



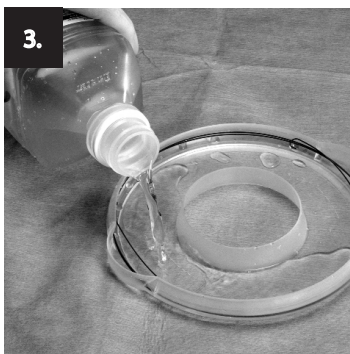
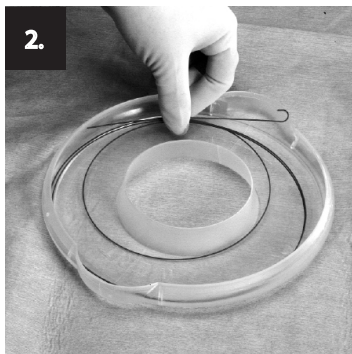
GUIDE WIRE BASIN

EN INSTRUCTIONS FOR USE

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1. Employ an aseptic technique during removal from the package during use.



INDICATION FOR USE

"The RingMaster™ Guide Wire Basin is a containment device that holds guide wires, catheters, and other associated medical devices such as balloons and stents."

CLINICAL BENEFITS

- Hydration - Keeps wires hydrated during procedure.
- Versatile - Holds catheters and wires.
- Identification - Separates wires for easy selection.

CONTRAINDICATIONS

None

PRECAUTIONS

None

In the EU, any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the applicable member state.

POTENTIAL COMPLICATIONS

All efforts have been made to reduce the risks from the use of this device as much as possible; however, the potential exists for complications exist including multiple use of the device and use with non-biocompatible materials materials/chemicals leading to infection , biologic exposure, inflammatory reaction.

DEVICE PREPARATION

1. Remove RingMaster from sterile packaging using aseptic technique and place on sterile procedure table.
2. Fill basin with sterile heparinized saline solution to desired level to cover devices in basin.
3. Carefully insert and remove devices as needed to prevent potential contamination.
4. Properly dispose of device according to hospital safety protocol.

WARNING

After use, dispose of device in a manner consistent with standard protocols for biohazard waste disposal.

SYMBOL	DESIGNATION
	Use By Date
	Lot Number
	Catalog Number
	Do Not Use if Package is Damaged and Consult Instructions for Use
	Single Use
	For electronic copy scan QR code or go to www.merit.com/ifu and enter IFU ID Number. For printed copy, call U.S.A or E.U Customer Service.
	Sterilized Using Ethylene Oxide
	Non-pyrogenic
	CAUTION: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.
	Manufacturer
	Medical Device
	Caution
	Date of Manufacturer
	Unique Device Identifier
	Single Sterile Barrier System



www.merit.com

**Manufacturer:**

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South Jordan, Utah 84095 U.S.A.
1-801-253-1600
U.S.A Customer Service 1-800-356-3748