

Orchestra®

Hydrophilic Nitinol Guidewires

Nitinol Shaft and Core

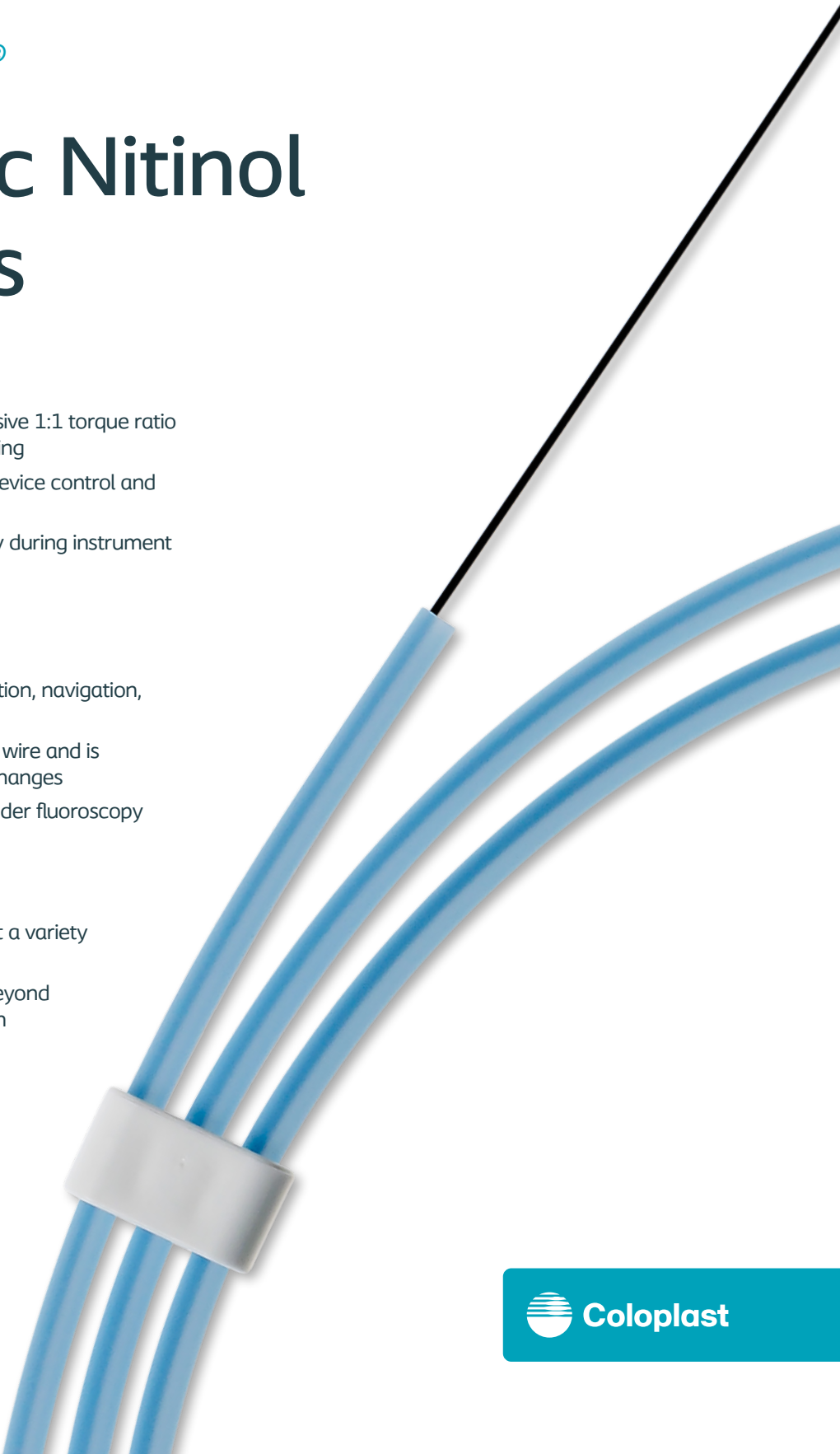
- Nitinol core design provides ultra-responsive 1:1 torque ratio for enhanced maneuverability and handling
- Ideal blend of stiffness and flexibility for device control and atraumatic navigation
- Elastic core provides support and stability during instrument advancement and exchange

Hydrophilic Outer Coating

- Reduces surface friction for ease of insertion, navigation, and withdrawal
- Coating exists on the entire length of the wire and is maintained after multiple instrument exchanges
- Outer jacket is radiopaque for visibility under fluoroscopy

Tip Talk

- Variety of available sizes and tips to meet a variety of procedural needs
- 3 cm flexible tip facilitates the passage beyond obstructions and minimizes trauma within urinary tract



Ordering Information

Coloplast Interventional Urology Service Support 800-258-3476

Diameter	Length	Shaft	Tip Shape	Quantity	Order Number
.035"	150 cm	Standard	Angled	Box of 5	AEAE35
.035"	150 cm	Stiff	Straight	Box of 5	AECD35

Orchestra Hydrophilic Guidewire Brief Statement

Indications

The hydrophilic guidewire is intended to facilitate the placement of devices through the urinary tract during endourologic procedures.

Contraindications

The hydrophilic guidewire is not intended for use other than for endourologic procedures.

Warnings

The hydrophilic guidewire should be used only by a physician, who is well trained in manipulation and observation of guidewires.

The following are risks with the use of the device: perforation of the ureter, damage to the urinary tract, iatrogenic burns.

The information provided is not comprehensive with regard to product risks. For a comprehensive listing of indications, contraindications, warnings, precautions, and adverse events refer to the product’s Instructions for Use. Alternatively, you may contact a Coloplast representative at 1-800-258-3476 and/or visit the company website at www.coloplast.com.

Caution: Federal law (USA) restricts this device to sale by or on the order of a physician.

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