The coronary Stent-On-A-Wire (SOAW)

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Background

Fixed wire angioplasty systems1,2 were introduced in the era of balloon angioplasty as these systems had a lower profile than the bulky, stiff, over-the-wire systems in use at the time. The fixed wire system allowed treatment of more complex lesions. As stenting procedures evolved in the mid 1990s, balloon catheters became mostly relegated to the pre- and post-dilatation of deployed stents which accelerated the use of over-the-wire devices. During this period, stent design and materials had not advanced sufficiently to provide low enough crossing profiles for direct stenting. The need to re-cross a lesion to permit further treatment with a stent and another balloon catheter represented a significant limitation of the fixed-wire balloon system. Reports of wire fractures and difficulties with balloon deflations with one of the manufactures systems (USCI Probe) may also have dampened enthusiasm for these early fixed-wire systems3-5. These early limitations of the technology, along with the development of lower profile over-the-wire systems, resulted in the limited use of the fixed-wire balloon catheter in recent years.

The Svelte stent-on-a-wire (SOAW) system represents the next generation of fixed wire technology and is unique in that it has a direct stenting indication by design. A direct stenting strategy offers the possibility of further reducing the risk of adverse events in high risk percutaneous coronary intervention (PCI) patients, decreasing radiation dose to the operator and patient, and decreasing procedural time and cost.6-10

Device description

The Svelte SOAW System is a high-performance coronary stent delivery system, consisting of a balloon expandable Svelte Stent pre-mounted on the Svelte SOAW single lumen fixed-wire delivery catheter platform (Figure 1). The Svelte Stent is a thin, flexible, bare-metal, L605 cobalt chromium coronary stent. The SOAW delivery catheter has a working length of 145 cm; the device includes a proximal stainless steel shaft with two proximal shaft markers (90 cm and 100 cm) to indicate the relative position of the delivery system to the end of a brachial or femoral guiding catheter; a flexible distal shaft, a nylon balloon with proximal and distal radiopaque markers located underneath to indicate the working length of the balloon and the expanded stent under fluoroscopy; a spring coil tip, proximal and distal Balloon Control Bands (BCBs) at each end of the balloon to protect leading stent edges from damage during delivery, as well as control balloon expansion and deflation; a standard luer fitting, and an integrated torque device located on the proximal shaft (Figure 1).

The SOAW System devices are compatible with 5 Fr guide catheters. The 0.74 mm (0.029") profile, leading tip diameter of 0.30 mm (0.012"), and excellent 1.1:1 torque response allows delivery to distal and tortuous anatomy. The stent reaches nominal size and is deployed at a balloon inflation pressure of 10-11 atm with a rated burst pressure at 18 atm. The SOAW System will be available in 25 models with diameters of 2.5 mm, 2.75 mm,
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3.0 mm, 3.5 mm, and 4.0 mm and lengths of 8 mm, 13 mm, 18 mm, 23 mm, and 28 mm.

Technical specifications

Material: L-605 cobalt chromium alloy
Stent lengths: 8 mm – 28 mm
Balloon diameters: 2.50 to 4.0 mm.
Guide catheter compatibility: 5 Fr (minimum ID 0.056")
Lesion entry profile: 0.30 mm (0.012")
Crimped stent profile: 0.74 mm (0.029")
Working catheter length: 145 cm
Shaft markers: 90 cm and 100 cm
Balloon: Proximal and distal radio-opaque markers; control bands at each end
Strut thickness: 84 μm
Length of the wire distal to the stent: 22 mm

Indications for use

- Elective direct stenting of a single stenotic lesion in a native coronary artery.
- Reference vessel ≥ 2.5 mm and ≤ 4.0 mm in diameter.
- Target lesion < 25 mm in length (the intention should be to cover the entire lesion with one stent of adequate length).
- Target lesion stenosis ≥ 50% and < 100%.
- Ability to cross the lesion with a wire.

The main exclusion criteria for the use of this device are:

- Lesion location that is aorto-ostial or within 5 mm of the origin of the LAD or LCX.
- Target lesion with side branches > 2.0 mm in diameter.
- Target lesion severely calcified.
- Target lesions that are likely to prevent complete inflation of an angioplasty balloon or proper placement of the stent or stent delivery system.

Tips and tricks for delivery

It is important to ensure the system is disconnected from the inflation device as the stent is being tracked through the vessel and across the lesion. If the inflation device is attached, it will negatively impact the torque response of the device as it will act as an anchor when steering into the vessel. Once the stent is in position, the inflation device can be attached and the stent can be deployed according to the instructions for use.

