Multimodality intravascular imaging of bioresorbable vascular scaffolds implanted in vein grafts

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Abstract

Introduction: There are no data presenting a serial assessment of vein graft healing after bioresorbable vascular scaffold (BVS) implantation at long-term follow-up.

Aim: To describe ABSORB BVS healing in vein grafts by optical coherence tomography (OCT) and high-definition intravascular imaging (HD-IVUS) at long-term follow-up.

Material and methods: The study group consisted of 6 patients. The first patient had serial OCT assessment of BVS implanted in the saphenous vein grafts (SVG) at baseline and at 3-, 6-, 18-month follow-up and the second patient had OCT assessment of BVS implanted in the SVG at baseline and 24-, 48-month follow-up. The second and the third patients had OCT and HD-IVUS imaging at baseline and 48-month follow-up. The last 3 patients had OCT imaging of BVS implanted in the native coronary artery at 48-month follow-up.

Results: There were no differences in neointimal hyperplasia after BVS implantation between each time point. However, complete scaffold coverage was observed only 48 months after implantation. Out of 202 analyzed scaffold struts, there were 67 (33%) black boxes detectable at 48-month follow-up. HD-IVUS presented plaque burden up to 67% at the segment of BVS implantation at 48-month follow-up. There was a difference in neointimal hyperplasia thickness (1.27 (0.953–1.696) vs. 0.757 (0.633–0.848), p < 0.001) between a native coronary artery and BVS scaffolds at 48-month follow-up.

Conclusions: Bioresorbable vascular scaffold implanted in SVG characterized moderate neointimal hyperplasia as excessive as compared to native coronary arteries at long-term follow-up. The complete scaffold coverage was observed only 48 months after implantation.

Key words: ABSORB, vein graft, optical coherence tomography, high-definition intravascular ultrasound.

Summary

This study sought to describe bioresorbable vascular scaffold (BVS) ABSORB healing in vein grafts by optical coherence tomography (OCT) and high-definition intravascular imaging (HD-IVUS) at long-term follow-up. Complete scaffold coverage was observed only 48 months after implantation. There was a difference in NIH thickness (1.27 (0.953–1.696) vs. 0.757 (0.633–0.848), p < 0.001) between a native coronary artery and BVS scaffolds at 48-month follow-up. HD-IVUS presented plaque burden up to 67% at the segment of BVS implantation in the vein graft at 48-month follow-up.

Introduction

In the last years, implantation of bioresorbable vascular scaffolds (BVS) have attracted worldwide interest as an equally valuable alternative to drug-eluting stents to treat coronary vessel disease. The outcomes of the first clinical trials were so promising that BVS implantation...
became a commonly accepted therapy of coronary artery disease [1, 2]. Biodegradable vascular scaffolds was not only used to treat simple coronary lesions but was also implanted in bifurcations, chronic total occlusions, in-stent restenosis and de novo lesion of saphenous grafts [3–6]. Unfortunately, the initial enthusiasm vanished because the long patient follow-up resulted in increased incidence of late scaffold thrombosis after BVS implantation [7, 8].

Although BVS implantation is not now recommended during percutaneous coronary interventions [9], its remodeling and healing in a different clinical setting have not been comprehensively described yet [10]. One such unusual clinical scenario is the implantation of BVS to de novo lesions of vein grafts [11, 12]. The first clinical observations presented promising results of BVS implantation in vein grafts, and intravascular imaging presented favorable vein graft healing after BVS implantation at short-term follow-up [5, 13, 14]. However, there were no data presenting a serial assessment of vein graft healing after BVS implantation at long-term follow-up.

**Aim**

The following study sought to describe BVS healing in vein grafts by optical coherence tomography (OCT) and high-definition intravascular imaging (HD-IVUS) at long-term follow-up. The obtained results were compared with BVS healing in native coronary arteries.

