Saphenous vein graft intervention with a bioresorbable vascular scaffold: a follow-up optical coherence tomography study at 40 months

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Introduction
Bioresorbable vascular scaffolds (BVS)1-3 were designed to be an effective substitute to conventional metallic stents and overcome limitations including positive vessel remodelling, stent fracture, neoatherosclerosis, being a perpetual nidus for stent thrombosis and inhibition of physiological vasodilation4,5. The most widely studied polymeric devices comprise poly-L-lactide (PLLA) struts coated with poly-D, L-lactide (PDLLA) and an antiproliferative agent4,5. Although late outcomes with the Absorb™ bioresorbable vascular scaffold (Abbott Vascular, Santa Clara, CA, USA) are disappointing6, this only confirms the need to gain a better understanding of the late behaviour of these scaffolds. The literature so far suggests that there is some evidence supporting their use in venous grafts7-10. This case description illustrates the unique findings of optical coherence tomography (OCT) imaging of a BVS in a vein graft at 40 months post implantation.

Methods
A 77-year-old lady had previously undergone two bypass operations, the latter following recurrent restenoses of conventional metallic stents. Approximately seven years post second bypass (April 2013), she had symptoms (angina) with inducible ischaemia on a stress echocardiogram. Angiography found high-grade proximal disease in a saphenous venous graft to the left anterior descending artery. The lesion was adequately prepared using a 3.0 mm predilatation balloon (Tazuna® [Terumo Corp., Tokyo, Japan] – a semi-compliant percutaneous transluminal coronary angioplasty [PTCA] balloon catheter). A 3.5×28 mm Absorb BVS was deployed at 14 atmospheres and post-dilation was performed with a 3.75 mm non-compliant balloon (Hiryu® [Terumo Corp.] – a non-compliant PTCA balloon catheter) to 21 atmospheres. A distal protection device was utilised (3.0 mm SpiderFX™ [Medtronic, Minneapolis, MN, USA] embolic protection device) and the patient was placed on dual antiplatelet therapy for six months (aspirin 100 mg daily and clopidogrel 75 mg daily).

Results
Approximately 40 months post implant, she again developed symptoms (angina); an exercise test was abnormal and repeat coronary angiography was performed. This demonstrated new flow-limiting disease in a small to medium-sized native marginal circumflex artery, successfully treated with plain balloon

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angioplasty using a 2.0 mm predilatation balloon (Tazuna). Given concerns as to whether the diseased circumflex artery explained her presentation and the fact that the proximal segment of the vein graft to the left anterior descending artery appeared to have reduced volume (Figure 1), an OCT study (Moving image 1, Figure 2) was performed to assess the BVS in the vein graft to the left anterior descending artery. This demonstrated good apposition of the scaffold, tissue coverage with a neointimal depth of 0.48 mm and five uncovered struts. There were several struts manifesting various stages of PLLA degradation ranging from early cellular infiltration of dissolving struts (very high peak intensity at the core with low median intensity) to late degradation of the strut (moderately high peak intensity with a high median intensity). No intervention was performed on the vein graft. Post procedure, she was placed on dual antiplatelet therapy for six months (aspirin 100 mg daily and clopidogrel 75 mg daily) and has remained well since.

Discussion
In September 2017, Abbott Vascular ceased supplying BVS, citing low demand but more likely due to disappointing clinical results. They also signalled their intention to release a newer improved device for future clinical trials. Accepting that initial poor implantation technique and inappropriate vessel selection played a role, the four-year ABSORB II data showed a statistically higher rate of target lesion failure (TLF) (11.1% vs. 5.6%). There were no cases of stent/scaffold thrombosis between three and four years and some see this as the potential start of scaffold benefit. Data do support a “learning curve”: the three-year data of ABSORB III showed only a trend towards a higher rate of TLF in the ABSORB arm (13.4% vs. 10.4%) but with a persistently higher rate of device thrombosis (2.3% vs. 0.7%), whilst the 30-day ABSORB IV (using an optimal implantation technique) reported reduced early events (30-day scaffold thrombosis reduced from 1.1% in ABSORB III to 0.4% in ABSORB IV at 30 days) to the extent that ABSORB IV shows non-inferiority at 30 days. Given that a “learning curve” probably exists, cases such as this and any knowledge gained remain clinically relevant.

The development of a “vascular restoration” strategy in the form of BVS has been considered by many as the “fourth revolution” in the percutaneous management of coronary artery disease. So far, seven trials have been published comparing everolimus-eluting bioresorbable scaffolds and everolimus-eluting metallic stents. The benefit of bioresorbable stents is thought to pay off in the long term: at around twelve months there is a loss of mechanical support that allows conditioning of tissue to enhance healing, and at three years full reabsorption is noted, that assists in physiological

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**Figure 1.** Saphenous vein graft to the left anterior descending artery pre implantation, immediately post implantation of the BVS and at 40 months post implantation of the BVS. BVS: bioresorbable vascular scaffold

**Figure 2.** Optical coherence tomography image of the BVS.
function and neo-media formation in coronary arteries. Based on trial evidence at 60 months\(^1\), the BVS struts disappear completely.

Recently, Nakatani and colleagues\(^1\) performed an intravascular ultrasound echogenicity analysis and light intensity analysis on OCT in a porcine model with a follow-up duration of 48 months. There was good correlation observed for the strut depolymerisation process and integration process post comprehensive bioresorption. In addition, late luminal enlargement was observed at three to four years and was associated with strut integration.

