Shifting the SFA Treatment Paradigm

Sponsored by Boston Scientific Corporation

IMPERIAL 12-Month Full Cohort and Long Lesion Sub-Study Results

Sustained drug-release in the SFA drives superior results.

BY WILLIAM A. GRAY, MD | IMPERIAL PRINCIPAL INVESTIGATOR

Coronary drug-eluting stents (DES) with both paclitaxel and limus drugs have demonstrated successful long-term clinical outcomes for patients, with single-digit reintervention rates at 1 year. While the limus family of drugs have been particularly successful in coronary stenting, two major clinical trials studying limus-eluting stents in the superficial femoral artery (SFA) failed to show clinical efficacy. The SIROCCO study evaluated a sirolimus-eluting version of the SMART stent (Cordis, a Cardinal Health company),\(^1\) and the STRIDES study evaluated an everolimus-eluting version of the Dynalink stent (Abbott Vascular).\(^2\) Neither study was able to show a statistically significant difference between the limus-eluting stents and their respective bare-metal counterparts.

Conversely, paclitaxel, which works by inhibiting cell proliferation and migration, has demonstrated safety and efficacy in the coronary arteries as well as the SFA. The randomized controlled trial for the Zilver PTX DES (Cook Medical) showed an 18% difference in primary patency between the Zilver PTX arm versus the bare-metal stent (BMS) arm.\(^3\) Additionally, the RANGER-SFA, LEVANT, and IN.PACT trials demonstrated the efficacy of paclitaxel-coated balloons over PTA in the SFA.\(^4-6\)

RESTENOSIS IN THE SFA

The treatment of disease in the SFA presents a considerable challenge due to the unique mechanical forces in this vessel bed and the high degree of severe calcium and occlusions. Based on the clinical literature, smooth muscle cell proliferation can occur for up to 100 days or longer, and the final phase of restenosis can last for well beyond 1 year.\(^7\)

Unlike disease in the coronary arteries, where restenosis usually peaks within 3 to 6 months, restenosis tends to peak later in the SFAs, usually between 9 to 12 months. The experience with peripheral first-generation nitinol BMS was disappointing, with 1-year primary patency rates well below 80%, far less than patency rates observed in the coronary arteries.

THE ZILVER PTX DRUG-COATED STENT

The Zilver PTX stent is a paclitaxel drug-coated stent, which received FDA approval in 2012. This stent does not have a polymer or carrier, so the drug is simply applied to the stent. It has a 3 \(\mu\)g/mm\(^2\) dose of paclitaxel, which upon implantation is immediately released during the first few days and remains in the vessel wall up to 56 days in preclinical testing.\(^8\) The Zilver PTX stent is well-studied and has demonstrated improved patency and target lesion revascularization (TLR) rates over its BMS counterpart, with a 12-month primary patency rate of 82.7% in the Zilver PTX randomized controlled trial.\(^3\) The Zilver PTX stent has a 5-year primary patency rate of 66.4% versus a percutaneous transluminal angioplasty (PTA) primary patency rate of 43.4%, demonstrating a durable patency effect with antiproliferative therapy that improves outcomes.\(^9\)

THE ELUVIA DRUG-ELUTING STENT

With the design of the Eluvia Drug-Eluting Stent System, Boston Scientific sought to improve upon existing clinical outcomes, targeting low single-digit reintervention rates and more durable long-term outcomes for patients. The Eluvia Drug-Eluting Stent represents a novel approach to the treatment of diseased femoropopliteal arteries as the first and only technology designed to sustain drug release beyond 1 year to match the restenotic process in the SFA (Figure 1). The stent platform is designed to withstand the mechanical forces of the SFA, balancing optimal strength and fracture resistance, while providing a uniform scaffolding for drug delivery. The polymer is a fluorinated polymer, which was intentionally designed to deliver optimized drug transfer with the lowest possible drug dose. The paclitaxel release is highly targeted to the lesion with virtually no drug lost downstream. Its safety has been studied in over 100,000 patients in clinical trials and implanted in over 20 million vessels commercially.\(^10\) The polymer allows the 0.167 \(\mu\)g/mm\(^2\) paclitaxel dose drug delivery to be tuned to sustain drug release beyond 1 year.

THE IMPERIAL CLINICAL TRIAL PROGRAM

The IMPERIAL trial represents the first head-to-head
trial comparing two antiproliferative stents in the SFA. The IMPERIAL randomized cohort is a global prospective single-blind multicenter randomized controlled trial comparing Boston Scientific’s Eluvia Drug-Eluting Vascular Stent to Cook Medical’s Zilver PTX drug-coated stent (2:1 randomization). The randomized cohort enrolled 465 patients across 64 sites around the world. Eligible patients had chronic, symptomatic lower limb ischemia and de novo or restenotic lesions up to 140 mm in length in the native SFA and/or proximal popliteal artery (PPA). As prespecified in the Statistical Analysis Plan, once both primary endpoints were met, a post-hoc superiority analysis could be performed. The IMPERIAL trial also included a single-arm 50-patient long lesion sub-study evaluating safety and effectiveness of Eluvia in lesions 140 to 190 mm in length (clinicaltrials.gov identifier NCT02574481).

