Bioresorbable scaffolds in the treatment of coronary artery disease

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Abstract: Drug-eluting stents have reduced the risk of in-stent restenosis and have broadened the application in percutaneous coronary intervention in coronary artery disease. However, the concept of using a permanent metallic endovascular device to restore the patency of a stenotic artery has inherited pitfalls, namely the presence of a foreign body within the artery causing vascular inflammation, late complications such as restenosis and stent thrombosis, and impeding the restoration of the physiologic function of the stented segment. Bioresorbable scaffolds (BRS) were introduced to potentially overcome these limitations, as they provide temporary scaffolding and then disappear, liberating the treated vessel from its cage. Currently, several BRSs are available, undergoing evaluation either in clinical trials or in preclinical settings. The aim of this review is to present the new developments in BRS technology, describe the mechanisms involved in the resorption process, and discuss the potential future prospects of this innovative therapy.

Keywords: bioresorbable scaffold, drug-eluting stent, biodegradable, design, mechanism, coronary artery disease

Introduction

In 1977, Andreas Grüntzig introduced percutaneous transluminal coronary angioplasty, opening new horizons in the treatment of coronary artery disease (CAD).1,2 Although the potential of this less invasive approach was apparent, the risk of acute occlusion and the high incidence of restenosis caused by elastic recoil and neointimal hyperplasia limited its application.3–5 Bare metal stents (BMSs) provided a partial solution to these problems, by eliminating the risk of acute occlusion and reducing the occurrence of restenosis.6–10 With the introduction of drug-eluting stents (DESs), the properties of the coated antiproliferative drug prolonged vessel wall healing, limited neointimal hyperplasia, and subsequently reduced the prevalence of target lesion revascularization (TLR) to less than 10%.11 This resulted in extensive application of DES, especially with the use of intravascular ultrasound for the patients with CAD.12–16 Conversely, delayed endothelization, and the hypersensitivity reaction of the vessel wall to the durable polymer, increased the risk of late and very late stent thrombosis (ST).17,18 Recent advances in DES technology with the use of either a bioresorbable or compatible polymer reduced the occurrence of ST but still failed to address other limitations of permanent metallic DES, namely the potential risk of neo-atherosclerosis, preclusion of surgical revascularization, and the distorted vessel wall physiology caused by the presence of a foreign body within the artery.15,19,20

For a number of decades, interventional cardiologists have pursued the possibility of a fully bioresorbable scaffold (BRS) that will initially safeguard the patency of
the vessel, and then will disappear, thus allowing the artery to restore its physiological integrity.\textsuperscript{21–24} The feasibility of this concept was first investigated in the 1990s with the Igaki-Tamai\textsuperscript{8} device (Igaki Medical Planning Co, Ltd, Kyoto, Japan); this process was however temporarily halted with the introduction of metallic DESs. Recently, BRS has been heralded as the fourth revolution, rendering a novel treatment termed vascular restoration therapy (VRT) in interventional cardiology.\textsuperscript{25} The clear feature of VRT is that vessel scaffolding is degraded evanescently, and therapeutic advantages may be derived from allowing natural vasomotor response in the vessel. In addition, the underlying benefits involve decreased or eliminated late ST, facilitated re-intervention due to the complete reabsorption of the implanted scaffold, improved noninvasive imaging of the target site using computed tomography or magnetic resonance during post-intervention follow-up, and shortened prescription of dual anti-platelet therapy.\textsuperscript{26} Accordingly, for BRS, it is potentially a ideal therapy for CAD, although there is still a long way to go to make improvements.

Over the last 5–10 years, considerable efforts were made to develop new fully bioresorbable devices. Currently, BRS technology has been gradually matured, and there are numerous devices available which undergo preclinical or clinical testing. In this review article, we present the mechanisms of BRS technology, the evolution of individual devices, discuss the developments of the currently available BRS, and underline their potential value in the treatment of CAD.

