Groshong*
Central Venous Catheters

Long Term

Instructions For Use
Introduction

Description

Groshong* Catheters consist of soft, medical grade silicone tubing with a closed rounded tip. Unlike open-ended catheters, the closed end has a patented three-position valve (or valves) which allows liquids to flow in or out, but remains closed when not in use.

In addition to the Groshong* Valve, Groshong* Catheters have the following features:

- Soft, Medical Grade Silicone Tubing
- Radiopaque Tip
- Radiopaque Stripe
- SureCuff* Tissue Ingrowth Cuff
- Winged Connector(s)
- Connector Locking Sleeve(s)
- Depth Markings
- VitaCuff* Antimicrobial Cuff
- Attachable Suture Wings
- Large Lumen(s)
- Multiple sizes and configurations

Placement

The catheter is placed into one of the large central veins so the tip lies in the superior vena cava above the right atrium. It is tunnelled subcutaneously for several inches to the desired exit site. The SureCuff* Tissue Ingrowth Cuff, attached to the long-term catheter, is positioned 3-5 cms below the skin exit site in the tunnel. The cuff promotes tissue ingrowth to secure the catheter in place.
The benefits provided by the Groshong* valve are:

1. Increased patient safety due to reduced risk of air embolism or bleedback.
2. Virtual elimination of heparin flushing to maintain catheter patency.
3. Reduced need for catheter clamping.
4. Reduced need for flushing when the catheter is not in use (only flushed every seven days with sterile normal saline when not in use).

Groshong* multi-lumen catheters have Groshong* valves which are rotated and staggered, allowing the concurrent infusion of incompatible drugs. Each lumen of a multi-lumen catheter is treated separately for maintenance and irrigation purposes.

**Indications For Use**

Groshong* Long-Term Catheters are designed for long-term vascular access and for use in patients that lack adequate peripheral venous access. They are available in single lumen and multi-lumen catheters.

All Groshong* central venous catheters are designed for the administration of I.V. fluids, blood products, drugs, and parenteral nutrition solutions, as well as blood withdrawal.
VitaCuff* Antimicrobial Cuff

Description

The VitaCuff* device is designed to help provide protection against infections related to vascular access catheters. The outer, tissue-interfacing surface of the VitaCuff* device may help reduce the incidence of infection by incorporating an antimicrobial agent into the porous collagen matrix.

The VitaCuff* device is comprised of two concentric layers of material. The internal layer is constructed of specially formulated and processed medical grade silicone. The external, tissue-interfacing layer is VitaGuard* antimicrobial collagen matrix. The antimicrobial activity of the VitaGuard* material is attributable to the silver ions bound to the collagen matrix. The activity lasts until the VitaGuard* matrix is completely absorbed by the tissue in four to six weeks.

The VitaGuard* collagen sponge is initially in a compressed state for ease of insertion. After placement, the matrix absorbs physiological fluids, quickly expands to approximately twice its original size, and helps provide an antimicrobial barrier and a physical barrier at the exit site. Tissue ingrowth into the VitaGuard* collagen matrix occurs in a few days, further securing the catheter in place, and reducing catheter movement.

Caution: The antimicrobial cuff is not intended to be used as a treatment for catheter related infections. The antimicrobial cuff does not provide protection against “blood seeding” infection or infusate-related infection. It is not intended to provide protection from bacteria for longer than one month. The antimicrobial cuff should not be used on patients with known sensitivities to silver ions or collagen.

Contraindications, Warnings, Cautions and Precautions

Contraindications

The device is contraindicated whenever:

- The presence of device related infection, bacteremia, or septicemia is known or suspected.
- The patient’s body size is insufficient to accommodate the size of the implanted device.
- The patient is known or is suspected to be allergic to materials contained in the device.
- Severe chronic obstructive lung disease exists (percutaneous subclavian placement only.)
- Past irradiation of prospective insertion site.
- Previous episodes of venous thrombosis or vascular surgical procedures at the prospective placement site.
• Local tissue factors will prevent proper device stabilization and/or access.
• Do not use the antimicrobial cuff in patients with known sensitivities to silver or collagen.

