Transjugular mitral valve repair with the MitraClip: A step-by-step guide

INTRODUCTION

- This report describes MitraClip™ device (Abbott) implantation via the transjugular approach in a patient with tortuous inferior vena cava (IVC) and prior mitral valve annuloplasty.

METHODS

- Initial attempts to engage the septum from the right or left femoral veins failed despite using a large introducer, as well as various fixed and steerable catheters and needles.
- Computed tomography confirmed extreme tortuosity in the iliofemoral veins but not internal jugular vein (IJ).
- Right IJ access was evaluated at the bench and accessed under ultrasound guidance.

Transseptal puncture

- SupraCross™ RF Wire was advanced into the IVC through the SupraCross™ Steerable Sheath (Baylis Medical*).
- Dilator was extended just 1 cm beyond the tip of the sheath.
- Transseptal system was rotated to 8 o’clock, retracted, and flexed to engage and tent the septum in the mid-superior position.
  - Note: Transjugular approach does not have the same requirements for transseptal height as from a femoral approach.
- Tenting was optimized by further exposing the dilator (Figure 1) and confirmed on transesophageal echocardiography.
- Flex and forward advancement were applied to the SupraCross™ Sheath and Wire assembly during RF transseptal puncture.
- The RF wire assumed a pigtail confirmation immediately upon entry in the left atrium (LA).

MitraClip™ procedure

- ProTrack™ Pigtail Wire (Baylis Medical*) was used to introduce the MitraClip™ Steerable Guide Catheter.
- MitraClip™ delivery system was miskeyed at 90° counterclockwise and advanced into the LA under-straddled to navigate toward the desired position.
  - M Knob resulted in medial and anterior deflection
  - Rotating counterclockwise resulted in posterior positioning to counteract anterior deflection of M Knob
  - Anterior Knob resulted in both anterior and lateral deflection, while the Posterior Knob resulted in both posterior and lateral deflection

DISCUSSION AND CONCLUSIONS

- Right IJ access is a reasonable alternative to performing MitraClip™ device procedures in the absence of femoral access.
- SupraCross™ Steerable Sheath and dilator, respectively, allowed adjustment of the sheath trajectory and tenting of the septum from the transjugular route.
- The RF wire used for transseptal puncture provided further support for advancing the steerable sheath across the septum.
- Intentional miskeyed insertion and under-straddle improved steerability and co-axial positioning of the MitraClip™ system within the LA from the IJ approach.

Figure 1. SupraCross™ Steerable Sheath was flexed and maneuvered to engage the septum using internal jugular vein access. Further optimization of tenting was possible by extending the dilator (Figure adapted from Yap et al, 2020).
Brief Summary | SupraCross™ Steerable Sheath

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete “Instructions for Use” for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator’s Instructions.

INDICATIONS FOR USE: The SupraCross™ Steerable Sheath kit is indicated for introducing various cardiovascular catheters to the heart, including the left side of the heart through the interatrial septum.

CONTRAINDICATIONS: There are no known contraindications for this device.

WARNINGS: • Laboratory staff and patients can undergo significant x-ray exposure during an interventional procedure 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 SupraCross™ Steerable Sheath kit is intended for single patient use only. Do not attempt to sterilize and reuse the SupraCross™ Steerable Guiding Sheath kit. Reuse can cause the patient injury and/or the communication of infectious disease(s) from one patient to another. • Care should be taken to ensure that all air is removed from the sheath before infusing through the side port. • Do not attempt direct percutaneous insertion of the sheath without the dilator as this may cause dilator injury. DO NOT attempt to insert or retract the guidewire through a metal cannula or a percutaneous needle, which may damage the guidewire and may cause patient injury.

PRECAUTIONS: • Careful manipulation must be performed to avoid cardiac damage, or tamponade. Sheath, dilator and guidewire advancement should be done under fluoroscopic guidance. If resistance is encountered, DO NOT use excessive force to advance or withdraw the device. • Avoid deflecting distal end of sheath during delivery and removal, otherwise damage to vessels may occur. • The SupraCross™ Steerable Sheath kit is not compatible with transseptal needles such as the “NER™ Transseptal Needle”. • Do not reshape the skid tip or curve of the guidewire. Excessive bending or kinking of the distal curve may damage the integrity of the wire or coating and lead to patient injury.

ADVERSE EVENTS: • Adverse events that may occur while using the SupraCross™ Steerable Sheath include: infection, air embolism, local nerve damage, vasovagal reaction, dissection, vessel spasm, all fistula formation, atrial septal defect, pseudoaneurysm, aortic puncture, arrhythmias, perforation and/or tamponade, hematoma, hemorrhage, catheter entanglement, embolic events, stroke, valve damage, myocardial infarction, paramagnet/debrillator lead displacement, pulmonary edema, coronary artery spasm and/or damage, vessel trauma, pericardial effusion

Brief Summary | SupraCross™ RF Wire

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete “Instructions for Use” for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator’s Instructions.

INDICATIONS FOR USE: The SupraCross™ RF Wire is indicated for creation of an atrial septal defect in the heart.

CONTRAINDICATIONS: The SupraCross™ RF Wire is not recommended for use with any conditions that do not require the creation of an atrial septal defect. The Connector Cable is not recommended for use with any other RF Generator or any other device.