**Design and resorption of the BRSs**

For a BRS to be as effective as the currently available DES, it should have an increased radial strength that should be maintained for at least 3–6 months post device implantation and incorporate an antiproliferative drug that would control neointimal formation and prevent restenosis. The prolonged radial strength can be provided by an appropriate scaffold design (eg, increased strut thickness, closed cell design) and by the incorporation of a resilient resorbable material (polymer or metallic alloy) which would gradually resorb so as to prevent late recoil and safeguard the patency of the vessel. Although the first generation BRS did not include an antiproliferative drug, more recent updated revisions incorporate a fully biodegradable polymer that controls the release of a drug elution with antiproliferative properties.

The first polymeric (nonbioresorbable polyethylene-terephthalate) braided mesh stent was designed in 1992.\textsuperscript{27} Preclinical evaluation in porcine models demonstrated that the developed polymeric stent had comparable safety and efficacy to the cotemporary metallic stent. In 1996, van der Giessen et al introduced the Wiktor stent, coated with five different fully bioresorbable polymers.\textsuperscript{28} The experimental study showed an excessive inflammatory reaction of the vessel wall to the device, resulting in increased neointimal proliferation and thrombus formation. One year later, Lincoff et al reported the first high molecular weight poly-L-lactic acid (PLLA, 321 kDa) coated stent, that in contrast to the low molecular weight (80 kDa) PLLA coated stent, appeared to be well tolerated by the vessel wall.\textsuperscript{29} Yamawaki et al were the first to develop a fully BRS, consisting of PLLA and a tyrosine kinase inhibitor. Evaluation of the device in porcine models demonstrated the safety and efficacy of the device and supported its implantation in humans in 1998.\textsuperscript{30} These preliminary encouraging results attracted interest and drove researchers and industry towards the development of several BRSs with different bioresorbable properties and composition.\textsuperscript{19,31–33}

Most of the currently available BRSs (eg, Igaki-Tamai scaffold, Absorbtm [Abbott Vascular, Santa Clara, CA, USA] bioresorbable vascular scaffold [BVS]) are composed of PLLA. The catabolism of the PLLA incorporates five stages, with the resultant degradation of the polymer to carbon dioxide and water (Figure 1).\textsuperscript{34,35} The first phase includes hydration of the polymer, which starts to absorb water from the surrounding tissue. The absorbed water catalyzes a chain scission at an ester bond, resulting in the degradation of the polymer (second stage). Gradually the polymer loses its cohesive strength and fragments into segments with a lower molecular weight (third stage). Polymer chains become progressively hydrophilic through their hydrolysis to shorter lengths. The outcomes of the abovementioned process are hydrophilic monomers that can be phagocytized by macrophages (fourth phase). After phagocytosis, the soluble monomer (L-lactate) is catabolized to pyruvate and ultimately to carbon dioxide and water through the Krebs cycle (fifth stage).\textsuperscript{36,37}

Tyrosine polycarbonate is another polymer used in BRS technology (REVA BRS). Its catabolism includes hydrolysis of the polymer to carbon dioxide and iodinated-desaminotyrosyl-tyrosine ethyl esters, which are further hydrolyzed to ethanol and iodinated-desaminotyrosyl-tyrosine. Cleavage of the latter results in tyrosine molecules and iodinated-desaminotyrosine, which are finally catabolized into carbon dioxide and water through the Krebs cycle (Figure 2).

Another material used in BRSs to date is magnesium, the only metal that has been implemented in BRS technology
(absorbable metallic stent [AMS]). A limitation of magnesium is its fragility; thus, it has been mixed with several elements such as zirconium, yttrium, and other rare earth metals to provide it with adequate radial strength. An advantage of the AMS is the fact that the degradation of the magnesium alloy to inorganic salts (Figure 3) triggers only a minor inflammatory response, and creates an electronegative charge that has been shown to have an anti-thrombogenic effect.38

Currently available BRSs

Currently, there are more than 14 BRSs available (Table 1 and Figure 4). Most are still either under preclinical evaluation or being examined in the clinical setting. Of note, two have already acquired certificate Conformité Européenne (CE) mark approval and are used in clinical practice: the Igaki-Tamai stent for the treatment of peripheral vascular disease, and the Absorb BVS for CAD. The following sections describe the BRSs that are under clinical or preclinical evaluation, and present the evidence from the first applications of BRSs in clinical settings or experimental studies.