**Warnings:**

• Intended for **Single Patient Use. DO NOT REUSE.**
  Bard Access Systems products are single use devices and should never be reimplanted. Reuse carries with it the attendant concern of cross-infection regardless of the cleaning or sterilization method. Resterilization of incompletely cleaned devices may not be effective. Any device that has been contaminated by blood should not be reused or resterilized.

• After use, this product may be a potential biohazard. Handle and discard in accordance with accepted medical practice and applicable local, state and federal laws and regulations.

• **Pinch-off Prevention:** Catheters placed percutaneously or through a cut-down, into the subclavian vein, should be inserted at the junction of the outer and middle thirds of the clavicle, lateral to the thoracic outlet. The catheter should not be inserted into the subclavian vein medially, because such placement can lead to compression of the catheter between the first rib and the clavicle, which can cause damage and even severance of the catheter. A radiographic confirmation of catheter placement should be made to ensure that the catheter is not being pinched by the first rib and clavicle. 1,2

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**Signs of Pinch-off**

**Clinical:**
• Difficulty with blood withdrawal
• Resistance to infusion of fluids
• Patient position changes required for infusion of fluids or blood withdrawal

**Radiologic:**
• Grade 1 or 2 distortion on chest X-ray.
Pinch-off should be evaluated for degree of severity prior to explantation. Patients indicating any degree of catheter distortion at the clavicle/first rib area should be followed diligently. There are grades of pinch-off that should be recognized with appropriate chest x-ray as follows: 3,4

<table>
<thead>
<tr>
<th>Grade</th>
<th>Severity</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade 0</td>
<td>No distortion</td>
<td>No action.</td>
</tr>
<tr>
<td>Grade 1</td>
<td>Distortion present without luminal narrowing</td>
<td>Chest x-ray should be taken every one to three months to monitor progression of pinch off to grade 2 distortion. Shoulder positioning during chest x-rays should be noted as it can contribute to changes in distortion grades.</td>
</tr>
<tr>
<td>Grade 2</td>
<td>Distortion present with luminal narrowing</td>
<td>Removal of the catheter should be considered.</td>
</tr>
<tr>
<td>Grade 3</td>
<td>Catheter transection or fracture</td>
<td>Prompt removal of the catheter.</td>
</tr>
</tbody>
</table>

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**Cautions:**

• Carefully read and follow all instructions prior to use.
• Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.
• Only qualified healthcare practitioners should insert, manipulate and remove these devices.
Precautions:

Follow Universal Precautions when inserting and maintaining the catheter.
Follow all contraindications, warnings, cautions, precautions and instructions for all infusates as specified by its manufacturer.

Use aseptic techniques whenever the catheter lumen is opened or connected to other devices. Povidone-iodine is the suggested antiseptic to use with this device and components. Acetone and tincture of iodine should not be used because they could adversely affect the performance of the catheter and connectors. 10% acetone/70% isopropyl alcohol swabsticks used for dressing changes should not adversely affect the catheter.

I. Prior to beginning placement procedure, do the following:

- Examine package carefully before opening to confirm its integrity and that the expiration date has not passed. The device is supplied in a double sterile package and is non-pyrogenic. Do not use if package is damaged, opened or the expiration date has passed. Sterilized by ethylene oxide. Do not Resterilize.
- Inspect kit for inclusion of all components.
- When device includes an antimicrobial cuff, do not expose the cuff to fluids prior to insertion. Handle carefully to avoid cuff damage.
- Fill (prime) the device with normal saline solution to help avoid air embolism.
- When using an introducer kit, verify that the catheter fits easily through the introducer sheath.

II. To avert device damage and/or patient injury during placement:

- Avoid accidental device contact with sharp instruments and mechanical damage to the catheter material. Use only smooth-edged atraumatic clamps or forceps.
- Avoid perforating, tearing or fracturing the catheter when using a guidewire.
- Do not use the catheter if there is any evidence of mechanical damage or leaking.

