

ReTrace[®]

Ureteral Access Sheath

*Safe, secure access and
procedural efficiency*

Safe, Secure Access

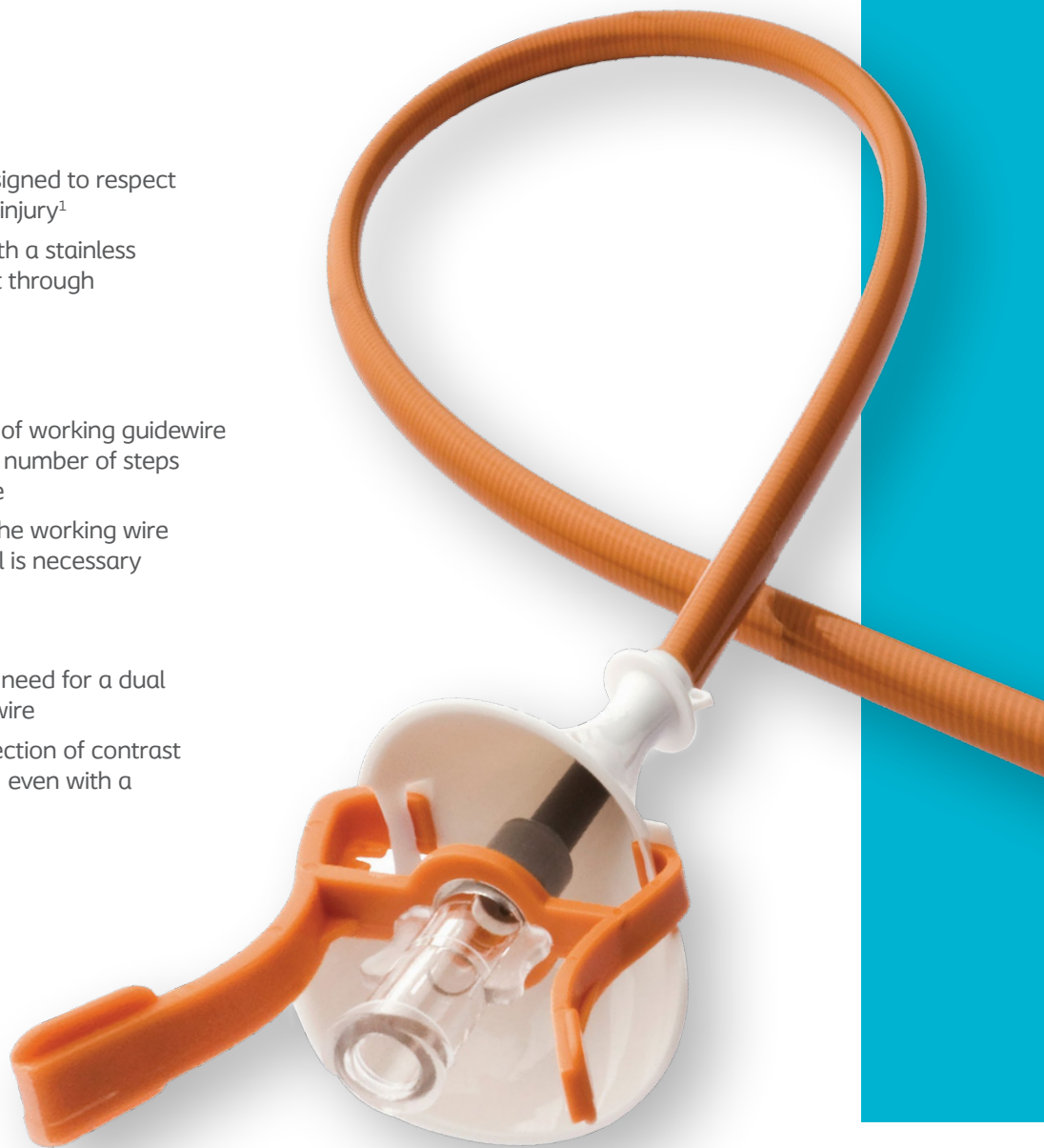
- Flexible introducer tip carefully designed to respect the anatomy and prevent ureteral injury¹
- Kink resistant sheath reinforced with a stainless steel coil to facilitate advancement through tortuous anatomy