The average lesion length in the IMPERIAL randomized cohort was 87 mm in the Eluvia arm and 82 mm in the Zilver PTX arm. In the Eluvia arm, 40% of the lesions were severely calcified, 31% were total occlusions, and 84% extended into the distal portion of the SFA and/or PPA. In the Zilver PTX arm, 32% of the lesions were severely calcified, 30% were total occlusions, and 78% extended into the distal portion of the SFA and/or proximal popliteal artery. There were no statistically significant differences in terms of patient or lesion characteristics between the two study arms.

**SUPERIOR RESULTS IN THE FIRST HEAD-TO-HEAD DES SFA TRIAL**

Eluvia demonstrated superiority in primary patency over Zilver PTX in the prespecified post-hoc analysis. The Kaplan-Meier estimated primary patency rates were 88.5% versus 79.5%, respectively ($P = .0119$). IMPERIAL reported a 4.9% major-adverse event rate in the Eluvia arm and a 9% rate in the Zilver PTX arm ($P = .0975$), most of which were comprised of TLR rates at 1 year in both arms. Patients in the Eluvia arm underwent half as many TLRs compared to those in the Zilver PTX arm (4.5% versus 9%, $P = .0672$). Both arms of the study reported strong patient outcomes, with 85.8% of the patients presenting with no or minimal claudication at 12 months in the Eluvia arm compared to 84.5% in the Zilver PTX arm, but at a cost of twice the reintervention rate in the Zilver PTX arm to achieve these outcomes. Baseline clinical improvement was sustained at 12 months in 89.6% of the Eluvia patients and 83.1% of the Zilver PTX patients.

**CONSISTENT RESULTS INDEPENDENT OF LESION LENGTH**

In the long lesion cohort of the IMPERIAL trial, Eluvia demonstrated an 87.9% primary patency rate in lesions with a mean length of 162.8 mm (Figure 2). This patient group also had lesion characteristics of 70% moderate/severe calcium, nearly a third total occlusions, and 76% extended into the distal portion of the SFA and/or PPA. Freedom from major adverse events was observed at 93.5% at 12 months with a TLR rate of 6.5% (Table 1). These results with Eluvia completely counter the notion that as lesion length and complexity increases, stent patency decreases. These data are consistent with the results from the smaller independent Münster registry, which observed an 87% primary patency rate at 12 months in a highly complex patient population with 80% chronic total occlusions, 48% critical limb ischemia, and an average lesion length of 200 mm.

The IMPERIAL trial is a landmark trial that advances the peripheral vascular space and provides the physician...
Shifting the SFA Treatment Paradigm

Sponsored by Boston Scientific Corporation

INSIGHTS ON THE DATA WITH AN IMPERIAL PRINCIPAL INVESTIGATOR

With Prof. Stefan Müller-Hülsbeck, MD, EBIR, FCIRSE, FICA, FSIR | IMPERIAL Co-Principal Investigator

As the lead Principal Investigator for the MAJESTIC trial and Co-Principal Investigator for IMPERIAL, what were your initial reactions to the IMPERIAL results?

Prof. Müller-Hülsbeck: The results were excellent. My expectations, which were based on the encouraging 1-year data from MAJESTIC, were met for primary patency and rate of freedom from TLR. I felt somewhat relieved that a larger study population achieved excellent data with a polymer-coated DES.

Why is it important for the endovascular community to have a head-to-head trial comparing two DES technologies?

Prof. Müller-Hülsbeck: Having a head-to-head comparison of two available devices is something new that the interventional community has been waiting for. Comparing an established DES like Zilver PTX with the new Eluvia DES technology is outstanding, because convincing 5-year Zilver PTX data have already been published. Having more robust data from a head-to-head comparison might strengthen the acceptance of this technology in general when there is a need for an implant such as a self-expanding stent or DES.

Eluvia demonstrated superiority over Zilver PTX in IMPERIAL. How should physicians performing endovascular procedures think about this superiority data when making device decisions?

Prof. Müller-Hülsbeck: Physicians must keep in mind that all trial data are collected under ideal "trial conditions," meaning dedicated inclusion and exclusion criteria need to be fulfilled before the study device is allowed to be used; all patients are under more controlled follow-up, including stricter drug regimens; and last but not least, all participating physicians are well trained to serve as investigators in the trial. That means data are obtained under optimized conditions, which strengthens the results reached.