**Material and methods**

It was a single center study evaluating the vessel healing after BVS ABSORB (Abbott Laboratories, USA) [1] implantation in vein grafts by multimodality imaging including optical coherence imaging (OCT) and high-definition intravascular ultrasound (HD-IVUS) in patients with stable coronary artery disease (CAD) and acute coronary syndromes (ACS). The study was approved by the Ethics Committee of the Medical University of Silesia (KNW/0022/KB1/39/18) and conforms to the Declaration of Helsinki. All of the patients were enrolled in the study only after the patients gave their informed written consent.

**Inclusion/exclusion criteria**

The study included patients with a history of coronary artery bypass grafting (CABG) utilizing vein grafts with recent stable CAD or ACS to implant and to perform serial intravascular imaging of BVS. Furthermore, the study included patients 48 months after BVS implantation in native coronary arteries to compare vessel healing after BVS implantation between SVG and native coronary arteries. The study exclusion criteria were as follows: age < 18 years old, glomerular filtration rate less than 45 ml/min/1.73 m², severe valve disease warranting redo cardiac surgery and contrast allergy.

**Optical coherence tomography imaging**

Optical coherence tomography imaging was performed after the scaffold implantation in SVG and at the follow-up. It was also performed 48 months after BVS implantation in native coronary arteries. The St Jude iLumien OPTIS Medical system was used for OCT imaging. The OCT probe (a mid marker of the OCT Dragonfly catheter) was positioned 5 mm distally to the scaffold intended to analyze. All OCT imaging was performed using automated pullback triggered by the manual injection of contrast.

**Optical coherence tomography image analysis**

CASS intravascular software 2.0 (Pie medical company) was used for offline analysis of the implanted BVS. The region of interest was selected between the proximal and distal edges of the BVS visible by OCT as struts occupying more than 180 degrees of the lumen’s circumference. The analysis was performed every 1 mm to measure lumen area (LA), lumen diameters (LD) and endoluminal and out scaffold area (SA). Endoluminal SA was measured at the inter circumference of polymeric struts and out SA was measured at the outer rim of polymeric struts. The eccentric index (EI) was measured as follows: 

\[
EI = 1 - \left(\frac{\text{minimal lumen area}}{\text{maximal lumen area}}\right)
\]

Lessions with EI > 0.3 were defined as eccentric lesions.

Polymeric struts apposition was also assessed, and if there was a gap between the polymeric strut and the vessel’s lumen contour, malapposition was diagnosed [15]. At the follow-up, polymeric struts’ coverage by neointima was also assessed. The complete coverage of BVS by neointima was identified if four corners of the polymeric strut had lost the right-angle shape with signs of tissue coverage [16]. To measure the tissue thickness the distance from every black box to the lumen contour was measured. Since the thickness of the BVS contours is 30 μm, this value was subtracted from the final tissue measurement to present neointimal hyperplasia (NIH, μm). Neointimal area was measured by subtracting as follows: endoluminal SA – (lumen area + malapposition area).

**HD-IVUS image analysis**

HD-IVUS imaging was performed using the ACIST HDi system and ACIST Kodama IVUS catheter device. The region of interest was the segment of the artery where the BVS was previously implanted. Quantitative grey-scale IVUS measurements were performed every millimeter in scanned coronary segments. Cross-sectional images were quantified for lumen diameters and area, external elastic membrane (EEM) diameters and area, total plaque area (TPA) and plaque burden (PB). Since all HD-IVUS imaging was performed at 48 months after implantation no BVS...
strut detection was performed. TPA was calculated as the difference between EEM area, and PB was calculated as total plaque area (TPA) divided by EEM area × 100 (%).

**Statistical analysis**

Continuous parameters were reported as mean with standard deviation and median with the first and the third quartiles. Discrete data were summarized as frequencies and group percentages. Wilcoxon signed-rank test and the χ² test with Rao and Scott adjustment were used for comparison of continuous and categorical data, respectively. P-value < 0.05 was considered statistically significant. Analyses for statistical computing were performed using MedCalc version 18.6 (MedCalc Software, Belgium).

