Instructions for Use

VersaCross® Large Access Transseptal Dilator

Carefully read all instructions prior to use. Observe all warnings and precautions noted in these instructions. Failure to do so may result in patient complications. Baylis Medical Company relies on the physician to determine, assess and communicate to each individual patient all foreseeable risks of the procedure.

DEVICE DESCRIPTION

The VersaCross® Large Access Transseptal Dilator consists of two components: a dilator and a J-tipped guidewire. The VersaCross® Large Access Transseptal Dilator is designed for safe and easy catheterization and angiography of specific heart chambers and locations. The dilator provides superior torque control and is flexible. The dilator shaft in its entirety is coated with a hydrophobic lubricious coating for smoother device manipulation. No pre-conditioning is required for this coating. The dilator features a tapered tip and a shaft that can be reshaped manually. The echogenic shaft and tip and radiopaque tip maximizes visualization of the dilator during manipulation.

INDICATIONS FOR USE

Canada: The VersaCross® Large Access Transseptal Dilator is used for the percutaneous introduction of various types of cardiovascular catheters and guidewires to all heart chambers, including the left atrium via transseptal perforation / puncture.

United States: The VersaCross® Large Access Transseptal Dilator is indicated for use in procedures where access to the left atrium via the transseptal technique is desired.

WARNINGS

Laboratory staff and patients can undergo significant x-ray exposure during interventional procedures due to the continuous usage of fluoroscopic imaging. This exposure can result in acute radiation injury as well as increased risk for somatic and genetic effects. Therefore, adequate measures must be taken to minimize this exposure. The use of echocardiography is recommended.

- The VersaCross® Large Access Transseptal Dilator and accompanying guidewire are intended for single patient use only. Do not attempt to sterilize and reuse the VersaCross® Large Access Transseptal Dilator or accompanying guidewire. Reuse can cause the patient injury and/or the communication of infectious disease(s) from one patient to another. Failure to do so may result in patient complications.

- The VersaCross® Large Access Transseptal Dilator and accompanying guidewire are supplied STERILE using an ethylene oxide process. Do not use if the package is damaged.

- The dilator shaft is coated with a lubricious coating. The following warnings must be considered:
  - Use with sheaths smaller than the size listed in the section below may result in a tight fit that affects device performance, including coating integrity.
  - Excessive wiping and/or wiping with a dry gauze may damage the coating.
  - Manual shaping of the distal curve shall be done with smooth motions along the curve without applying excessive force and/or pressure. Excessive manual bending and/or shaping of the shaft may affect the coating integrity.

- Care should be taken to ensure all air is removed from the dilator before infusing through the proximal hub.
- Care should be taken when inserting or removing the dilator from introducer sheaths.
- Do not attempt direct percutaneous insertion of the dilator without a guidewire as this may cause vessel injury.
- Damage to guidewire may result if withdrawn through a metal needle cannula.
- Careful manipulation must be performed to avoid cardiac damage or tamponade. Dilator advancement should be performed under imaging guidance. Fluoroscopic or echocardiographic imaging is recommended.

PRECAUTIONS

- Do not attempt to use the VersaCross® Large Access Transseptal Dilator or accompanying guidewire before thoroughly reading the accompanying Instructions for Use.
- Careful manipulation must be performed to avoid cardiac damage, or tamponade. Dilator and guidewire advancement should be performed under imaging guidance, such as fluoroscopy or echocardiography. If resistance is encountered, DO NOT use excessive force to advance or withdraw the device.
- The sterile packaging, dilator, and guidewire should be visually inspected prior to use. Do not use the device if it has been compromised or damaged.
- Only physicians thoroughly trained in the techniques of the approach to be used should perform interventional procedures.
- Note product “Use By” date.
- The VersaCross® Large Access Transseptal Dilator is compatible with introducer sheaths 12.5Fr or larger.
- The VersaCross® Large Access Transseptal Dilator is compatible with .035” transseptal devices and guidewires
- The VersaCross® Large Access Transseptal Dilator is NOT compatible with transseptal needles such as the “NRG® Transseptal Needle”.