It is important to torque the device and get a feel for the torque response when it exits the guiding catheter. Therefore when the wire tip exits the guide, the catheter shaft can be rotated in order to confirm the appropriate rotation of the catheter tip.

If the tip of the device is not rotating, one should make sure the torquer is tight on the shaft of the SOAW system. The device can also be rotated at the hub to check the torque control of the tip.

If it is needed to cross the deployed stent with a guidewire, it is best to prolapse the tip of the wire prior to crossing the stent to ensure the wire goes through the centre of the stent and not between the struts.

When the stent has been fully deployed, it is recommended to push the balloon just distal to the lesion and then make a final angiogram to check result. By doing so, re-crossing the stent if needed with another device is simplified. It is also possible to post-dilate the lesion with the SOAW if required.

Pre-clinical studies and initial clinical experience

The Svelte SOAW System has been evaluated against the MultiLink Vision (Abbott Vascular, Santa Clara, CA, USA) system in both non-good laboratory practice (GLP) and GLP pre-clinical assessments. These studies relied on porcine models, specifically Yorkshire pigs, with no gender bias. The Svelte SOAW performed equivalently to the MultiLink Vision stent with respect to deliverability and ease of use.

Static x-ray images of implanted stents demonstrate the similarity between the Svelte and MultiLink Vision stents, as can be seen in the Figure 2. These images were captured by Svelte Medical Systems, Inc. after the stents were implanted in coronary arteries of porcine models. The imaging system is a Faxitron #43855B, (Faxitron X-Ray LLC, Lincolnshire, IL, USA) and the images were captured through a 45 s exposure at 45 kV. (Figure 2)

In a recently completed GLP study, eight juvenile Yorkshire swine (30 day cohort) and seven Yucatan mini-swine (90 day cohort) were implanted with either Svelte SOAW or control MultiLink Vision balloon expandable stents. Stented vessels were examined for standard histomorphometric and histopathologic measures of tissue response. Scanning electron microscopy (SEM) was also conducted to assess endothelialisation.
Histopathological assessment demonstrated minimal injury and inflammation at both 30 and 90 days with Svelte SOAW and MultiLink Vision stents. Endothelialisation was complete by 30 days for both stent treatments as was neointimal maturation (Figure 3). Histomorphometry demonstrated no differences between Svelte SOAW and MultiLink Vision stents in neointimal growth, lumen area, percent occlusion, or wall area ratios at either 30 or 90 days post implant (Figure 4). SEM analysis of both stents showed 100% confluent mixture of polygonal and spindle shaped endothelial cells with minimal inter-cellular spacing at 30 days and cobblestone-like polygonal endothelial cells with tight junctions by 90 days (Figure 5).

Based on the animal studies findings, a prospective, single arm, first-in-man, multicentre study is ongoing at Thoraxcenter Erasmus MC, Rotterdam, The Netherlands. The first-in-man implantation of the device has recently been performed in a patient with a mid-left
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Circumflex coronary artery lesion. The 3-D quantitative coronary angiogram (QCA) pre- and post-stenting, and the 3-D optical coherence tomography (OCT) post-stenting are shown in Figure 6 (A, B) and Figure 7. The 3-D angiographic reconstruction with virtual navigation through the centreline of the vessel and the coronary angiogram showing the positioning of the device through the lesion is displayed on www.eurointervention.com.

Conclusions

The Svelte SOAW system represents the next generation iteration of fixed wire technology, and is unique in that it has a direct stenting indication by design.

Figure 5. Scanning electron microscopy (SEM) of vessels implanted with either Svelte SOAW or ML Vision stents showing 100% confluent mixture of polygonal and spindle shaped endothelial cells with minimal inter-cellular spacing at 30 days and cobblestone-like polygonal endothelial cells with tight junctions by 90 days.

Figure 6. A) 3-D quantitative coronary angiogram, QCA pre- and post-stenting. B) 3-D QCA post-stenting.

Figure 7. 3-D optical coherence tomography (OCT) post-stenting showing well apposed Svelte stent struts.

Figure 6. A) 3-D quantitative coronary angiogram, QCA pre-stenting. B) 3-D QCA post-stenting.
It may enable a further reduction in procedure time, contrast use and cost compared to conventional devices. Results from a GLP study demonstrate an acceptable healing response when the Svelte SOAW is implanted in porcine coronary arteries with complete endothelialisation by 30 days that was no different than that observed with the MultiLink Vision stent. The potential downsizing of sheath sizes and improved outcomes with direct stenting may make this approach appealing.

References