Safety and Efficacy Considerations:
The product is designed for single use only. Do not resterilize or reuse. Do not alter the sheath or any other kit/set component during insertion, use, or removal. Procedure must be performed by trained personnel well versed in anatomical landmarks, safe technique and potential complications.

Indications for Use:
The Arrow percutaneous sheath introducer permits venous access and catheter introduction to the central circulation. The SLIC® is a two-piece assembly consisting of an infusion catheter and an obturator. With SLIC® obturator removed, the Arrow SLIC® permits access to the central venous circulation through an indwelling sheath/hemostasis valve. With the SLIC® obturator in place, the SLIC® occludes the hemostasis valve minimizing the risk of air entry and blood loss through the valve.

Contraindications:
None known.

Warnings and Precautions:
1. Warning: Practitioners must be aware of complications associated with percutaneous sheath introduction including vessel wall perforation, pleural and mediastinal injuries, air embolism, sheath embolism, thoracic duct laceration, bacteremia, septicemia, thrombosis, inadvertent arterial puncture, nerve damage, hematoma formation, hemorrhage, and dysrhythmias.
2. Warning: Do not apply excessive force in removing guide wire, dilator or sheath. If withdrawal cannot be easily accomplished, a chest x-ray should be obtained and further consultation requested.
3. Warning: The practitioner must be aware of potential air embolism problems associated with leaving open needles, sheaths, or catheters in venous puncture sites or as a consequence of inadvertent disconnects. To lessen the risk of disconnects, only securely tightened Luer-Lock connections should be used with this device. Follow hospital protocol for all sheath and side port maintenance to guard against air embolism.
4. Warning: Hemostasis valve/side port assembly to SLIC® connection and SLIC® to obturator connection must be secured and routinely examined to avoid disconnection and possible air embolism, hemorrhage, or exsanguination.
5. Warning: Hemostasis valve must be occluded at all times to minimize the risk of air embolism or hemorrhage. If catheter introduction is delayed, or catheter is removed, temporarily cover valve opening with sterile-gloved finger until catheter or SLIC® assembly is inserted. Use Arrow SLIC® assembly, included with this product, as dummy catheter with valve/side port assembly and sheath. This will ensure that leakage does not occur and inner seal is protected from contamination.
6. Warning: Care should be exercised in passing spring-wire guide. Use of excessive length of the guide wire into the right heart can cause dysrhythmias, right bun-
dle branch block, and vessel wall, atrial or ventricular perforation.

7. Warning: Due to the risk of exposure to HIV (Human Immunodeficiency Virus) or other blood borne pathogens, health care workers should routinely use universal blood and body-fluid precautions in the care of all patients.

8. Precaution: Do not suture directly to the outside diameter of the sheath to avoid cutting or damaging the sheath or impeding sheath flow.

9. Precaution: Indwelling sheaths should be routinely inspected for desired flow rate, security of dressing, correct position and for proper Luer-Lock connection.

10. Precaution: Maintain the insertion site with regular meticulous redressing using aseptic technique.

11. Precaution: Alcohol and acetone can weaken the structure of polyurethane material. Therefore, care should be taken when instilling drugs containing alcohol or when using high concentration of alcohol or acetone when performing routine insertion site care and maintenance. Alcohol should not be utilized to declot polyurethane sheaths.

Carefully read all warnings and precautions throughout procedure instructions.

A Suggested Procedure:
Use sterile technique.

1. Precaution: Place patient in slight Trendelenburg position as tolerated to reduce the risk of air embolism. If femoral approach is used, place patient in supine position.

2. Prep area of anticipated venipuncture.

3. Drape puncture site as required.

4. Perform skin wheal using desired needle. In kits where provided, a SharpsAway® disposable cup is used for the disposal of needles. Push needles into foam after use. Discard entire cup at completion of procedure. Precaution: Do not reuse needles after they have been placed into the disposal cup. Particulate matter may adhere to needle tip.

