Novel Coronary Stent Platforms

Bon-Kwon Koo, MD1,2 and Peter J. Fitzgerald, MD2
1Department of Internal Medicine, Seoul National University College of Medicine, Cardiovascular Center and Cardiovascular Research Institute, Seoul National University Hospital, Seoul, Korea, 2Stanford University Medical Center, Stanford, California, USA

ABSTRACT

The stent has been a key part of percutaneous coronary intervention. The advent of drug-eluting stents has further expanded the indication for this technology with a lower overall rate of restenosis. However, the limitations of the current generation stent platforms have become more apparent as more complex are being treated with percutaneous coronary intervention. Coronary stenting sometimes results in a suboptimal outcome for challenging lesion subsets such as tortuous, calcified, bifurcating and multiple and long lesions. In this review, novel stent systems that have been developed to overcome the challenges surrounding current stent designs will be discussed. (Korean Circ J 2008;38:393-397)

KEY WORDS: Coronary artery stenosis; Angioplasty, transluminal, percutaneous, coronary; Stents; Drug-eluting stents.

Introduction

Through the marked reduction of in-stent restenosis, the use of drug-eluting stents has definitely expanded the field of percutaneous coronary intervention. However, as more and more complex lesions are being treated with drug-eluting stents, the limitations of the current stent designs have become more apparent. Complex lesion characteristics challenges stent delivery and the designs of the current stents are not always suitable for the complex anatomy, such as bifurcation lesions. Furthermore, the use of multiple stenting results in increased costs for both the patients and healthcare providers. In this review, recently developed novel stent systems that have been designed to overcome these limitations will be described.

Improvement in Deliverability

Despite the recent progress in coronary stents and delivery system, a failure to cross the stent through calcified, tortuous or stented segments is still one of the main causes of procedural failure. A novel stent and delivery system named Sparrow™ (CardioMind, Sunnyvale, CA, USA) consists of a baremetal, self-expanding nitinol stent incorporated into an assembly miniaturized to a 0.014-inch diameter guidewire platform. As this system has a guidewire-level crossing profile and excellent flexibility (Fig. 1), it allows stenting for complex lesions that are inaccessible with current stent systems. A feasibility study (CARE 1 trial, n=22) in patients with small vessel disease showed no six-month major adverse cardiac events with an angiographic late loss of $0.73 \pm 0.57$ mm. The drug-eluting version of this stent system has been also developed and is currently under clinical evaluation.

In Situ Stent Customization for Long/Multiple Lesions

Treatment of long or multiple lesions using currently available fixed-length stent systems requires several devices and a separate exchange for each stent. This procedure increases the complexity, time and cost of stent usage. The Custom NX® DES system (XTENT® Inc., Menlo Park, CA, USA) is composed of multiple inter-digitated cobalt chromium stent modules up to 60 mm, each 6 mm in length (Fig. 2). This stent has a biodegradable polymer that releases Biolimus A9, a sirolimus analog. An operator can select the appropriate stent length in situ and the deployment of multiple stents is allowed with a single delivery device. In addition, this
system also allows the operator to shorten the balloon length to perform higher pressure post-dilatation within the stented segment. Therefore, long and multiple lesions of variable lengths and diameters, in one or more arteries, can be treated with a single device. The two-year follow-up results of a feasibility study (CUSTOM I trial, n = 30) showed no major adverse cardiac event with an eight-month angiographic late loss of 0.26 ± 0.23 mm.9) With the favorable results of the CUSTOM I-III trials, the pivotal clinical trial for this stent will soon be started.

Dedicated Stents for Bifurcation Lesions

Percutaneous coronary intervention for bifurcation lesions is technically difficult due to the anatomical variability and complexity of the lesions.10) Bifurcation intervention is associated with more procedural complexity, a lower procedural success rate and a higher clinical event rate.11-13)

As the overall design of stents has been optimized for straight, non-bifurcating coronary artery lesions, dedicated stents for bifurcation lesions have been developed. In general, these stents can be divided into three categories: Y-type complete bifurcation stents, stents with side branch access and side branch stents. As there has been limited progress in the development of Y-type stents, novel stents of the other categories will be described.