WARNINGS: • Laboratory staff and patients can undergo significant x-ray exposure during RF puncture 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 SupraCross™ RF Wire and Connector Cable are intended for single patient use only. Do not attempt to sterilize and reuse either device. Use of the device, if not disposed of properly, may affect the patient’s treatment. • Reuse can cause patient injury and/or the communication of infectious disease(s) from one patient to another. • Reuse may result in patient complications. • The SupraCross™ RF Wire must be used with the Connector Cable provided. Attempts to use it with other connector cables can result in electrocution of the patient and/or operator. • Do not use the SupraCross™ RF Wire with electrocautery or electrocautery generators, connector cables or accessories as attempted use can result in patient and/or operator injury. • The Connector Cable must only be used with the RFP-100A Baylis RF Generator and the included SupraCross™ RF Wire. Attempts to use it with other RF Generators and devices can result in electrocution of the patient and/or operator. • The SupraCross™ RF Wire must be used with the 0.035” compatible transseptal sheath and/or dilator device. Use of incompatible accessory devices may damage the integrity of the SupraCross™ RF Wire or accessory devices and may cause patient injury. • The SupraCross™ RF Wire has only been validated for transseptal puncture use through SupraCross™ dilators which have been demonstrated to provide the required support for optimal function. • The SupraCross™ RF Wire is not intended for use with neonatal patients (i.e. less than one month of age). Do not attempt to treat neonatal patients with the SupraCross™ RF Wire.

PRECAUTIONS: • In order to prevent the risk of ignition, ensure that flammable materials are not present in the room during RF power application. • Do not bend the SupraCross™ RF Wire or the Connector Cable. Excessive bending or kinking of the wire shaft, distal curve of the wire and/or the Connector Cable may damage the integrity of the device components and may cause patient injury. Care must be taken when handling the SupraCross™ RF Wire and Connector Cable. • Careful manipulation of the SupraCross™ RF Wire must be performed to avoid vessel trauma. If resistance is encountered, DO NOT use excessive force to advance or withdraw the SupraCross™ RF Wire or ancillary sheath and/or dilator assembly. Excessive force may lead to bending or kinking of the device limiting advancement and retraction of sheath and/or dilator device. • Avoid RF energy delivery of the SupraCross™ RF Wire with incompatible dilator or cannula devices, which may lead to patient burns, ineffective puncture or failure to puncture. • The Baylis RF Generator is capable of delivering significant electrical power. Patient or operator injury can result from improper handling of the SupraCross™ RF Wire and/or DIP electrode, particularly when operating the device. • During power delivery, the patient should not be allowed to come in contact with ground metal surfaces.

ADVERSE EVENTS: • Adverse events that may occur while creating an atrial septal defect include: tamponade, sepsis, infection, thromboembolic episodes, vessel perforation, atrial fibrillation, myocardial infarction, vessel spasm, sustained arrhythmia, atrial flutter, hemorrhage, vascular thrombosis, perforation of the myocardium, hematoma, allergic reactions to contrast medium, ventricular tachycardia, pain and tenderness, arteriovenous fistula, pericardial effusion, tachycardia, vascular trauma, additional surgical procedure, wire entanglement, entrapment, foreign body/wire fracture

Brief Summary | ProTrack™ Pigtail Wire

CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. Rx only. Prior to use, please see the complete “Instructions for Use” for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator’s Instructions.

INDICATIONS FOR USE: The ProTrack™ Pigtail Wires are intended for use in percutaneous transseptal procedures to introduce and position catheters and other interventional devices within the left heart. The device is not intended for use in the coronary arteries.

CONTRAINDICATIONS: There are no known contraindications for this device.

WARNINGS: • DO NOT push, tug or withdraw the pigtail wire. Reuse can cause patient injury against until the cause of the resistance has been determined. Applying excessive force against unresistant resistance may cause damage to the pigtail wire, interventional device and/or vessel/or gan. • When the pigtail wire is exposed to the vascular system, it should be manipulated while under high-resolution imaging guidance including fluoroscopy and/or echocardiography. Improper visualization of the guidewire may lead to misplacement, dissection, or perforation. • Inspect the pigtail wire prior to use for coil separation, kinking, appropriate tip flexibility or breakage. If the pigtail wire is damaged or defective, do not use it. • Using a damaged or deflected pigtail wire may cause vasculature damage and/or compromise pigtail wire performance. • Laboratory staff and patients can undergo significant x-ray exposure during interventional procedures due to the continuous usage of fluoroscopic imaging. The 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.

ADVERSE EVENTS: • Potential complications associated with the use of the pigtail wire include, but are not limited to: vessel perforation, dissection, trauma or damage, vessel spasm, hemorrhage, access site complications, hematoma, thromboembolism, allergic reaction, vascular complication, cardiac tamponade, cardiac perforation, laceration, constriction disorder, embolism, additional surgical procedure, pericardial/pleural effusion, sepsis, infection, inflammation, foreign body/wire fracture, hemolysis, hypovolemia, myocardial ischemia and/or infarction, stroke, transient ischemic attack, vessel occlusion, wire entanglement, embolization, valve complication

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