- Avoid sharp or acute angles during implantation which could compromise the patency of the catheter lumen(s).
- Use suture wings to secure catheters.
- Do not place sutures directly around the catheter.
- When using percutaneous introducers:
  - Carefully insert the introducer and catheter to avoid inadvertent penetration to vital structures in the thorax.
  - To avoid blood vessel damage, do not allow the percutaneous introducer sheath to remain indwelling in the blood vessel without the internal support of a catheter or dilator.
  - Simultaneously advance the sheath and dilator with rotational motion to help prevent sheath damage.
- During insertion of catheter with antimicrobial cuff:
  - Minimize the exposure of the cuff to pooled blood by sponging the intended cuff placement site.
  - The entire collagen (tan) portion of the cuff must be in the subcutaneous tissue at the catheter exit site.

III. After insertion, observe the following precautions to avoid device damage and/or patient injury:

- Do not use the catheter if there is any evidence of mechanical damage or leaking. Damage to the catheter may lead to rupture, fragmentation and possible embolism and surgical removal.
- Accessories and components used in conjunction with this device should incorporate Luer lock connections.
- If signs of extravasation exist, discontinue injections. Begin appropriate medical intervention immediately.
- Infusion pressure greater than 25 psi (172 kPa) may damage blood vessels and viscus and is not recommended. DO NOT USE A SYRINGE SMALLER THAN 10 ml!
**Possible Complications**

The use of an indwelling central venous catheter provides an important means of venous access for critically ill patients; however, the potential exists for serious complications including the following:

- Air Embolism
- Allergic Reaction to Silver or Collagen (Catheters with VitaCuff* Antimicrobial Cuff only)
- Bleeding
- Brachial Plexus Injury
- Cardiac Arrhythmia
- Cardiac Tamponade
- Catheter or Cuff Erosion Through Skin
- Catheter Embolism
- Catheter or Cuff Occlusion
- Catheter Occlusion, Damage or Breakage due to Compression Between the Clavicle and First Rib
- Catheter-related Sepsis
- Endocarditis
- Exit Site Infection
- Exit Site Necrosis
- Extravasation
- Fibrin Sheath Formation
- Hematoma
- Hemotherox
- Hydrothorax
- Intolerance Reaction to Implanted Device
- Laceration of Vessels or Viscus
- Perforation of Vessels or Viscus
- Pneumothorax
- Spontaneous Catheter Tip Malposition or Retraction
- Thoracic Duct Injury
- Thromboembolism
- Venous Thrombosis
- Ventricular Thrombosis
- Vessel Erosion
- Risks Normally Associated with Local and General Anesthesia, Surgery, and Post-Operative Recovery

These and other complications are well documented in medical literature and should be carefully considered before placing the catheter.

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**Groshong* Central Venous Catheter Placement Procedures**

Before beginning procedure, read the “Contraindications, Warnings, Cautions and Precautions” and “Possible Complications” sections of this manual.

**Section A: Cutdown**

1. Create sterile field and open tray.
2. Prep cutdown area, tunnel and tunnel exit areas.
3. Perform local anesthetic infiltration in cutdown area and along pathway chosen for the subcutaneous catheter tunnel.
4. Irrigate the catheter with sterile normal saline via the flushing hub:
   - **Single-Lumen Catheters** have catheter-stylet attached to the flushing hub.
   - **Multi-Lumen Catheters** have a flushing hub and no stylet.
5. Place patient in the Trendelenburg position with head turned away from the intended venipuncture site.
6. Surgically isolate the desired vessel through a small skin incision.
7. Insert the catheter assembly and advance to desired position in vessel by using the graduations on the catheter which are in 2.5 cm (3.5 Fr catheter) or 5 cm (all other catheters) intervals from the distal tip.

Refer to the “Warnings” section concerning Catheter Pinch-off.
8. Verify catheter tip location radiographically. The preferred location of the catheter tip is at the junction of the superior vena cava and the right atrium. **Caution:** Avoid positioning the catheter tip in the right atrium.

9. Measure catheter against chest wall of patient to determine desired location of SureCuff® tissue ingrowth cuff and exit site. Mark locations.

**For Tunneling Instructions refer to Section C.**

**Section B: Percutaneous**

1. Create sterile field and open tray.
2. Prep venipuncture, tunnel and exit site areas. Position drape.
3. Perform local anesthetic infiltration in venipuncture, tunnel and tunnel exit areas.
4. Irrigate the catheter with sterile normal saline via the flushing hub:
   - **Single-Lumen Catheters** have catheter-stylet integrated in the flushing hub.
   - **Multi-Lumen Catheters** have a flushing hub and no stylet.
5. Place patient in the Trendelenburg position with head turned away from the intended venipuncture site.
6. Locate desired vessel using a small needle attached to a syringe. **Note:** The subclavian vein is entered percutaneously at the point that identifies the junction of the outer and middle thirds of the clavicle using the needle and syringe.