CONTRAINDICATIONS

- There are no known contraindications for this device.

SPECIAL STORAGE AND/OR HANDLING INSTRUCTIONS

Keep away from sunlight.

ADVERSE EVENTS

Adverse events that may occur while using the VersaCross® Large Access Transseptal Dilator and accompanying guidewire include:

- Infection
- Local nerve damage
- Vessel spasm
- AV fistula formation
- Arrhythmias
- Hematoma
- Catheter entrapment
- Valve damage
- Air embolus
- Vessel trauma
- Pseudoaneurysm
- Atrial septal defect
- Perforation and/or tamponade
- Hemorrhage
- Embolic events
- Pericardial/pleural effusion

PREPARATION FOR USE

Prior to use of the VersaCross® Large Access Transseptal Dilator and accompanying guidewire, the individual components should be carefully examined for damage or defects, as should all equipment used in the procedure. Do not use defective equipment. Do not reuse the device.
**SUGGESTED DIRECTIONS FOR USE**

- Carefully read all instructions prior to use. Failure to do so may result in complications.
- Thoroughly flush the dilator and guidewire with heparinized saline solution prior to use.
- Perform a standard vein puncture of the right femoral vein using an access needle (not supplied).
- Introduce the guidewire and advance to required depth. If resistance is encountered, DO NOT use excessive force to advance or withdraw the guidewire. Determine the cause of resistance before proceeding.
- Leaving the guidewire in place, withdraw the needle. Enlarge the cutaneous puncture site as necessary.
- Note that a compatible access introducer sheath may be used at the venous cutaneous puncture site if desired. Refer to the compatible introducer sheath’s Instructions for Use for details and directions.
- The distal curvature of the dilator may be adjusted manually if desired. Manual shaping of the distal curve shall be done with smooth motions along the curve. Do not use excessive force of any type when reshaping.
- Thread the dilator over the guidewire using a slight twisting motion into the superior vena cava (SVC) under imaging guidance such as fluoroscopy or echocardiography. If resistance is encountered, DO NOT use excessive force to advance or withdraw the dilator over the guidewire. Determine the cause of resistance before proceeding.
- Use standard technique to position the dilator/guidewire assembly into the desired heart chamber.
- If transseptal puncture is required, refer to the Instructions for Use of the transseptal puncture device.
- Ensure the dilator is clear of air. To aspirate blood, use the dilator hub.
- Monitor the location of the radiopaque tip frequently under imaging guidance, such as fluoroscopy or echocardiography.
- Deliver a continuous heparinized solution infusion or aspirate periodically. This may help reduce the risk of thromboembolic complications due to thrombus formation, as there may be a possibility of thrombus development at the distal dilator tip or inside the dilator lumen. Also aspirate when removing the transseptal device or dilator.
- After removal of the dilator, use standard technique to achieve hemostasis.

**CLEANING AND STERILIZATION INSTRUCTIONS**

Do not clean or re-sterilize the VersaCross® Large Access Transseptal Dilator or the accompanying guidewire. The VersaCross® Large Access Transseptal Dilator and the accompanying guidewire are intended for single use only.

**DISPOSAL OF WASTE**

Treat the used device(s) as biohazardous waste and dispose of in compliance with standard hospital procedures.

**CUSTOMER SERVICE AND PRODUCT RETURN INFORMATION**

If you have any problems with or questions about Baylis Medical Equipment contact our technical support personnel.

Baylis Medical Company Inc.
5959 Trans-Canada Highway
Montreal, Quebec, Canada, H4T 1A1
Phone: (514) 488-9801 or (800) 850-9801
Fax: (514) 488-7299
www.baylismedical.com

**NOTES:**

1. In order to return products you must have a return authorization number before shipping the products back to Baylis Medical Company. Product Return Instructions will be provided to you at this time.
2. Ensure that any product being returned to Baylis Medical has been cleaned, decontaminated and/or sterilized as indicated in the Product Return Instruction before returning it for warranted service. Baylis Medical will not accept any piece of used equipment that has not been properly cleaned or decontaminated as per the Product Return Instructions.