**Results**

**Study group**

The study group consisted of 6 patients. Forty-eight months imaging was performed in 5 patients. The first one could not have the intravascular imaging at 48-month follow-up because she died 40 months after BVS implantation due to a non-cardiac cause (lymphoma). The others did not experience any adverse cardiac events at 48-month follow-up. All patients received DAPT for the 12 months after BVS implantation. Patients and BVS characteristics, and the study flow chart are presented in Figure 1.

**Serial OCT follow-up of BVS after its implantation in SVG**

The first BVS was observed at baseline and at 3, 6 and 18 months after implantation. The MLA and MLD decreased after 3 months, and then it significantly increased at 18 months. There were no differences in NIH and the neointimal area between each time point respectively. Interestingly, endoluminal SA was systematically increasing at each time point, but the outer SA increased only in the sixth month after BVS implantation (Figure 1, Table I). The second BVS was observed at baseline, 24 and 48 months after its implantation. MLA and MLD decreased at 24- and 48-month follow-up as compared to baseline values (Table I). There were no differences in NIH and the neointimal area between each time point. The endoluminal SA and outer SA increased over the time of observation (Figure 2, Table I).

| Vein grafts | Coronary arteries |
|-------------|------------------|
| 1st patient, female, 73 y.o., 3 years after CABG, HA, HL, DM, GFR 57 ml/min, non-smoker, ABSORB 3.0 x 18 mm | 4th patient, male 57 y.o., no CABG, HA, HL, DM, GFR 63, smoker, ABSORB 3.0 x 12 mm |
| Baseline OCT | Baseline OCT |
| 3-month OCT | 3-month OCT |
| 6-month OCT | 6-month OCT |
| 18-month OCT | 18-month OCT |
| 24-month OCT | 24-month OCT |
| 48-month OCT and HD-IVUS | 48-month OCT and HD-IVUS |
| 48-month OCT | 48-month OCT |
| 48-month OCT | 48-month OCT |
| 48-month OCT | 48-month OCT |

**Figure 1.** Study chart flow. The study chart flow presents the time of optical coherence tomography (OCT) and high definition intravascular (HD-IVUS) imaging of ABSORB implanted in vein grafts and coronary arteries.

HA – hypertension, HL – hyperlipidemia, DM – diabetes mellitus, CABG – coronary artery bypass grafting, GFR – glomerular filtration rate (ml/min/1.73 m²).
48-month follow-up after BVS implantation in SVG

There were 26 OCT cross-sections analyzed at baseline and 17 OCT cross-sections analyzed at 48-month follow-up for both BVS. Out of 202 scaffold struts, there were only 67 (33%) black boxes visible at 48-month follow-up. The MLA (mm²), 7.45 (4.65–8.75) vs. 4 (3.51–3.01), p < 0.001 and MLD (2.81 (2.25–3.05) vs. 1.91 (1.81–2.21), p < 0.001) were smaller at 48-month follow-up as compared to baseline. The endoluminal SA (7.3 (4.85–7.9) vs. 5.5 (5.2–5.7), p = 0.464) and out SA (9.35 (6.37–10.25) vs. 7.16 (6.86–7.83), p = 0.691) decreased at 48-month follow-up. HD-IVUS was not able to visualize struts at 48-month follow-up and presented MLA = 3.4 mm², MLD = 2 mm, PB = 44% and TPA = 2.7 mm² and TPV = 33.7 m² for the first patient, and MLA = 4.6 mm², MLD = 2.3 mm, PB = 62% with TPA = 6.2 mm² and TPV = 68.7 mm² for the second patient. HD-IVUS also presented the echogenic rim of neointima remaining after the BVS implantation (Figure 3).