5. Insert tip of desired catheter through Tuohy Borst adapter end of catheter contamination shield. Advance catheter through tubing and hub at other end (refer to Fig. 1).

6. Slide entire catheter contamination shield to proximal end of catheter.

7. If flow directed catheter is used, inflate and deflate balloon with syringe to ensure integrity. Precaution: Do not exceed balloon catheter manufacturer’s recommended volume. Place catheter and catheter contamination shield on sterile field awaiting final sheath placement.

8. Insert entire length of dilator through hemostasis valve into sheath pressing hub of dilator firmly into hub of hemostasis valve/side port assembly. Place assembly on sterile field awaiting final sheath placement.

9. Insert introducer needle with attached Arrow Raulerson Syringe into vein and aspirate. (If larger introducer needle is used, vessel may be pre-located with 22 Ga. locator needle and syringe). Remove locator needle.

Alternate technique:
Catheter/needle may be used in the standard manner as alternative to introducer needle. If catheter/needle is used, Arrow Raulerson Syringe will function as a standard syringe, but will not pass spring-wire guide. If no free flow of venous blood is observed after needle is removed, attach syringe to the catheter and aspirate until good venous blood flow is established. Precaution: The color of the blood aspirated is not always
a reliable indicator of venous access. Do not reinsert needle into introducer catheter.

10. Because of the potential for inadvertent arterial placement, one of the following techniques should be utilized to verify venous access. Insert the fluid primed blunt tip transduction probe into the rear of the plunger and through the valves of the Raulerson Syringe. Observe for central venous placement via a waveform obtained by a calibrated pressure transducer. Remove transduction probe (refer to Fig. 2).

**Fig. 2**

Alternate technique:
If hemodynamic monitoring equipment is not available to permit transducing a central venous waveform, check for pulsatile flow by either using the transduction probe to open the syringe valving system or by disconnecting the syringe from the needle. Pulsatile flow is usually an indicator of inadvertent arterial puncture. **Precaution:** The color of the blood aspirated is not always a reliable indicator of venous access.

11. Using the two-piece Arrow Advancer™, advance spring-wire guide through syringe into vein. **Warning:** Aspiration with spring-wire guide in place will cause introduction of air into syringe. **Precaution:** To avoid leakage of blood from syringe cap do not reinfuse blood with spring-wire guide in place.

**Arrow Two-Piece Advancer™ Instructions:**
- Using your thumb, straighten the “J” by retracting the spring-wire guide into the Advancer™ (refer to Fig. 3). When the tip is straightened, the spring-wire guide is ready for insertion. Centimeter marks are referenced from “J” end. One band indicates 10 cm, two bands 20 cm, and three bands 30 cm.

**Fig. 3**

**Introducing the Spring-Wire Guide:**
- Place the tip of the Advancer™ – with “J” retracted – into the hole in the rear of the Raulerson Syringe plunger (refer to Fig. 4).

**Fig. 4**

- Advance spring-wire guide into the syringe approximately 10 cm until it passes through the valves.
- Lift your thumb and pull the Advancer™ approximately 4 cm to 8 cm away from the syringe. Lower thumb onto the Advancer™ and while maintaining a firm grip on the spring-wire guide, push the assembly into the syringe barrel to further advance the spring-wire guide. Continue until spring-wire guide reaches desired depth (refer to Fig. 5).

**Fig. 5**

**Alternate technique:**
If a simple straightening tube is preferred, the straightening tube portion of the Advancer™ can be disconnected from the unit and used separately. Separate the Advancer™ tip or straightening tube from the blue Advancer™ unit. If the “J” tip portion of the spring-wire guide is used, prepare
for insertion by sliding the plastic tube over the “J” to straighten. The spring-wire guide should then be advanced in the routine fashion to the desired depth.

12. Advance the guide wire until triple band mark reaches rear of syringe plunger. Advancement of “J” tip may require a gentle rotating motion. **Warning:** Do not cut spring-wire guide to alter length. Do not withdraw spring-wire guide against needle bevel to avoid possible severing or damaging of spring-wire guide.