Igaki-Tamai scaffold

The Igaki-Tamai scaffold was the first fully bioresorbable stent implanted in humans and was made of PLLA. The first revision of scaffold had a helical zigzag design, was mounted on a standard angioplasty balloon, and was both thermal self-expanding and balloon expandable. Scaffold expansion was performed with the use of heated contrast (up to 70°C). The device continued expanding after its implantation over a 20–30 minute period, until it reached its final dimensions.

The first-in-man (FIM) study of the Igaki-Tamai scaffold included 15 patients treated with 25 scaffolds, and showed no major adverse cardiac event (MACE) at
the first 30 days and only one TLR at 6-month follow-up.\textsuperscript{39} Angiographic follow-up performed at day 1 after the device implantation revealed a similar percentage diameter stenosis and minimal lumen diameter (MLD) compared with the post-procedural values (12% ± 8% versus 13% ± 11%; 2.59 ± 0.35 mm versus 2.58 ± 0.32 mm). At 3 months, the MLD decreased (1.88 ± 0.59 mm), with minimal progression thereafter at 6 months follow-up (1.84 ± 0.66 mm). Serial intravascular ultrasound (IVUS) examinations demonstrated a gradual increase in the scaffold cross-sectional area (from 7.42 ± 1.51 mm\textsuperscript{2} post stenting, to 8.18 ± 2.42 mm\textsuperscript{2} at 3 months, and 8.13 ± 2.52 mm\textsuperscript{2} at 6 months; \(P < 0.1\), \(P < 0.1\); respectively), whilst the luminal dimensions decreased at 3 months, and remained unchanged at 6 months (from 7.42 ± 1.51 mm\textsuperscript{2} after stenting to 5.67 ± 2.42 mm\textsuperscript{2} at 3 months, and 5.63 ± 2.70 mm\textsuperscript{2} at 6 months; \(P < 0.005\), \(P < 0.001\); respectively).

Recently, Nishio et al reported the long-term follow-up (>10 year) of an observational prospective study which enrolled 50 patients (63 lesions) implanted with Igaki-Tamai scaffolds (84 scaffolds).\textsuperscript{40,41} The 10-year cumulative event-free survival rates of all-cause death, cardiac death, and MACE were 87%, 98%, and 50%, respectively, while the TLR rate was 16% at 1 year, 18% at 5 years, and 28% at 10 years. Angiographically, the late lumen loss (LLL) was 0.91 ± 0.69 mm at 6 months, which improved to 0.67 ± 0.45 mm at 1 year, and was maintained at 3 years follow-up (0.59 ± 0.50 mm). IVUS examinations showed a reduction in the minimal lumen area at 6 months (from 5.44 ± 1.89 mm\textsuperscript{2} after stenting to 3.64 ± 1.68 mm\textsuperscript{2}), which started increasing at 1 (4.06 ± 1.61 mm\textsuperscript{2}) and 3 (5.18 ± 2.09 mm\textsuperscript{2}) years. IVUS echogenicity revealed raised hyper-echogenicity of the scaffolded segment post device implantation, and then subsequently started decreasing to reach the pre-implantation values at 3 years follow-up.

Although the short- and long-term follow-up results were encouraging, this device failed to progress, mainly due to the large guide catheters (8F) required for implantation, and the concerns that the heated contrast used for device deployment may cause vessel wall injury. The new generation Igaki-Tamai scaffold potentially overcomes these pitfalls, as it can be implanted through a 6F guide catheter without the need for

![Figure 2](image-url)\textsuperscript{42} The metabolism of tyrosine-polycarbonate scaffold. Initial hydrolysis of the tyrosine-polycarbonate produces iodinated-desaminotyrosyl-tyrosine ethyl esters (I\textsubscript{2}DTE), and releases carbon dioxide.