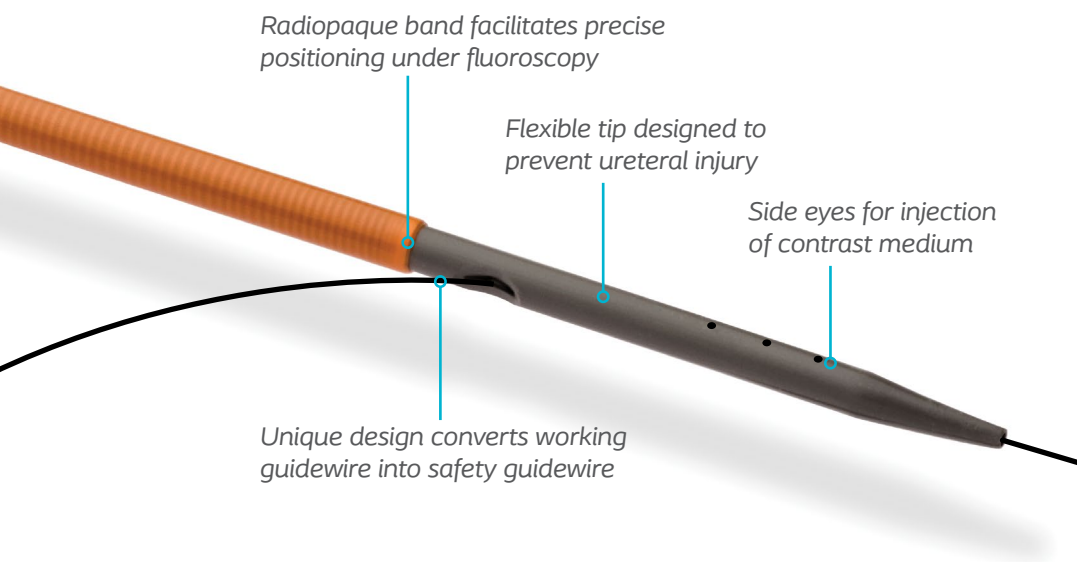
Improves Procedure Time

- Unique design enables conversion of working guidewire into safety guidewire, reducing the number of steps within the ureteroscopic procedure
- Safety guidewire may be used as the working wire to reintroduce sheath if withdrawal is necessary

Reduces Procedure Cost

- Single wire concept eliminates the need for a dual lumen catheter and second guidewire
- Side eyes on introducer enable injection of contrast medium for retrograde pyelogram, even with a guidewire in place





Ordering Information

Coloplast Interventional Urology Surgical Support **800-258-3476**

Description	Order Number
10-12 Fr, 28 cm length	ASXL10
10-12 Fr, 35 cm length	ACXL10
10-12 Fr, 45 cm length	AXXL10
10-12 Fr, 55 cm length	ALXL10
12-14 Fr, 28 cm length	ASXL12
12-14 Fr, 35 cm length	ACXL12
12-14 Fr, 45 cm length	AXXL12
12-14 Fr, 55 cm length	ALXL12

1 Doizi S, Knoll T, Scaffone CM, Breda A, Brehmer M, Liatsikos E, Cornu JN and Traxer O. (2014). First clinical evaluation of a new innovative ureteral access sheath (ReTrace®): a European study. *World J Urol*, 32(1), 143-7.

Indications for Use: To establish a continuous conduit during urological endoscopic procedures facilitating the in and out passage of endoscopes and other instruments into the urinary tract.

Warnings: Reuse of this single use product may create a potential risk to the user. Reprocessing, cleaning, disinfection and sterilization may compromise product characteristics which in turn create an additional risk of physical harm to or infection of the patient.

See Instructions for Use for detailed information regarding warnings/precautions, adverse events prior to using this product. For further information contact Coloplast Corp at 1-800-258-3476 and/or consult the company website at www.coloplast.us.

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

Ostomy Care
 Continence Care
 Wound & Skin Care
 Interventional Urology

Coloplast Corp. Minneapolis, MN 55411 / Urology Care Surgical Support 1-800-258-3476

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