Long-Term Dialysis Catheter Repair Kit

BD (Becton Dickinson) offers a repair kit that lets you easily repair our line of polyurethane dialysis catheters. The repair kit replaces cracked or broken luer-lock connectors, clamps, and damaged extensions when a minimum of 4.5 cm of viable extension tubing exists.

The kit allows you to repair the following products:

- **DECATHLON® DF** Long-Term Hemodialysis Catheter
- **EQUISTREAM®** Long-Term Hemodialysis Catheter
- **EQUISTREAM® XK** Long-Term Hemodialysis Catheter
- **GLIDEPATH®** Long-Term Dialysis Catheter
- **HEMOSPLIT®** Long-Term Hemodialysis Catheter
- **HEMOSTAR®** Long-Term Hemodialysis Catheter
- **HEMOSTAR® XK** Long-Term Hemodialysis Catheter
- **RELIANCE XK™** Long-Term Dialysis Catheter
- **SOFT-CELL®** Long-Term Hemodialysis Catheter

**Catheter Repair Kit Contents:**
- 2 Instructions for Use
- 1 CSR Wrap
- 1 Label
- 1 Scissors
- 1 Temporary Slide Clamp (Green)
- 1 End Cap/Needleless Connector
- 1 Two-Piece Connector
- 2 Clamps: 1 Red, 1 Blue
- 1 Drape
Catheter Repair Kit

Indications
To replace: Cracked or broken female luer lock connectors or repair damaged extension where there is a minimum of 4.5 cm of viable extension tubing on the following catheters:
- Decathlon® DF Long-Term Dual Lumen Catheter
- Equistream® Long-Term Dual Lumen Catheter
- Equistream® XK Long-Term Dual Lumen Catheter
- Glidepath® Long-Term Dual Lumen Catheter
- HemoSplit™ Long-Term Dual Lumen Catheter
- HemoSplit™ XK Long-Term Dual Lumen Catheter
- HemoStar® Long-Term Dual Lumen Catheter
- HemoStar® XK Long-Term Dual Lumen Catheter
- Reliance XK™ Long-Term Dual Lumen Catheter
- Soft-Cell® Long-Term Dual Lumen Catheter

Contraindications
- Do not use to repair catheters other than those indicated above. Do not attempt connector replacement if usable length of remaining connector extension tubing is less than 4.5 cm. Do not replace connector if tubing is swollen or displays signs of degradation.

Warnings
- Alcohol or alcohol-containing antiseptics (such as chlorhexidine) may be used to clean the catheter/skin site; however, care should be taken to avoid prolonged or excessive contact with the solution(s). - Acetone and Polyethylene Glycol-containing ointments can cause failure of this device and should not be used with polyurethane catheters. Chlorhexidine patches or bacitracin zinc ointments (e.g., Polysporin™ ointment) are the preferred alternatives. - Intended for Single Use. DO NOT REUSE. Reuse and/or repackaging may create a risk of patient or user infection, compromise the structural integrity and/or essential material and design characteristics of the device, which may lead to device failure, and/or lead to injury, illness or death of the patient. - Place all clamps near the center of the polyurethane extension pieces. Clamping near or on the luer lock connectors may cause tubing fatigue and possible disconnection. - Polyurethane may develop cuts or tears if subjected to excessive pulling or contact with rough edges. - Accessories and components used in conjunction with this connector must incorporate luer lock adapters in order to avoid inadvertent disconnection. - The green slide clamp is provided for use during the repair procedure only. DO NOT REUSE the green slide clamp as it is not permanently attached and could separate from catheter, resulting in excessive bleeding. Dispose of the green slide clamp following the repair procedure. - After use, this product may be a potential biohazard. Handle and dispose of in accordance with accepted medical practice and applicable local, state and federal laws and regulations. - Failure to clamp could lead to air embolism or blood loss. - ALL CATHETER REPAIRS ARE DONE AT THE DISCRETION OF THE ATTENDING PHYSICIAN. THE MANUFACTURER WILL NOT ASSUME ANY LIABILITY FOR THE SAFETY OF A CATHETER AS A RESULT OF CATHETER REPAIR. - Do not use heparin in patients with heparin allergy.

Cautions
- Carefully read and follow all instructions prior to use. - Only qualified healthcare practitioners should repair damaged catheters. - Sterile and non-pyrogenic only if packaging is not opened, or broken. Sterilized with Ethylene Oxide. - Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician. - Strict aseptic technique must be used during the catheter repair procedure in accordance with Centers for Disease Control and National Kidney Foundation Clinical Practice Guidelines such as sterile gloves, sterile gown, surgical masks and caps (for patient and healthcare provider), and large sterile sheet. - Be sure to pull on the extension tubing and the connector only and not on the implanted portion of the catheter. - The priming volume of the repaired lumen will decrease by 0.1 mL for initial repair and 0.5 mL for every additional cm of extension tube removed. (See priming volume worksheet on back of IFU). - In the rare event of a leak, the catheter must be clamped immediately. Necessary remedial action must be taken prior to resuming dialysis treatment.

References

Please consult product labels and instructions for use for indications, contraindications, hazards, warnings and precautions.

Ordering Information

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<tr>
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Catheter Repair Kit With Replacement Connector

Please see instruction for Single Use. DO NOT REUSE. Reuse and/or repackaging may create a risk of patient or user infection, compromise the structural integrity and/or essential material and design characteristics of the device, which may lead to device failure, and/or lead to injury, illness or death of the patient. Please consult product labels and instructions for use for indications, contraindications, hazards, warnings and precautions.