\textbf{Notes:} I\textsubscript{2}DTE is hydrolyzed to iodinated-desaminotyrosyl-tyrosine (I\textsubscript{2}DT) and ethanol. Cleavage of I\textsubscript{2}DT produces tyrosine, and iodinated-desaminotyrosine (I\textsubscript{2}DAT), which enters into the Kreb’s cycle.

![Figure 3](image-url)\textsuperscript{43} Schematic representation of the resorption process in the drug-eluting absorbable magnesium scaffold.

\textbf{Notes:} The release of the anti-proliferative drug occurs within the first 3 months after device implantation. Hydrolysis of the scaffold affects the radial strength of the scaffold, resulting in a gradual degradation of the device into a soft amorphous hydroxyapatite at 9 months follow-up.

\textbf{Abbreviation:} DREAM, drug-eluting absorbable metallic stent.
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<tr>
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<th>Company</th>
<th>Status</th>
<th>Strut material</th>
<th>Coating material</th>
<th>Drug elution</th>
<th>Design</th>
<th>Strut thickness (μm)</th>
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**Abbreviations:** BRS, bioresorbable scaffold; BVS, bioresorbable vascular scaffold; CE, Conformité Européenne; DREAMS, drug-eluting absorbable metallic stents; PDLA, poly-D, L-lactide acid; PLCL, poly-L-lactide-co-ε-caprolactone; PLGA, poly-lactide-co-glycolide; PLLA, poly-L-lactic acid.
a heated contrast agent. The device is currently undergoing preclinical evaluation in Germany.

Absorb BVS

The Absorb BVS has a backbone composed by PLLA, and is coated with a poly-D,L-lactide (PDLA) polymer that contains the anti-proliferative drug everolimus. Histology based studies in porcine models have demonstrated that scaffold resorption is completed within 3 years post device implantation (Figure 5).42

The first generation of Absorb BVS (BVS 1.0) was examined in the ABSORB Cohort A Trial (A Clinical Evaluation of the Biodegradable Everolimus Eluting Coronary Stent System in the Treatment of Patients with de Novo Native Coronary Lesions). The trial aimed to assess the feasibility and safety of the scaffold in patients with single de novo coronary artery lesion. In this prospective, open-label study, 30 patients who had stable, unstable, or silent ischemia were enrolled.33 The composite endpoint of MACE was cardiac death, myocardial infarction (MI), and ischemia-driven TLR. The cumulative estimated incidence of MACE was 3.3%, with only one patient having a non-Q-wave MI and no TLR at 1-year follow-up.43 Numerous invasive imaging modalities were used to assess the efficacy of the device, including serial (at baseline, 6 months, and 2 years) coronary angiography, IVUS, IVUS echogenicity, IVUS radiofrequency backscatter analysis, palpography and optical coherence tomography (OCT). Angiographically, the in-scaffold LLL did not differ between the 6-month and 2-year follow-up (0.44 ± 0.35 mm, 0.48 ± 0.28 mm). Intravascular ultrasound imaging demonstrated a reduction in the lumen area between baseline and follow-up (6.04 ± 1.12 mm² at baseline versus 5.19 ± 1.33 mm² at 6 months and 5.47 ± 2.11 mm² at 2 years), which was partially attributed to the scaffold shrinkage (6.94 ± 1.70 mm² immediately post-procedure and 6.29 ± 1.47 mm² at the 6 months follow-up).44 Given this drawback, the scaffold was redesigned, and the new revision had a more uniform strut distribution and provided increased radial strength to vessel wall.45

