

T-3[®] CT

SHORT-TERM HEMODIALYSIS

FEATURES & BENEFITS

INTERNAL LUMEN DESIGN

Power Injectable Triple



MATERIAL

Polyurethane

TIP DESIGN

Step Tip



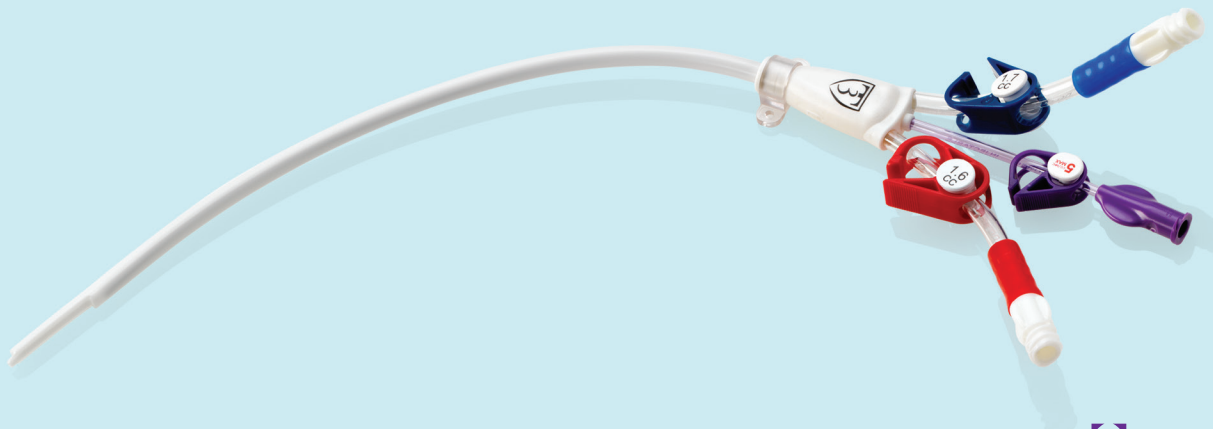
FRENCH SIZE

15.5F



CONFIGURATION

Straight



T-3[®] CT

SHORT-TERM HEMODIALYSIS CATHETER

| BASIC SET | | |
|-----------|--|-------|
| MC013501 | 15.5F X 15CM T-3 [®] CT, STRAIGHT | 5/BOX |
| MC013502 | 15.5F X 20CM T-3 [®] CT, STRAIGHT | 5/BOX |
| MC013503 | 15.5F X 24CM T-3 [®] CT, STRAIGHT | 5/BOX |
| MC013504 | 15.5F X 28CM T-3 [®] CT, STRAIGHT | 5/BOX |
| MC013505 | 15.5F X 32CM T-3 [®] CT, STRAIGHT | 5/BOX |

Basic Sets Contents: (1) Catheter (1) Guidewire (1) Introducer Needle (1) Scalpel (1) Suture (1) Valved Peelable Introducer (3) Dilator (3) End Cap (1) Adhesive Wound Dressing

IMPORTANT RISK INFORMATION

Indications For Use: The Medcomp[®] T-3[®] CT Catheter is a triple lumen catheter indicated for use in attaining short-term vascular access for hemodialysis, apheresis. The third internal lumen is intended for infusion, power injection of contrast media and central venous pressure monitoring. The catheter is intended to be inserted in the jugular, femoral or subclavian vein as required. The maximum recommended infusion rate is 5ml/sec for power injection of contrast media.

Contraindications: This catheter is intended for short-term (less than 30 days) vascular access only and should not be used for any purpose other than indicated in these instructions. This device is also contraindicated: When the presence of device related infection, bacteremia, or septicemia is known or suspected. When the patient's body size is insufficient to accommodate the size of the implanted device. When the patient is known or is suspected to be allergic to materials contained in the device. If the prospective insertion site has been previously irradiated. If the prospective placement site has previously suffered episodes of venous thrombosis or vascular surgical procedures. If local tissue factors may prevent proper device stabilization and/or access.

Refer to Instructions for Use provided with the product for complete instructions, warnings, and contraindications. Observe all Instructions for Use prior to using products. Failure to do so may result in patient complications.

For patient information please visit www.medcomp.net/patientinformation



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Certified Quality System

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