48 months after BVS implantation in SVG vs. native coronary artery

There were 41 OCT cross-sections and 268 struts analyzed in native coronary arteries and 17 cross-section and 67 struts analyzed in SVG. There were no differences in MLA (4.8 (3.6–8.9) vs. 4.0 (3.5–5), p = 0.228), endoluminal SA (6.15 (5.1–11.1) vs. 5.5 (5.2–5.7), p = 0.228) and out SA (7.72 (6.61–13.38) vs. 7.16 (6.87–7.83), p = 0.3664) between native and SVG scaffolds. However, there was a difference in NIH thickness (1.27 (0.953–1.696) vs. 0.757 (0.633–0.848), p < 0.001) between a native coronary artery and BVS scaffolds at 4 years follow-up.

Discussion

It is the first description of BVS healing by serial OCT imaging at long-term follow-up. The main finding of the study was as follows: 1) BVS healing in vein grafts was a dynamic process with the reduction of lumen area during

Table I. Serial OCT assessment of ABSORB scaffold implanted in SVG

| Parameter          | First patient | Second patient |
|--------------------|---------------|----------------|
| Parameter          | Time-point    | Time-point     |
|--------------------|---------------|----------------|
| Cross-sections, n  | First patient | Second patient |
| Baseline           | 12            | 20             |
| 3M                 | 19            | 15             |
| 6M                 | 24            | 20             |
| 18M                | 10            | 15             |
| Baseline           | 11            | 13             |
| 24M                | 13            | 13             |
| 48M                | 13            | 13             |
| Lumen area [mm²]   | 6.55          | 7.45           |
| (6.21–7.15)        | (4.65–8.75)   | (3.91–4.42)    |
| 3M                 | 5.91          | 7.81           |
| (5.80–6.45)        | (6.92–8.53)   | (3.51–4.12)    |
| 6M                 | 7.03          | 4.22           |
| (6.12–7.40)        | (4.65–8.75)   | (3.51–4.12)    |
| 18M                | 7.81          | 4.82           |
| (6.92–8.53)        | (4.65–8.75)   | (3.51–4.12)    |
| Min lumen diameter [mm] | 2.62        | 2.81            |
| 3M                 | 2.42          | 2.00           |
| (2.51–2.65)        | (2.25–3.05)   | (1.71–2.00)    |
| 6M                 | 2.62          | 2.19           |
| (2.35–2.80)        | (2.25–3.05)   | (1.71–2.00)    |
| 18M                | 2.92          | 2.00           |
| (2.71–3.12)        | (2.25–3.05)   | (1.71–2.00)    |
| Endoluminal SA [mm²] | 5.71          | 5.92 |
| 3M                 | 7.72          | (7.5–7.75)     |
| (7.5–7.75)         | (4.65–8.75)   | (3.51–4.12)    |
| 6M                 | 7.73          | 8.22           |
| (7.5–7.75)         | (4.65–8.75)   | (3.51–4.12)    |
| 18M                | 8.22          | 8.81           |
| (7.65–9.12)        | (4.65–8.75)   | (3.51–4.12)    |
| Out SA [mm²]       | 7.74          | 7.45           |
| (7.45–7.95)        | (5.91–6.72)   | (5.51–6.37)    |
| Endoluminal SA [mm²] | 7.73          | 8.22           |
| 3M                 | 9.31          | 10.15          |
| (8.26–9.81)        | (9.39–11.07)  | (5.43–6.79)    |
| 6M                 | 9.31          | 10.15          |
| (8.26–9.81)        | (9.39–11.07)  | (5.43–6.79)    |
| 18M                | 9.31          | 10.15          |
| (8.26–9.81)        | (9.39–11.07)  | (5.43–6.79)    |
| Eccentricity index | 0.19          | 0.19           |
| (0.14–0.24)        | (0.16–0.22)   | (0.13–0.17)    |
| EI > 0.3, n (%)    | 0%            | 0%             |
| Covered struts, n (%) | 110         | 113            |
| Scaffold struts, n | 110           | 113            |
| Covered struts, n (%) | 158         | 151            |
| Malapposed struts, n (%) | 6 (5) | 4 (1) |
| NIH [μm]           | 700           | 725            |
| (460–1040)         | (542–1007)    | (683–1006)     |
| Neointimal area [mm²] | 0.22         | 0.22           |
| (0–0.51)           | (0.15–0.81)   | (0.51–1.05)    |