Refer to the “Warnings” section concerning Catheter Pinch-off.

7. Attach the introducer needle to syringe and insert into vessel alongside the small needle. Remove small needle.
8. Aspirate gently as the insertion is made. **Warning:** If the artery is entered, withdraw the needle and apply manual pressure for several minutes. If the pleural space is entered, withdraw the needle and evaluate patient for possible pneumothorax.

9. When the subclavian vein has been entered, remove the syringe leaving the needle in place. **Place a finger over the hub of the needle to minimize blood loss and the risk of air aspiration.** The risk of air aspiration is reduced by performing this part of the procedure with the patient performing the Valsalva maneuver.

10. Straighten “J” tip of guidewire with tip straightener and insert tapered end of tip straightener into the needle. **Tip straightener should not be advanced over the guidewire beyond the guidewire tip.** **Caution:** Do not insert guidewire beyond the bevel of the needle while removing straightener from the needle, in order to prevent guidewire shearing. Remove the tip straightener and advance the guidewire into the superior vena cava. Advance the guidewire as far as appropriate for the procedure. **Verify correct positioning radiographically.**
11. Gently withdraw and remove needle.
   **Caution:** If the guidewire must be withdrawn while the needle is inserted, remove both the needle and guidewire as a unit to help prevent the needle from damaging or shearing the guidewire.

12. Make a small (approx. 1 cm wide) incision parallel to the clavicle, positioning the guidewire at the center of the incision to permit proper entry of vessel dilator and sheath introducer.

(For Peel-Apart Introducer instructions proceed to step 20)

**Intro-Eze® Introducer Instruction:**

13. Advance the vessel dilator and sheath introducer as a unit over the exposed guidewire using a rotational motion. Advance it into the subclavian vein as a unit, leaving at least 2 cms of sheath exposed. **Warning:** Avoid vessel perforation.

14. Withdraw the vessel dilator and “J” guidewire, leaving the sheath in place. **Warning:** Hold thumb over exposed orifice of sheath to prevent air aspiration. The risk of air aspiration is reduced by performing this part of the procedure with the patient performing the Valsalva maneuver.

15. Advance the catheter through the sheath and into the vein.

16. Verify catheter tip location radiographically. **Warning:** Avoid positioning the catheter tip in the right atrium. The preferred location of the catheter tip is at the junction of the superior vena cava and the right atrium.

17. Pull the storage tube from the slitter. Place the channeled portion of the slitter onto the catheter near the proximal end of the introducer sheath.

18. Grasp the proximal end of the slitter between the thumb and index finger of one hand. With the tips of the fingers, reach around the slitter and secure the catheter into the channeled portion.
19. Withdraw the sheath over the catheter, sliding the proximal opening of the sheath over the nose of the channel and into the blade. Continue to withdraw the sheath, pulling it away from the catheter until it is completely slit, remove and discard the slit sheath and slitter.

20. Advance the vessel dilator and sheath introducer as a unit over the exposed guidewire using a rotational motion. Advance it into the subclavian vein as a unit, leaving at least 2 cms of sheath exposed. **Warning:** Avoid vessel perforation.

21. Squeeze the hub handles together releasing the locking mechanism and gently withdraw the vessel dilator and “J” guidewire, leaving the sheath in place.

22. **Warning:** Hold thumb over exposed orifice of sheath to prevent air aspiration. The risk of air aspiration is reduced by performing this part of the procedure with the patient performing the Valsalva maneuver.

23. Insert catheter-stylet assembly into lumen of sheath and advance to desired position in vessel by using the graduations on the catheter which are in 2.5 cm (3.5 Fr catheter) or 5 cm (all other catheters) intervals from the distal tip.

24. Verify catheter tip location radiographically. **Warning:** Avoid positioning the catheter tip in the right atrium. The preferred location of the catheter tip is at the junction of the superior vena cava and the right atrium.

