

INTERLOCK[™]-35 COIL Fibered IDC[™] Occlusion System

THE EVOLUTION BEGINS ONE ADVANCEMENT AFTER ANOTHER



Accuracy, Precision and Power

The Interlock[™]-35 Coil introduced the unique combination of powerful Dacron[™] Fibers and precise detachability. Now we redefine deliverability with advancements that are designed to improve the consistency of coil deployment. Technology advances again.

Improved Deliverability

Side holes on coil introducer allow for hydration during advancement and retraction

Nitinol pusher wire engineered for improved kink resistance

Flushing luer intended to facilitate pre-deployment hydration



Technology advances again.



Precision

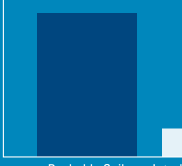
Interlocking arms for precise placement

- The Interlock -35 Coil features a simple interlocking connection between the pusher wire and the coil.
- While inside the lumen of the catheter, the interlocking connection remains attached.
- When pushed outside of the catheter, the connection detaches.

Power

Dacron Fibers for rapid occlusion

Number of Coils Used Per Procedure



Pushable Coils (n=27)

Interlock Fibered IDC Occlusion System (n=23)

- Dudek, et al, Embolization of the Gastroduodenal Artery Before Selective Internal Radiotherapy: A Prospectively Randomized Trial Comparing Standard Pushable Coils with Fibered Interlock Detachable Coils. CVIR, April 14, 2010.
- 83.6% of cases (19 of 23) achieved complete occlusion with one Interlock Fibered IDC Occlusion Coil.
- A shorter procedure time was observed for procedures utilizing detachable coils when compared to pushable coils.
- No coil migration was observed in the detachable coil group, whereas one incidence of migration was observed in the pushable coil group.

INTERLOCK[™] –35 COIL Fibered IDC[™] Occlusion System



Standard Length 2D Configurations							
UPN	Description	Diameter (mm)	Length (cm)	Shape			
M001363500	Interlock - 35 Coil	3	4	2D			
M001363520	Interlock - 35 Coil	4	10	2D			
M001363540	Interlock - 35 Coil	6	10	2D			
M001363550	Interlock - 35 Coil	6	20	2D			
M001363570	Interlock - 35 Coil	8	10	2D			
M001363580	Interlock - 35 Coil	8	20	2D			
M001363590	Interlock - 35 Coil	8	40	2D			
M001363600	Interlock - 35 Coil	10	20	2D			
M001363610	Interlock - 35 Coil	10	40	2D			
M001363620	Interlock - 35 Coil	12	20	2D			
M001363630	Interlock - 35 Coil	12	40	2D			
M001363640	Interlock - 35 Coil	15	20	2D			
M001363650	Interlock - 35 Coil	15	40	2D			
M001363660	Interlock - 35 Coil	18	20	2D			
M001363670	Interlock - 35 Coil	18	40	2D			



Cube Configurations								
UPN	Description	Diameter (mm)	Length (cm)	Shape				
M001363700	Interlock - 35 Coil	4	6	Cube				
M001363720	Interlock - 35 Coil	6	10	Cube				
M001363730	Interlock - 35 Coil	6	20	Cube				
M001363760	Interlock - 35 Coil	8	20	Cube				
M001363790	Interlock - 35 Coil	10	25	Cube				
M001363800	Interlock - 35 Coil	10	40	Cube				
M001363810	Interlock - 35 Coil	15	25	Cube				
M001363820	Interlock - 35 Coil	15	40	Cube				
M001363830	Interlock - 35 Coil	20	40	Cube				



Diamona Configurations							
Description	Diameter (mm)	Length (cm)	Shape				
Interlock - 35 Coil	4	4.5	Diamond				
Interlock - 35 Coil	6	9	Diamond				
Interlock - 35 Coil	8	14	Diamond				
	Description Interlock - 35 Coil Interlock - 35 Coil	DescriptionDiameter (mm)Interlock - 35 Coil4Interlock - 35 Coil6	DescriptionDiameter (mm)Length (cm)Interlock - 35 Coil44.5Interlock - 35 Coil69				

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete "Directions for Use" for

more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator's Instructions.

INTENDED USE/INDICATIONS FOR USE: The Interlock - 35 Fibered IDC Occlusion System is indicated for obstructing or reducing blood flow in the peripheral vasculature during embolization procedures. This device is not intended for neurovascular use. CONTRAINDICATIONS: None known. PRECAUTIONS: Do not attempt to use the Interlock - 35 Fibered IDC Occlusion System with a soft-walled delivery catheter. Do not advance the Interlock - 35 Fibered IDC Occlusion System if it becomes lodged within the catheter. Determine the cause of the resistance and replace the catheter and coil if necessary. ADVERSE EVENTS: The complications that may result from a peripheral embolization procedure include, but are not limited to: • Complications related to catheterization (e.g., hematoma at the site of entry, clot formation at the tip of the catheter and subsequent dislodgement, nerve and vessel dissection or perforation, etc.) • Pain • Hemorrhage • Infection necessitating medical intervention • Foreign body reactions necessitating medical intervention • Emboli • Ischemia • Vasospasm • Tissue necrosis • Undesirable clot formation of the vasculature • Recanalization • Death • Temporary neurological deficit

Interlock and Fibered IDC are unregistered trademarks of Boston Scientific Corporation or its affiliates. All other trademarks are property of their respective owners



Peripheral Interventions

300 Boston Scientific Way Marlborough, MA 01752-1234 www.bostonscientific.com

To order product or for more information contact customer service at 1.888.272.1001.

© 2015 Boston Scientific Corporation or its affiliates. All rights reserved.

PI-181206-AC JUN2015