The second generation of the Absorb BVS (BVS 1.1) was tested in the ABSORB Cohort B trial. A total of 101 patients (102 lesions) treated with 3.0 mm × 18 mm Absorb BVS devices were enrolled in this study and divided into two groups: Cohort B1 and Cohort B2.46,47 Cohort B1 had invasive follow-up examinations (quantitative coronary angiography [QCA], IVUS, and OCT) at 6 months and 2 years, while Cohort B2 had the same investigations at 1 and 3 years. Computed tomographic coronary angiography was performed in both groups at 18 months follow-up. The rate of MACE in 101 patients was 9.0% (three non-Q-wave MI, six ischemia-driven TLR) at 2-year follow-up.48 In the Cohort B1 group, QCA analyses demonstrated an LLL of 0.19 ± 0.18 mm at the 6-month, and 0.27 ± 0.20 mm at the 2-year follow-up.49 IVUS examinations performed in 33 patients demonstrated the minimum lumen area to be reduced at 6 months (from 5.45 ± 1.08 mm² post-procedure to 5.12 ± 1.01 mm²), with no changes at 2 years (5.13 ± 1.25 mm²),
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Abbreviations: OCT, optical coherence tomography; BVS, bioresorbable vascular scaffold.

whereas the mean lumen area and mean scaffold area decreased at 6 months (from $6.53 \pm 1.24$ mm$^2$ post-procedure to $6.36 \pm 1.18$ mm$^2$ and from $6.53 \pm 1.23$ mm$^2$ post-procedure to $6.42 \pm 1.17$ mm$^2$, respectively), and then increased at 2 years ($6.85 \pm 1.78$ mm$^2$, $7.08 \pm 1.73$ mm$^2$, respectively). OCT examinations showed similar results in 23 patients. In the Cohort B2 group, the 1-year follow-up results demonstrated the LLL to be $0.27 \pm 0.32$ mm. The mean scaffold area and mean lumen area detected by IVUS were similar between post-procedure and 1 year follow-up ($6.29 \pm 0.92$ mm versus $6.33 \pm 0.98$ mm and $6.31 \pm 0.95$ mm versus $6.33 \pm 1.17$ mm, respectively). OCT examinations demonstrated that the mean scaffold area did not change, whereas the mean lumen area decreased significantly by 23.4% ($P < 0.001$).

Apart from the ongoing ABSORB Cohort B trial, three other clinical trials are underway: the ABSORB Physiology, the ABSORB II, and the ABSORB EXTEND. The ABSORB Physiology study aims to estimate the short- and long-term effects of an Absorb BVS and a Xience V* (Abbott Vascular, Santa Clara, USA) stent on the physiology of the vessel wall, and will include the following metrics: vascular compliance, distensibility, endothelial responsiveness, and shear stress distribution. The ABSORB II study is the first randomized trial designed to compare the safety and efficacy of the Absorb BVS and the analogous metallic stent (Xience prime, Abbott Laboratories) in 500 patients with de novo coronary artery disease. Finally, the ABSORB EXTEND registry aims to recruit 1000 patients with de novo single or two-vessel disease and test the efficacy of the device in clinical settings. In contrast to the previous studies, this single-arm study will include long lesions and small caliber vessels with a reference vessel diameter of 2.0–2.5 mm.

Magnesium metallic stents (AMSs)

An AMS (Biotronik, Berlin, Germany) is a tubular, slotted balloon-expandable stent sculptured by laser from a tube of magnesium alloy. The device has similar mechanical strength to other metallic stents. In the first generation AMS 1.0, the resorption process was almost completed at 2 months after
implantation, resulting in late vessel recoil. To address this limitation, the scaffold was redesigned, and in the latest revision (drug-eluting AMS, DREAM 2.0) the process was prolonged to 6 months.

The efficacy of the first generation AMS was validated in the PROGRESS-AMS (Clinical Performance and Angiographic Results of Coronary Stenting) study, which was a nonrandomized, multicenter trial that included 63 patients who had de novo lesions with lengths of 10–15 mm and reference diameters of 3.0–3.5 mm. During a follow-up period of 4 months, there was a 23.8% clinically driven TLR incidence, associated with a high LLL (1.08 ± 0.49 mm). From 4 to 12 months, only one further patient sustained TLR, resulting in a cumulative incidence of MACE at 12 months of 26.7%. These results were attributed to neointimal proliferation and vessel recoil caused by the premature reduction in the radial strength of the implanted devices. To overcome these limitations, the company added the antiproliferative drug paclitaxel, and changed the design of the scaffold and the composition of the magnesium alloy, thus providing the device with increased strength, and a prolongation of its resorption process. The efficacy of the new revision DREAMS (drug-eluting absorbable metallic stents) 1.0 was evaluated in the BIOSOLVE-1 trial. In this prospective multicenter FIM trial, 46 patients with a single de novo coronary artery lesion were implanted with 47 scaffolds. At 1-year follow-up, the MACE rate was 7% (two TLR and one MI), whereas the LLL, although improved, remained high (0.52 ± 0.39 mm).