25. Grasp the two handles of the peel-apart sheath and pull outward and upward at the same time.

Proceed to section C.
26. Peel the sheath away from the catheter completely. Make sure the catheter is not dislodged from vessel as sheath is removed.

27. Use the depth markings on the catheter to assure proper insertion depth.

Section C: Tunneling
(For multi-lumen catheter proceed to step 31)

Inserting a Single Lumen Catheter:
28. Slide flushing hub out of catheter. Gently withdraw stylet while firmly holding catheter. Do not permit catheter to be withdrawn from vessel while removing the stylet.

   **WARNING:** Do not leave stylet in catheter.

29. Create subcutaneous tunnel to exit site using tunneler.
   a. Continue pushing sharp point of tunneler through skin to create exit site.
   b. Attach catheter to tunneler extension.
   c. Secure catheter to tunneler with suture.
   d. Pull tunneler with secured catheter out of exit site. Initial resistance may be met as **VitaCuff** Antimicrobial Cuff or **SureCuff** Tissue Ingrowth Cuff first enters the tunnel. Gently holding the catheter distal to the cuff while pulling the tunneler and catheter through subcutaneous tunnel should result in smooth passage of the cuff(s) through the subcutaneous tunnel. **Caution:** The catheter must not be forced.
   e. Avoid subcutaneous kinking of catheter between the venipuncture site and tunnel curvature point.

30. Cut catheter from tunneler extension and trim external segment of catheter to desired length using 90° cut.

   Proceed to section D.

Inserting a Multi-Lumen Catheter:
31. The multi-lumen catheter has no stylet that needs to be removed.

32. Create subcutaneous tunnel to exit site using tunneler.
   a. Continue pushing sharp point of tunneler through skin to create exit site.
   b. Cut flushing hub assembly from proximal end of catheter.
   c. Attach catheter to tunneler extension.
   d. Secure catheter to tunneler with suture.
   e. Pull tunneler with secured catheter out of exit site. Initial resistance may be met as **VitaCuff** Antimicrobial Cuff or **SureCuff** Tissue Ingrowth Cuff first enters the tunnel. Gently holding the catheter distal to the cuff while pulling the tunneler and catheter through subcutaneous tunnel should result in smooth passage of the cuff(s) through the subcutaneous tunnel. **Caution:** The catheter must not be forced.
   f. Avoid subcutaneous kinking of catheter between the venipuncture site and tunnel curvature point.

33. Cut the extension tubes at the “Y” connection adjacent to the tunneler to allow separation of the catheter lumens.

Section D: Connector Attachment
34. Secure connector(s) to catheter per the following instructions:
   a. **During Catheter Placement:** Cut catheter at a 90° angle and trim external segment of catheter to desired length. For multi-lumen catheters, match the connector hub color to the color of the respective tubing stripes. Proceed to b.
   
   **When replacing connector.** Using aseptic technique, cut the catheter off at a 90° angle, 0.5 in. (12.7 mm) distal to the
Section E: Catheter Securement

37. Suture venipuncture site as necessary, taking care not to damage the catheter.

38. Suture catheter at exit site using the provided movable suture wing. Remove suture wing from card. Pinch the movable suture wings together to open the split underside of the wing body. Place the wing body onto the catheter at the exit site and release. Secure wing in place with suture through holes in wing (avoid nicking catheter with suture needle) or sterile skin closure tape.

39. Secure catheter at exit site with a sterile dressing. The external segment of the catheter should be coiled and taped. Many institutions have recommended the use of an extension set. Avoid tension on the external segment to prevent dislodging the catheter.

35. Draw blood through catheter to insure patency after placement is complete, but before suturing the venipuncture site. If catheter is not patent, adjust catheter at curvature point to relieve possible restriction. Flush catheter with 10ml sterile saline solution to clear catheter of blood and create a saline lock.

Multi-Lumen — Draw blood through red lumen to insure patency and irrigate all lumens with 10ml of sterile saline.