13. Hold spring-wire guide in place and remove introducer needle and Raulerson Syringe (or catheter). **Precaution:** Maintain firm grip on spring-wire guide at all times. Use centimeter markings on spring-wire guide to adjust indwelling length according to desired depth of indwelling sheath placement.

14. Enlarge cutaneous puncture site with cutting edge of scalpel positioned away from the spring-wire guide. **Precaution:** Do not cut guide wire.

15. Thread tapered tip of dilator/sheath/valve assembly over spring-wire guide. Grasping near skin, advance assembly with slight twisting motion to a depth sufficient to enter vessel. Dilator may be partially withdrawn to facilitate advancement of sheath through tortuous vessel. **Precaution:** Do not withdraw dilator until the sheath is well within the vessel to minimize the risk of damaging the sheath tip.

16. Advance sheath/valve assembly off dilator into vessel, again grasping near skin and using slight twisting motion.

17. To check for proper sheath placement within the vessel, remove side port end cap and attach syringe for aspiration. Hold sheath/valve assembly in place and withdraw spring-wire guide and dilator sufficiently to allow venous blood flow to be aspirated into side port. **Precaution:** Maintain firm grip on spring-wire guide at all times.

18. Holding sheath/valve assembly in place, remove guide wire and dilator as a unit. Place sterile-gloved finger over hemostasis valve. **Warning:** To minimize the risk of possible vessel wall perforation do not leave vessel dilator in place as an indwelling catheter. **Warning:** Although the incidence of spring-wire guide failure is extremely low, practitioner should be aware of the potential for breakage if undue force is applied to the wire. Flush and connect side port to appropriate line as necessary.

19. Feed catheter through sheath/valve assembly into vessel. Advance catheter to desired position. **Warning:** Hemostasis valve must be occluded at all times to minimize the risk of air embolism or hemorrhage. If catheter introduction is delayed, temporarily cover valve opening with sterile-gloved finger until SLIC® assembly is inserted. Use Arrow SLIC® assembly, included with this product, as dummy catheter with hemostasis valve/side port assembly and sheath. Refer to detailed SLIC® instructions on page 5 and 6. This will ensure that leakage does not occur and inner seal is protected from contamination.

20. Hold catheter in place and reposition catheter contamination shield so that distal hub is approximately five inches (12.7 cm) from hemostasis valve/side port assembly (refer to Fig. 6).

21. Hold rear hub (Tuohy-Borst adapter end) of catheter contamination shield in place. Disengage distal hub from inner feed tube by moving forward. Advance distal hub for-
ward toward hemostasis valve/side port assembly. Hold assembly in place (refer to Fig. 7).

Fig. 7

22. Press distal hub of catheter contamination shield over assembly cap. Twist to lock (refer to Fig. 8).

Fig. 8

- Orient slot in hub with locking pin on assembly cap.
- Slide hub forward over cap and twist.

23. Grasp catheter through front portion of catheter contamination shield and hold in place while repositioning Tuohy-Borst adapter end as desired (refer to Fig. 9). Precaution: Do not reposition Tuohy-Borst adapter end on catheter once moved to this final position.

Fig. 9

24. Tighten Tuohy-Borst adapter by pressing down on cap and simultaneously turning clockwise to secure hub to catheter. Gently pull catheter to verify securement. Precaution: Do not overtighten Tuohy-Borst adapter to minimize the risk of lumen constriction or catheter damage.

25. Tuohy-Borst adapter end of catheter contamination shield should be secured with sterile tape to inhibit catheter movement (refer to Fig. 10). Precaution: Do not apply tape to the transparent sheathing on the shield to avoid tearing material.

Fig. 10

26. Use suture tab to secure sheath and/or anchor with a purse string suture around the sheath suture ring. Precaution: Do not suture directly to the outside diameter of the sheath to avoid cutting or damaging the sheath or impeding flow.