Stents With Side Branch Access

Side branch accessible main branch stents are designed to preserve access to a side branch during a procedure and to provide adequate scaffolding of the side branch ostium. In most stents, the aperture is located at the center of a stent and the proximal part of side branch balloon is mounted within the main branch stent (Multilink Frontier™, Abbott Vascular, Santa Clara, CA, USA; Twin Rail™, Invatec, Roncadelle, Italy; Nile Croco, Minvasys, Genevilliers, France). In the FRONTIER Stent Registry (n = 105), the procedural success rate of the Multilink Frontier™ stent (Fig. 3) was 93% and the six-month target lesion revascularization rate was 13.3%.14) A drug-eluting version of this stent will soon be available.

The TAXUS Petal™ bifurcation stent (Boston Scientific, Natick, MA, USA) has a unique petal structure in the middle of the stent to provide consistent mechanical support to the side branch ostium (Fig. 3). This stent system has the same anti-proliferative drug and polymer as the TAXUS stent. The TAXUS PETAL I First Human Use trial has recently begun.

The Antares™ Sidebranch Adaptive System (Trireme Medical, Pleasanton, CA, USA) is also designed to faci-
litate side branch access and maximize side branch ostial scaffolding. It is a balloon-expandable stainless steel stent with a unique side branch ostial preservation structure in the center of the stent. This ostial preservation structure opens into the side branch ostium and adapts to the asymmetric anatomy of the side branch with variable angles (Fig. 4). A side branch stabilizing wire is encased in a wire management system and it allows direct side branch wiring and access after stent deployment into the parent vessel, and thus eliminates the need for stent strut re-crossing. The first human clinical trial of this stent is ongoing.

The Stentys™ coronary bifurcation stent (Stentys, SAS, Paris, France) is a self-expanding, drug-eluting stent designed to treat Y-type bifurcation lesions with various calibrations. Unlike most other dedicated bifurcation stents, this stent does not require precise positioning of the side branch segment. As the Z-shaped mesh of the stent is linked by small interconnections, the struts can be disconnected using an angioplasty balloon to create side branch access and to achieve side branch ostial scaffolding (Fig. 5). Therefore, any strut can be selected as a side branch opening after main branch stent implantation.

**Side Branch Stents**

The Tryton Side-Branch Stent™ (Tryton Medical, Newton, MA, USA) has a unique design to cover the side branch completely in all bifurcation lesions with different angulations. This stent is a cobalt chromium stent composed of three different zones—side branch region with a standard stent design, a transition zone and the main vessel region (Fig. 6). In the first human study (Tryton I, n=30), the six-month major adverse cardiac events rate was 9.9% with an angiographic late luminal loss of the side branch of 0.17±0.35 mm. The procedural success rate was 93.3%. In this trial, the side branch was treated first with the Tryton stent and then a standard drug-eluting stent was subsequently implanted in the main branch. The Sideguard™ Ostium protection device™ (Cappella, Auburndale, FL, USA) is a self-expanding stent with a pre-formed, trumpet-shaped nitinol frame (Fig. 6). This is an anatomically-designed side branch stent for complete ostial coverage. In the Sideguard I FIM study (n=15), the acute angiographic success rate of this device was 80%.

**Stents to Cover the Proximal Part of a Bifurcation Lesion**

The Axess™ stent (Devax, Irvine, CA, USA) consists of a self-expanding nickel titanium platform with a bioabsorbable polymer that releases Biolimus A9. The stent is designed to be placed in the proximal part of the main vessel, at the level of the carina. It has a straight and conical configuration to expand properly into the complex anatomy of a bifurcation lesion (Fig. 7).
The unique design and expansion of this device enhances the interaction between the adequate mechanical scaffolding and accurate delivery of antirestenosis drugs. After Axxess™ stent implantation, additional stents can be easily placed at the distal main or side branches. In the AXXESS Plus trial (n=139), which was the first human trial of the Axxess stent for non-left main bifurcation lesions, the six-month target lesion revascularization rate was 7.5%, with an angiographic late luminal loss of Axxess stent of $0.09 \pm 0.56$ mm. This stent system also showed favorable outcomes for left main bifurcation lesions (AXXENT trial).

**Conclusion**

Due to an increasing demand to treat more complex...
lesions with percutaneous coronary intervention, a number of novel stent designs has been developed. As opposed to the one-design-fits-all concept, the current developments are directed toward the achievement of a lesion specific approach. Most of these novel devices have shown promising results from feasibility studies. Larger randomized trials are being planned or are ongoing to confirm the safety and efficacy of the devices further. These next generation technologies will expand the field of intracoronary stenting and improve clinical outcomes for patients with coronary artery disease.

Acknowledgments

The CardioVascular Research Foundation, Korea (CVRF), supported this work.

REFERENCES