INDICATION FOR USE
The Amplatzer™ Steerable Delivery Sheath is indicated to facilitate the delivery of the Amplatzer™ Amulet™ Left Atrial Appendage Occluder.

CONTRAINDICATIONS
The Amplatzer™ Steerable Delivery Sheath is contraindicated for patients with the presence of intracardiac thrombus with active infections including endocarditis. The Amplatzer™ Steerable Delivery Sheath is contraindicated for use with a power injection system.

WARNINGS
This device was sterilized with ethylene oxide and is for single use only. Do not reuse or resterilize this device. Attempts to resterilize this device can cause a malfunction, insufficient sterilization, or harm to the patient. Do not use this device if the sterile package is open or damaged. Inspect all components before use. Do not use if the package or items appear to be damaged or defective. This device should be used only by physicians who are trained in standard transcatheter techniques. The physician should determine which patients are candidates for procedures that use this device. Slowly remove the dilator and sheath from the patient to prevent an ingress of air. As this medical device uses an alternative small-bore connector (luer) design different from those of the ISO 80369 series, there is a possibility that a misconnection can occur between this medical device and a medical device using a different alternative small-bore connector, which can result in a hazardous situation causing harm to the patient. Special measures need be taken by the user to mitigate these reasonable, foreseeable risks. See call-out 6 in Figure 2 for an illustration of the small-bore connector (luer).

PRECAUTIONS
To prevent air ingress, maintain a continuous flush of saline to the flushing port throughout the procedure: The hemostasis valve in the closed position minimizes blood loss, but does not prevent air ingress. Capping of the bypass hub may not be sufficient to prevent air ingress. Do not aspirate from the sheath flushing port. DO NOT use the Amplatzer™ Steerable Delivery Sheath after the Use-by date stated on the package label. The physician should exercise clinical judgment in situations that involve the use of anticoagulants or antiplatelet drugs before, during, or after the use of the Amplatzer™ Steerable Delivery Sheath. Use caution when advancing the steerable delivery sheath and dilator to avoid damaging tissue and vessels or interfering with previously implanted medical devices. If resistance is felt, determine the cause before proceeding. Continuing to advance, retract, or manipulate the catheter under this circumstance has the potential to result in patient injury or delivery sheath damage.

POTENTIAL ADVERSE EVENTS
Potential adverse events that can occur during or after a procedure using this sheath may include but are not limited to: air embolism, allergic reaction/toxic effects (due to anesthesia, contrast media, etc.), arrhythmia, arteriovenous fistulae, bleeding, cardiac tamponade, cardiac perforation, death, dissection, embolism, foreign material embolous, heart failure, hematoma, hypotension/hypertension, infection, myocardial infarction/ischemia, pericardial effusion, peripheral embolism, pseudoaneurysm, stroke, thrombosis, transient ischemic attack, vascular access site injury (such as arteriovenous fistulae, hematoma, pseudoaneurysm), vessel trauma/damage.