137 1	Monodek™ Absorbable, monofilament PDO, double armed with TC tapercut needle (dart) and a T 26mm 1/2 circle taper needle, 48"	0	bx 12

Your Pelvic Floor Reconstructive Tool to Meet a Variety of Your Algorithm Needs:

- Native Tissue Repair
- Biologic Graft Augmentation
- Synthetic Mesh And More...

Repliform Tissue Regeneration Matrix complies with U.S. Regulations in 21 CFR part 1270 and 1271 Human Tissue for Transplantation.

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Caution: Federal Law (USA) restricts these devices to sale by or on the order of a physician. Refer to package insert provided with these products for complete Indications for Use, Contraindications, Warnings, Precautions, Adverse Events, and Instructions prior to using these products.

Data on file. Bench test results may not necessarily be indicative of clinical performance. Results from case studies are not predictive of results in other cases. Results in other cases may vary.



Boston Scientific Corporation One Boston Scientific Place Natick, MA 01760-1537 www.bostonscientific.com/gynecology

Ordering Information 1.888.272.1001

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