M – months, SA – scaffold area, NIH – neointimal hyperplasia measured for each scaffold, n – number.
In our study, we observed a nearly complete scaffold coverage at 3-month follow-up, which is in line with previous reports [13]. Long-term observation showed a stable thickness of BVS coverage in the vein graft, but full scaffold coverage was observed only at 48-month follow-up. It is in contrast with data obtained from BVS implanted in native coronary arteries [19]. Almost complete (~99%) coverage was found after BVS implantation in stable CAD and STEMI patients at 12-month follow-up [17, 18].

Interestingly, the vein graft lumen presented dynamic remodeling after BVS implantation, which is not in line with previous reports. The BVS lumen area remained stable after 12 months in both patients with stable CAD and ACS [18, 20]. Previous studies also showed constant remodeling of the vessel eccentricity in the first 24 months after BVS implantation in SCAD [10, 21]. However, a higher percentage of eccentric lesions was observed after the index procedure, within a time the vessel concentricity increased and remained stable after 2 years. Our study showed that BVS characterized stable scaffold concentricity up to 48 months after the implantation. It may be explained by the fact that vein graft lesions are less calcified and thus an appropriate scaffold shape was much more easily achieved as com-
pared to native coronary arteries [22, 23]. Additionally, the stable BVS shape may explain the lack of malapposition during the follow-up, which in line with the animal model observations [24], but in contrast to human studies. BVS malapposition was observed even up to 36 months after the implantation [20]. Interestingly, HD-IVUS presented an echogenic rim at 48 months after implantation. Since the OCT images did not present calcifications, the rim represents the acoustic tissue properties at the site of BVS implantation. Further studies are warranted to determine what is left in the vessel wall after BVS absorption.

This is the first report presenting the 48-month results of BVS implantation in vein grafts in comparison to native coronary arteries. It showed that NIH analyzed on the remaining scaffold was higher as compared to that observed in native coronary arteries. This, together with incomplete scaffold strut coverage, suggests that the BVS is less favorable to vessel healing in vein grafts. Greater NIH may be promoted by the lower shear stress in the SVG, as compared to native coronary arteries [25]. Furthermore, IVUS data revealed hypoechogenic lesions at the site of BVS implantation at 48-month follow-up, but their burden did not differ from those observed 3 years after BVS implantation in native coronary arteries [20]. The previous report documented progression of calcification at the segment of previous BVS implantation [26].

Figure 3. Representative angiography, OCT and HD-IVUS imaging of ABSORB implanted in the vein graft at 48-month follow-up. A – The coronary angiography presents the vein graft 48 months after ABSORB implantation. The dashed lines indicate (1, 2, 3) the position of HD-IVUS (B, D, F) and OCT cross-section images (C, E, G). The white arrows indicate the echogenic rim of the remaining neointima (B, D, F) and blue triangles indicate black boxes remaining (C, E, G) after ABSORB implantation.

Study limitations
The study enrolled a small number of patients, which is the main limitation. The serial intravascular imaging assessment was performed only in 2 patients, which makes it hard to draw any general conclusion from these observations. Unfortunately, it was impossible to enroll more patients because the distribution of ABSORB was stopped during the study period. The results come merely from the cross-sectional analysis. There was no propensity matching between patients with BVS implanted in vein grafts and native coronary arteries, which could also bias the comparison of vessel healing between these two clinical scenarios.

Conclusions
The serial intravascular imaging of BVS implanted in the SVG presented dynamic lumen remodeling but with stable scaffold concentric shape. The neointimal hyperplasia was not excessive, but the scaffold coverage was not complete even 24 months after the implantation.
HD-IVUS presented a remaining plaque at the site of BVS implantation at 48-month follow-up.

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Conflict of interest

The authors declare no conflict of interest.

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