36. Attach injection cap(s) or connect to intravenous fluid source.
**Groshong® Catheter Removal**

After tissue grows into the **SureCuff® Tissue Ingrowth Cuff** (2 to 3 weeks), catheters can be removed from the subcutaneous tunnel using one of several methods. The method used will depend upon physician preference and the amount of tissue/cuff ingrowth that is present. The catheter can usually be removed by traction on the external segment (see #1 below) if it is not sutured internally at the cuff or vessel insertion site. Surgical removal (see #2 below) may be necessary to prevent breaking the catheter if the catheter does not dislodge easily with traction or if there is no definite suture site information.

**Warning:** You should not feel any resistance when withdrawing the catheter from the vein. If you do encounter resistance, this may indicate that the catheter is being pinched between the clavicle and first rib (the “pinch-off” sign). Do not continue pulling against resistance as this may cause catheter breakage and embolism. Free up the resistance (e.g. by repositioning the patient) before proceeding further.

1. **Traction Removal**
   
   Pull the catheter external segment downward in a straight line away from the exit site with a series of gentle tugs. When separation of the cuff from the surrounding tissue and/or catheter occurs, there will be a “break-away” feeling. Continue to pull gently on the catheter to complete the removal. Apply pressure to the catheter/vein insertion site as needed to control bleeding. If the cuff remains in the subcutaneous tissue, dissect it out through a small incision utilizing local anesthesia.

2. **Surgical Removal (using aseptic technique)**
   
   a) Locate the position of the cuff either by palpation or by observing the position of “dimpling” when traction is applied to the catheter’s external segment.

   b) Make a short transverse incision at or below the external side of the cuff taking care not to transect the catheter. Reach under the catheter with a curved, smooth-jawed clamp and pull up on the catheter to remove the catheter tip from the vein. **Caution:** Do not grasp the catheter with any instrument that might sever or damage the catheter.

   c) Dissect out the cuff. Transect the catheter on the exterior side of the cuff and remove the interior portion of the catheter and cuff through the incision.  

   **Caution:** Do not cut the catheter before removal from vein to avoid catheter embolism.

   d) Remove the exterior segment of the catheter by pulling it from the skin exit site.

   e) Apply pressure to the catheter/vein insertion site as needed to control bleeding.

   f) Close the incision with a suture as needed. Apply antibiotic ointment to incision and skin exit sites and an occlusive dressing to prevent air embolism through the tract.

Catheter care and maintenance procedures are included in the **Groshong® CV Catheter Nursing Procedure Manual** available through **Bard Access Systems** Customer Service, 1-800-545-0890. For outside the U.S., contact your local sales representative or distributor.

**References**


Patient Information - Catheter Care and Maintenance

Catheter Damage
If the catheter or connection is damaged or dislodged during or after surgery, immediately clamp the catheter with an atraumatic catheter clamp or kink and tape it. The catheter should be repaired as soon as possible using the designated Groshong* repair kit for that particular catheter size. Damage close to the catheter hub can be repaired using the appropriate Groshong* replacement connector. Instructions are enclosed in the repair kit and replacement connector packages and are also available in the Groshong* CV Catheter Nursing Procedure Manual.

Site Care

Supplies you will need:
- Sterile gloves (if required)
- 3 Alcohol swabsticks
- Hydrogen peroxide
- 3 Povidone iodine swabsticks
- Povidone iodine ointment packet
- 1 Alcohol wipe
- Sterile 2 in. x 2 in. (5 cm x 5 cm) gauze dressing
- 1 Sterile pre-cut 2 in. x 2 in. (5 cm x 5 cm) gauze dressing
- Sterile cotton tipped applicators
- 1 Sterile cover dressing (transparent or tape)
- Tape