The second generation DREAMS (DREAMS 2.0) scaffold incorporates sirolimus elution instead of paclitaxel. Preclinical evaluation of the device revealed reduced inflammation, and a higher endothelization rate compared with the previous revision. Further evaluation is however needed to confirm these positive results in clinical settings and to examine its safety and efficacy in the treatment of CAD.

**REVA scaffold**

The REVA scaffold (Reva Medical Inc, San Diego, CA, USA) is made by a tyrosine-derived polycarbonate polymer. The first revision of the REVA scaffold had a distinctive slide-and-lock mechanism that provided for a robust radial strength to the device post implantation.

The long-term (55 months) performance of the first generation REVA scaffold was reported in a healthy porcine model recently. In this report, serial IVUS examinations showed a gradual increase in the lumen area during follow-up (lumen area change, defined as % lumen increase post-implantation: 86% ± 23% at 5 days, 94% ± 10% at 12 months, and 114% ± 23% at 55 months). The clinical performance of this first generation device was evaluated in the RESORB (The REVA endovascular study of a bioresorbable coronary stent) study. Twenty-five patients with single de novo lesions were implanted with the studied scaffolds. Angiographically, the MLD increased from 0.88 ± 0.39 mm to 2.76 ± 0.36 mm after device deployment, resulting in an acute gain similar to that reported with the current metallic stents. However, a high TLR rate was noted between 4- and 6-month follow-up, which was predominantly attributed to focal mechanical failures leading to the redesign of scaffold. The ReZolve is the second revision of REVA scaffold, has a spiral slide-and-lock mechanism, and contains the antiproliferative drug sirolimus. The new generation scaffold is currently undergoing evaluation in the RESTORE (ReZolve sirolimus-eluting bioresorbable coronary scaffold) clinical trial, which aims to investigate its safety and efficacy in 50 patients. The primary endpoints of this study are ischemic-driven TLR at 6 months, and quantitative measurements (QCA and IVUS) at 12 months. Concomitantly, a pivotal trial which aims to apply for the CE mark approval has been initiated.

**DE Solve™ bioresorbable coronary scaffold**

The DESolve BRS (Elixir Medical Corporation, Sunnyvale, CA, USA) is composed of a PLLA polymer and contains two novel antiproliferative drugs (Novolimus and Myolimus). The radial strength of the device is comparable with the Elixir’s BMS, and its resorption process takes approximately 2–3 years.

The safety and efficacy of the DESolve scaffold was evaluated in a prospective, multicenter FIM trial, which included patients with a single de novo coronary artery lesion, with a reference vessel diameter of 3.0 mm and lesion length < 10 mm. At present, 16 patients have been enrolled. Clinically indicated TVR was reported in one patient who underwent emergency coronary artery bypass graft for a procedurally related spiral dissection at 30 days follow-up. One clinically indicated TLR occurred between 30 days and 180 days follow-up, while no cardiac death, MI, and ST have been reported at 6-month follow-up. QCA revealed an LLL of 0.19 ± 0.19 mm at 6-month follow-up. IVUS examination showed a minor reduction in the lumen area and an increase in the scaffold area at 6 months (5.35 ± 0.78 mm² versus 5.10 ± 0.78 mm² at 6 months and 5.35 ± 0.78 mm² versus 5.61 ± 0.81 mm², respectively). OCT examination in ten patients...
showed that 98.68% ± 2.44% of the struts was fully covered at 6 months follow-up, and a mean neointimal hyperplasia area of 0.71 ± 0.36 mm². Based on this initial promising data, the DeSolve NX study has recently commenced, and aims to evaluate the safety and efficacy of the device. The study is expected to include 120 patients with single de novo coronary artery lesions. In-scaffold LLL at 6-month follow-up is the principal endpoint. In addition, the DeSolve NX II pivotal trial has been designed to evaluate the effectiveness of the device in a broader number of patients to provide sufficient evidence for CE mark approval and is expected to start in due course.⁵⁸