Some bioresorbable stents are still commercially available; however, we are still awaiting the results of ABSORB IV, due for release in the next few years. Newer-generation stents such as the Magmaris\(^\text{TM}\) (Biotronik, Bülach, Switzerland) are showing promising results after publication of BIOSOLVE II and III\(^1\), in which improved stent and study design adds strength to the argument that BVS may still play a role in future percutaneous interventions.

**Limitations**

Little is known about the role of bioresorbable vascular scaffolds in venous grafts. The publications\(^7-10,20\) so far in this genre are summarised in Table 1. Indications for use in venous grafts are beyond the scope of this report, and the stent inserted was done so prior to release of ABSORB II and III. Our rationale for implantation in this high-risk patient was that she had incessant restenosis of metallic stents and was willing to allow sequential non-invasive imaging with computed tomography.

**Conclusion**

This report is unique given that it is the longest follow-up period to date of such an intervention with an Absorb scaffold in a saphenous vein graft. In an era when the use of BVS has been questioned, in this particular case the scaffold demonstrated excellent late angiographic patency confirmed on invasive OCT imaging. Meticulous attention to Absorb scaffold implantation technique is critical to providing good short-term as well as long-term clinical results\(^21,22\). Despite the Absorb BVS being recalled due to an increased incidence of target vessel myocardial infarctions\(^11\), there is still potential for benefit after scaffold resorption, and newer-generation BVS devices remain a viable option. Published data are still lacking to debunk its use entirely\(^6\).

**Impact on daily practice**

Our findings suggest that late OCT images of the scaffold treatment site are consistent with the existing chronology of events in intracoronary imaging of Absorb scaffolds in native coronary arteries. Further research and ideally a robust randomised trial comparing Absorb bioresorbable vascular scaffolds to conventional stenting are needed to clarify their role in this cohort.

**Acknowledgements**

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### Table 1. Summary of bioresorbable vascular scaffolds in saphenous venous grafts.

<table>
<thead>
<tr>
<th>Author</th>
<th>Country</th>
<th>Cohort</th>
<th>Gender</th>
<th>Age</th>
<th>Size</th>
<th>Dilatation</th>
<th>Imaging</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ong(^7)</td>
<td>Singapore</td>
<td>1</td>
<td>M</td>
<td>52 years</td>
<td>3.0x18 mm</td>
<td>Pre</td>
<td>CTCA 3 months post insertion</td>
<td>Presentation with NSTEMI and patent stent at 3 months on CTCA</td>
</tr>
<tr>
<td>Roleder(^9)</td>
<td>Poland</td>
<td>1</td>
<td>M</td>
<td>83 years</td>
<td>2.5x15 mm</td>
<td>N/A</td>
<td>Repeat angiography 2 weeks post insertion</td>
<td>Re-presentation with unstable angina and patent stent at 2 weeks on conventional angiography</td>
</tr>
<tr>
<td>Roleder(^10)</td>
<td>Poland</td>
<td>6</td>
<td>M x4 F x2</td>
<td>Median 73 years (57-83)</td>
<td>Median RLD 3.13 mm (2.25-3.45). Median MLD 1.12 (0.52-1.86 mm)</td>
<td>Pre (100%) Post (100%)</td>
<td>Post-procedure OCT, surveillance OCT at 6 months and 18 months</td>
<td>Only one with 18-month and the rest 6-month follow-up with optical coherence imaging, with the majority showing neointimal formation.</td>
</tr>
<tr>
<td>Yew(^8)</td>
<td>Malaysia</td>
<td>1</td>
<td>M</td>
<td>53 years</td>
<td>3.0x28 mm</td>
<td>Pre + Post</td>
<td>Repeat PCI 4 weeks post angiogram</td>
<td>Presentation with ACS followed up with conventional angiography 4 weeks later</td>
</tr>
<tr>
<td>Wojakowski(*)</td>
<td>Poland</td>
<td>1</td>
<td>M</td>
<td>54 years</td>
<td>3.5x12 mm</td>
<td>Pre</td>
<td>Post-procedure OCT 3 months following insertion</td>
<td>54-year-old man with stable angina. Optical coherence imaging at 3 months showed complete apposition</td>
</tr>
<tr>
<td>Picard(^20)</td>
<td>Canada</td>
<td>10</td>
<td>M x9 F x1</td>
<td>Median 75 years (64-81)</td>
<td>Median diameter 3.5 mm (3-3.5). Median length 18 mm (18-28)</td>
<td>Pre (84%) Post (77%)</td>
<td>N/A</td>
<td>Target vessel revascularisation similar to conventional stents (13 lesions)</td>
</tr>
</tbody>
</table>

\(*\) TCT 2014 C-167; ACS: acute coronary syndrome; CTCA: computed tomography coronary angiography; F: female; M: male; MLD: minimal lumen diameter; mm: millimetres; N/A: not available; NSTEMI: non-ST elevation myocardial infarction; OCT: optical coherence tomography; PCI: percutaneous coronary intervention; Post: post-dilation; Pre: predilation; RLD: reference lumen diameter
Conflict of interest statement
The authors have no conflicts of interest to declare.

References


**Supplementary data**

**Moving image 1.** Optical coherence tomography study of the saphenous vein graft bioresorbable vascular scaffold.

The supplementary data are published online at: www.asiaintervention.org