However, statistical calculations for device safety and efficacy necessitate a minimum number of included subjects to draw any conclusions on the results, so the device performance is impressive. IMPERIAL showed that both available DES technologies provide good patient outcomes, but Eluvia performed better in terms of primary patency and 50% lower TLR rates. This might influence future decision making, meaning current Zilver PTX users may switch to Eluvia, and first-time potential users of DES technology will probably choose Eluvia from the beginning.

What are your thoughts on Eluvia's performance in IMPERIAL (88.5% primary patency), based on the very challenging lesion characteristics studied (40% severe calcium in the Eluvia arm and nearly a third chronic total occlusions)?

Prof. Müller-Hülsbeck: My preference in cases that need a stent is rather clear: implant a DES, ideally Eluvia. Doing the best for our patients is the goal. Since seeing the results of the first-in-human MAJESTIC trial, I believed that all lesions that need scaffolding should receive a DES such as Eluvia. Now seeing the results from IMPERIAL, this vision may come true. Calcium and chronic total occlusions shouldn't hinder us from using an Eluvia stent.

"Leave nothing behind" has become a popular saying in the endovascular space. However, Eluvia demonstrated a TLR rate of just 4.5% in IMPERIAL. Do you believe that outcome could cause some physicians to rethink a leave-nothing-behind strategy, when a DES provides such excellent TLR rates?

Prof. Müller-Hülsbeck: "Leave nothing behind" should still be a considered strategy, because a nitinol scaffold may not be appropriate for some patients. If a stentless strategy fails, we still have the option to implant a scaffold, or simply to repeat a leave-nothing-behind intervention. Stenting in all cases is not appropriate; however, many stents are still used because there is a strong need, and these cases should be privileged with a DES rather than a BMS. The promising results from IMPERIAL may liberalize stent usage, but this decision will ultimately be influenced by reimbursement, which varies from country to country.

How might the IMPERIAL data affect your SFA treatment algorithm moving forward?

Prof. Müller-Hülsbeck: I believe that if there is a need for an implant, all implants should be DES. This may be the end of the BMS era for femoropopliteal disease treatment.
DRUG-ELUTING THERAPIES IN PRACTICE

Antiproliferative therapies such as DESs and drug-coated balloons (DCBs) have demonstrated proven results in the SFA. We know they work. But how do we decide which therapy to use when and where? That question still needs to be answered. However, the patient populations studied in DCB trials represent primarily TASC A/B lesions, < 10 cm, and less calcification. Historical SFA stenting studies are typically inclusive of patient populations with more complex lesions and a high degree of calcification. Currently, interventionalists’ treatment algorithms typically start with vessel prep using a PTA balloon. If the result is optimal, then for shorter, simpler (TASC A and B) lesions, one might consider first-line therapy to be a DCB. If the result is suboptimal following PTA, a DES would be an appropriate treatment option.

The current available clinical data for long, highly calcified lesions seems to skew in favor of DES. Data from Dr. Fabrizio Fanelli demonstrated that the severity of calcification may impact DCB efficacy. As noted in the IMPERIAL long lesion sub-study results, Eluvia has demonstrated a 12-month patency of 87.9% in average lesion lengths of 162.8 mm and 70% moderate to severe calcium.

Today, technologies not only need to prove safety and efficacy, but total cost and overall value to the health care system must also be evaluated. With the progressive nature of peripheral artery disease, multiple reinterventions on the same patient becomes an expensive enterprise. Treating restenosis is often not an easy task, requiring multiple modalities depending on the location and nature of restenosis. Antiproliferative therapies greatly improve the reintervention rates not only at 12 months, but also provide durable results long-term. Balancing durable improved clinical outcomes with time spent in the lab during the procedure should be taken into consideration when evaluating the total value of a therapy. Mean procedure time for Eluvia was 57 minutes in the IMPERIAL trial. Eluvia has demonstrated consistent, reproducible results across multiple data sets in patients with claudication and critical limb ischemia, as well as short and long lesion lengths. A DES like Eluvia aims to provide patients with improved clinical outcomes and gives physicians an effective tool to help minimize costly reinterventions while keeping the procedure simple and efficient.

Table 1. Safety Results from IMPERIAL Long Lesion Sub-Study

<table>
<thead>
<tr>
<th></th>
<th>Eluvia (n=50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12-month MAE</td>
<td>6.5%</td>
</tr>
<tr>
<td>All Causes of Deaths at 1 Month</td>
<td>0.0%</td>
</tr>
<tr>
<td>Target Limb Major Amputation</td>
<td>0.0%</td>
</tr>
<tr>
<td>Target Lesion Revascularization</td>
<td>6.5%</td>
</tr>
<tr>
<td>Stent Thrombosis</td>
<td>0.0%</td>
</tr>
</tbody>
</table>

6.5% TLR in 162.8 mm Lesions