**LABELING AND SYMBOLS**

- Manufacturer
- Rx ONLY
- Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.
- Sterile using ethylene oxide
- Single Use – Do not reuse
- Use By
- Batch Code
- LOT
- Do Not Use if Packaging is Damaged
- Consult Instructions for Use
- Keep Away from Sunlight
- Model number
- Do Not Re-sterilize
- Non-Pyrogenic
- Maximum guidewire outside diameter that can be used with this device

**LIMITED WARRANTIES**

Baylis Medical Company Inc. (BMC) warrants its Disposable and Accessory products against defects in materials and workmanship. BMC warrants that sterile products will remain sterile for a period of time as shown on the label as long as the original package remains intact. Under this Limited Warranty, if any covered product is proved to be defective in materials or workmanship, BMC will replace or repair, in its absolute and sole discretion, any such product, less any charges to BMC for transportation and labor costs incidental to inspection, removal or restocking of product. The length of the warranty is: (i) for the Disposable products, the shelf life of the product, and (ii) for the Accessory products, 90 days from shipment date. This limited warranty applies only to new original factory delivered products that have been used for their normal and intended uses. BMC’s Limited Warranty shall not apply to BMC products which have been re-sterilized, repaired, altered, or modified in any way and shall not apply to BMC products which have been improperly stored or improperly cleaned, installed, operated or maintained contrary to BMC’s instructions.

**DISCLAIMER AND EXCLUSION OF OTHER WARRANTIES**

The limited warranty above is the sole warranty provided by BMC. Seller disclaims all other warranties, whether express or implied, including any warranty of merchantability or fitness for a particular use or purpose.

**LIMITATION OF LIABILITY FOR DAMAGES**

The remedy set forth herein shall be the exclusive remedy for any warranty claim, and additional damages, including consequential damages or damages for business interruption or loss of profit, revenue, materials, anticipated savings, data, contract, goodwill or the like (whether direct or indirect in nature) or for any other form of incidental or indirect damages of any kind. SHALL NOT BE AVAILABLE. Seller’s maximum cumulative liability relative to all other claims and liabilities, including obligations under any indemnity, whether or not insured, will not exceed the cost of the product(s) giving rise to the claim or liability. Seller disclaims all liability relative to gratuitous information or assistance provided by, but not required of seller hereunder. Any action against Seller must be brought within eighteen (18) months after the cause of action accrues. These disclaimers and limitations of liability will apply regardless of any other contrary provision hereof and regardless of the form of action, whether in contract, tort (including negligence and strict liability) or otherwise, and further will extend to the benefit of Seller’s vendors, appointed distributors and other authorized resellers as third-party beneficiaries. Each provision hereof which provides for a limitation of liability, disclaimer of warranty or condition or exclusion of damages is severable and independent of any other provision and is to be enforced as such.

In any claim or lawsuit for damages arising from alleged breach of warranty, breach of contract, negligence, product liability or any other legal or equitable theory, the Buyer specifically agrees that BMC shall not be liable for damages or for loss of profits, whether from Buyer or Buyer’s customers. BMC’s liability shall be limited to the purchase cost to Buyer of the specified goods sold by BMC to Buyer which give rise to the claim for liability.

No agent, employee or representative of Baylis Medical has the authority to bind the Company to any other warranty, affirmation or representation concerning the product.

This warranty is valid only to the original purchaser of Baylis Medical products directly from a Baylis Medical authorized agent. The original purchaser cannot transfer the warranty.

Use of any BMC product shall be deemed acceptance of the terms and conditions herein.

The warranty periods for Baylis Medical products are as follows:

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<th>Product Type</th>
<th>Shelf Life of the Product</th>
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<tr>
<td>Disposable Products</td>
<td>The shelf life of the product</td>
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<tr>
<td>Accessory Products</td>
<td>90 days from the shipment date</td>
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