**Ideal™ BioStent**

The Ideal BioStent (Xenogenesis Corp, Canton, MA, USA) is a fully bioresorbable balloon expandable scaffold. The device consists of a core backbone and a top coat. The backbone is synthesized from polylactide anhydride, and a trimer of two salicylic acid molecules joined by a sebacic acid, while the top coat part is comprised of salicylate and the antiproliferative drug sirolimus. Preclinical studies have demonstrated the potential anti-inflammatory and antiplatelet properties of salicylate, which appears to reduce restenosis and promote vessel healing during the polymer’s degradation.⁵⁹,⁶⁰

The first generation scaffold (BTI) was examined in the “WHISPER” study. Eleven patients were included in this prospective FIM trial that evaluated the safety and efficacy of the scaffold.⁶¹ Coronary angiography and IVUS, performed post-procedurally and at follow-up, revealed the absence of scaffold recoil. However, IVUS and OCT showed increased neointimal formation, which was attributed to the inadequate drug dose and fast drug elution. In view of the high restenosis rate, the device was redesigned. The revised device (Ideal BioStent) has a higher drug dose, slower drug release kinetics, and an easy to use peel-away sheath. The new generation Ideal scaffold is currently undergoing preclinical evaluation, with a plan for the initiation of clinical trials in the near future.

**ART BRS**

The ART BRS (Arterial Remodeling Technologies, Noisy le Roi, France), is a fully BRS made by a PLLA amorphous polymer, without any antiproliferative drug. The new revision device was named ART18Z. The resorption process of the scaffold starts at 3 months and is expected to be completed between 18 and 24 months. To date, more than 250 devices have been implanted in porcine models, and no MACE (defined as a composite of thrombosis, death, and MI) have been reported.⁶² QCA analyses have shown that the ART18Z scaffold has similar acute recoil and LLL, at 6 and 9 months, in comparison to the Multilink Vision stent (Abbott Vascular, Santa Clara, CA, USA). At 9 months follow-up, the LLL decreased in the ART18Z group, a finding that was attributed to late lumen enlargement and positive arterial remodeling.⁶³ The performance of the ART18Z scaffold is currently being investigated in the clinical setting, in the “ARTDIVA” (Arterial Remodeling Transient Dismantling Vascular Angioplasty) FIM trial, which commenced at five clinical centers in the third quarter of the year 2012.

**Amaranth bioresorbable stent**

The Amaranth BRS (Amaranth Medical Inc, Mountain View, CA, USA) is made of PLLA, with proprietary tube fabrication. The structural integrity of the Amaranth scaffold lasts 3–6 months, and the resorption process of scaffold requires roughly 1–2 years. Preliminary experimental studies have shown that the scaffold has a high radial strength, prolonged mechanical stability, and exhibits minimal recoil.⁶⁴ A recent report evaluated the performance of the scaffold in porcine models compared with the Liberte BMS (Boston Scientific, Natick, MA, USA). A similar LLL was reported at 28- and 90-day follow-up. However, thereafter the LLL started to decrease in the Amaranth group, whereas this increased in the Liberte BMS group, from 90-day to 180-day follow-up. OCT examination at 90 days follow-up revealed a lower neointimal thickness in the Amaranth scaffold compared with the Liberte group. The company is currently designing the FIM study that will recruit 30 patients in order to evaluate the safety and efficacy of the Amaranth scaffold, with a plan for the conduction of a prospective, multicenter CE mark trial.