1. Clean the work surface by wiping with a paper towel that has been moistened with alcohol. Wipe dry or allow to air dry. Then place supplies on the cleaned surface.
2. **Wash your hands thoroughly using warm soapy water.** Rinse completely and dry using a clean towel or fresh paper towels.
3. Carefully open the dressing kit, or unwrap supplies, without touching the inside surfaces of the kits or wrappers.
4. Carefully remove the old dressing, starting from the top of the dressing and working downward. Remove the tape or dressing carefully to avoid irritating your skin or pulling on the catheter. **Caution:** Do not use scissors or any sharp-edged instruments as they could damage the catheter.
5. Wash your hands again.
6. Do a careful observation of the exit site and the skin around it. If you notice anything unusual, finish the dressing procedure and then call your doctor.
7. If you are instructed to use gloves, put on the pair of sterile gloves following the procedure you were taught. Be careful to not touch anything except the supplies being used for site care.
8. Carefully clean the catheter exit site with an alcohol swabstick or sterile cotton tipped applicator, soaked in hydrogen peroxide, starting at the exit site and spiraling outward until a circle, at least 8 cm. in diameter, has been cleaned. Do not return to the catheter exit site with a swabstick that has touched any skin away from the exit site.
9. Repeat this step using the other two swabsticks. Look at the color of the swabsticks for signs of drainage.
10. Repeat step 8 using three of the povidone iodine swabsticks to clean the same skin area again as well as the part of the catheter that will be lying on the cleaned skin.
11. Gently clean the outside of the catheter with the inside surface of an alcohol wipe, starting from the exit site to the catheter connector sleeve. You may hold the catheter at the exit site with another alcohol wipe to prevent pulling on the catheter. **DO NOT PULL ON THE CATHETER.**
Clamping the Catheter

Under normal circumstances, your catheter will not need to be clamped. If damage to the catheter occurs, the catheter should be clamped immediately.

- Use only smooth-edged clamps.
- Follow the directions or your doctor or nurse regarding when to clamp.

When should you clamp?

You should clamp if there is any damage to the catheter or the catheter connector or if there is any separation of the catheter and the catheter connector: \textit{Always have a clamp available for emergencies.}

Flushing the Catheter

Supplies you will need:

- Alcohol or povidone iodine wipe.
- 10ml syringe or larger with attached one inch needle filled with 5ml of normal saline, prepared for use.
- Tape.

The steps in the procedure are:

1. Collect your supplies in a convenient place.
2. Wash your hands thoroughly.
3. Remove the tape that is around the cap.
4. Clean the cap with an alcohol or povidone iodine wipe. If you use the iodine wipe, allow the cap to air dry for two minutes—be sure not to touch the cap during this time. Do not blow on the area or allow the clean cap to dangle since this increases the chance of contamination of the area with germs.
5. Remove the needle cover and carefully insert the needle into the center of the catheter injection cap.
6. Inject the normal saline into the catheter. As you inject the last 0.5 ml, withdraw the needle from the injection cap. If you are...
flushing the catheter of a child, do not flush too rapidly because the child’s circulatory system is small and sensitive to rapid changes in volume and pressure.

where the cap is connected to the catheter. Allow to air dry.

5. **While holding the catheter connector below the level of your heart**, unscrew the old cap and discard. (The fluid level in the catheter will drop part-way into the catheter if the connector is held above the level of your heart.)

6. Pick up the new cap only by the top and remove the sterile tip protector. Attach the new cap by firmly screwing it onto the catheter connector.

7. Cut a 5 cm piece of tape and make tabs on each end by folding back 1 cm. Apply the sticky part of the tape around the connection of the cap and catheter and fasten securely.

8. Press ends of the tape together. The tabs on the end of the tape will enable you to remove it very easily.

If you have a multi-lumen catheter, use a separate syringe to flush each lumen with sterile normal saline. **Your doctor or nurse will give you additional information for the care of multi-lumen catheters.**

### Changing the Injection Cap

**Supplies you will need:**
- Sterile injection cap.
- Alcohol or povidone iodine wipe.
- Tape.

**The procedure to change the cap:**

1. Wash your hands thoroughly.

2. Open the package of the new injection cap and prepare according to your instructions. Be sure the cap does not touch the outer surface of the package.

   **NOTE:** You may need to prefill the injection cap with sterile normal saline if it is a long cap with significant air space. Your doctor or nurse will teach you this additional procedure.

3. Remove the old tape from around the cap by unpeeling the tape. **NEVER** attempt to cut the tape with scissors as you may damage the catheter.

4. Using an alcohol or povidone iodine wipe, clean around the place
An issued or revision date for these instructions is included for the user’s information. In the event two years have elapsed between this date and product use, the user should contact Bard Access Systems to see if additional product information is available.

Revised Date: February 2007

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Covered by one or more of following U.S. Patents: D498,844; 5,160,325; 5,221,263.

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