27. Dress puncture site per hospital protocol. Precaution: Maintain the insertion site with regular, meticulous redressing using aseptic technique.

28. Record the insertion procedure on the patient’s chart.

Single-Lumen Infusion Catheter (SLIC®)
A Suggested Procedure:
Use sterile technique.

1. Precaution: Place the patient in slight Trendelenburg position, as tolerated, to reduce the risk of air embolism. If femoral approach is used, place patient in supine position.

2. Warning: Hemostasis valve must be occluded at all times to minimize the risk of air embolism or hemorrhage. If catheter is removed from sheath or catheter insertion is delayed, temporarily cover valve opening with sterile-gloved finger followed by insertion of SLIC® assembly. Twist to lock (refer to Fig. 11).
• Orient slot in hub with locking pin on assembly cap.

• Slide hub forward over cap and twist.

Precaution: When SLIC® assembly is inserted after catheter removal, use an appropriate antiseptic to prep the hemostasis valve housing prior to inserting the SLIC®. Include the exposed portion of the valve on the top of the housing.

The SLIC®, with the obturator in place, occludes the hemostasis valve minimizing the risk of air entry or blood loss through the valve. Warning: Connection between SLIC® obturator and SLIC® must be tightened securely and routinely examined to minimize the risk of disconnection and possible air embolism, hemorrhage or exsanguination.

3. To use the SLIC® for intravenous infusion, remove the blue-capped SLIC® obturator by twisting counterclockwise. Hold the infusion port to maintain positive lock to hemostasis valve housing. Pull the SLIC® obturator from the infusion catheter (refer to Fig. 12). Immediately attach desired line to Luer-Lock hub. Warning: Exposure of the central vein to atmospheric pressure may result in entry of air into the central venous system. Inspect the SLIC® obturator to ensure the entire length has been withdrawn. Document the SLIC® obturator withdrawal and start of infusion.

4. If the infusion through the SLIC® is discontinued, the hub should be capped with a Luer-Lock injection cap and handled per hospital flushing protocol, or the SLIC® should be withdrawn and replaced with a sterile Arrow obturator, sold separately, to ensure that leakage does not occur and inner seal is protected from contamination. Warning: Cover the lumen during any manipulation to minimize the risk of blood loss or the introduction of air into the sheath.

Catheter Removal Procedure:
1. Precaution: Place the patient in a supine position.

2. Remove dressing, if applicable. Precaution: To avoid cutting of the sheath, do not use scissors to remove the dressing.

3. Twist distal hub of catheter contamination shield to allow removal from locking pin on hemostasis valve/side port assembly. Withdraw catheter from sheath. Warning: Hemostasis valve must be occluded at all times to minimize the risk of air embolism or hemorrhage. Temporarily cover valve opening with sterile-gloved finger until catheter or obturator is inserted.

Sheath Removal Procedure:
1. Precaution: Place the patient in a supine position.

2. Remove dressing, if applicable. Precaution: To avoid cutting of the sheath, do not use scissors to remove the dressing.

3. If applicable, remove sutures from sheath. Precaution: Be careful not to cut the sheath.

4. Withdraw device from sheath. Cover hemostasis valve with sterile-gloved finger. Warning: Hemostasis valve must be occluded at all times to minimize the risk of air embolism or hemorrhage.

5. Warning: Exposure of the central vein to atmospheric pressure may result in entry of air into the central venous system. Remove sheath slowly, pulling it parallel to the skin. As sheath exits the site, apply pressure with a dressing impermeable to air, e.g., vaseline gauze. Because the residual sheath track remains an air entry point until completely sealed, the occlusive dressing should remain in place for at least 24-72 hours dependent upon the amount of time the sheath was indwelling.10,14,15,17.
6. Upon removal of the sheath, inspect it to make sure that the entire length has been withdrawn.

7. Verify that the sheath was intact upon removal.


References:


Arrow International, Inc. recommends that the user be acquainted with the reference literature. *If you have any questions or would like additional reference information, please contact Arrow International, Inc.*