**Xinsorb BRS**

The Xinsorb BRS (Huaan Biotechnology Co, Ltd, Hangzhou, Zhejiang, People’s Republic of China) is a fully bioresorbable sirolimus-loaded scaffold, consisting of poly(aspartic acid-co-lactide), poly(ε-caprolactone), and polyglycolide. The device is balloon expandable, and has radiopaque markers to facilitate its deployment. The available scaffold size ranges from 2.75 to 4.0 mm in diameter, and from 12 to 28 mm in length.

A recent experimental study evaluated the feasibility and efficacy of the Xinsorb scaffold compared with the Excel DES (JW Medical Co, Shandong, People’s Republic of China).⁶⁵ Sixteen Xinsorb scaffolds and 16 Excel stents were implanted in the coronary arteries of porcine models.⁶⁶ Histomorphometry demonstrated a lower percentage diameter restenosis in the Xinsorb scaffold compared with
the Excel DES (at 30-day follow-up, 18.6% ± 5.2% versus 21.4% ± 7.2%, P > 0.05; at 90-day follow-up, 24.5% ± 4.7% versus 27.7% ± 5.6%, P > 0.05, respectively). Although these preliminary results are encouraging, further extensive preclinical studies are necessary to investigate the safety and efficacy of the scaffold. The company is expected to organize an FIM trial in 2013.

**Acute BRS**
The Acute BRS (OrbusNeich, Fort Lauderdale, FL, USA) is a tube-shaped lockable and balloon-expandable BRS. The device consists of three polymeric materials (poly-L-lactide-co-ε-caprolactone, PDLA, and PLLA) and incorporates a partitioned coating technology, which allows for the scaffold to be covered by abluminal sirolimus and luminal endothelial progenitor cell capture (+CD34) antibodies. Preliminary preclinical evaluation demonstrated an optimal device implantation without evidence of fracture.65 Updated data are expected to be presented in the EuroPCR Focus Group 2013.

**MeRes™ BRS**
MeRes BRS (Meril Life Sciences, Vapi, Gujarat, India) is a merilimus eluting bioresorbable coronary scaffold. The device is comprised of a new PLA formulation and has a hybrid scaffold geometry structure, which gives the scaffold a high radial strength. The performance of the device was examined in an experimental animal study. Initial results showed that the elution of the coated drug merilimus lasts more than 30 days, with no evidence of inflammatory reaction during the polymer’s biodegradation.64

**FADES BRS**
The FADES scaffold (Zorion Medical, Indianapolis, IN, USA) is a fully bioresorbable drug-eluting scaffold. The polymer of the scaffold involves a hybrid material of magnesium alloy that includes rare earth elements and PLGA. Preclinical studies showed that the device was completely absorbed with little to no inflammatory tissue response within 90 days.64 Further preclinical evaluations are needed before the conduction of the FIM study.

**Other BRSs**
Three other devices are under development: the Avatar BRS (S3V Vascular Technologies, Bangalore, Karnataka, India), the Sahajanand BRS (Sahajanand Medical Technologies, Surat, Gujarat, India), and Stanzat™ BRS (480 Biomedical, Boston, Massachusetts, USA). All these three BRSs are in their infancy, and preclinical and clinical evaluation is pending.64

**Conclusion**
BRSs introduce a unique potential in the treatment of coronary lesions, as they provide temporary vessel scaffolding, and then they disappear, thereby allowing for the restoration of the vessel wall physiology and vasomotion. Initial preclinical and clinical results appear promising. However, it should be mentioned that our clinical data are limited to simple lesions and there is no evidence in the context of randomized control trials that would allow direct comparison of the efficacy of the BRS with the effectiveness of the new generation DES. Furthermore, BRS technology has significant limitations such as the bulky nature of the devices, the fact that they cannot be implemented to treat bifurcation lesions requiring a two stent strategy, the concerns about an increased risk of restenosis in case of overlapping, and their limited extensibility and the risk of scaffold fracture that limit their applications in the current clinical practice. Therefore further development is necessary so as to reduce the thickness of the struts and increase the extensibility of the devices so as to overcome the current pitfalls of BRS technology and become the workhorse device for the treatment of CAD.

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