Current Management of In-Stent Restenosis

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ABSTRACT

Despite the advent of the drug-eluting stents (DES) and improved stent design, in-stent restenosis (ISR) remains a challenging problem. The currently available options for treatment of ISR include angioplasty alone, repeat stenting with DES or drug-coated balloons. Several recent studies have compared the available options for treating ISR in an attempt to identify the preferred therapeutic strategy. In this review, we will discuss the currently available therapeutic strategies for the management of patients with ISR and the evidence supporting their use.

Keywords: Restenosis; Angioplasty; Balloon; Coronary; Drug-eluting stent

INTRODUCTION

Although the use of coronary stents brought about a dramatic improvement in patients’ clinical and procedural outcomes, the long-term outcome of stent implantation remains significantly constrained by the risk of developing in-stent restenosis (ISR) over time. ISR, which is currently defined as a >50% stenosis of a previously stented segment, occurs in as many as 30% of all patients receiving bare metal stents (BMS). Despite the advent of the drug-eluting stents (DES) and improved stent design, the rates of ISR in patients treated with DES are as high as 10%. Specifically, the widespread adoption of DES for small arteries, long lesions, complex coronary lesions, diabetes, and a history of bypass surgery has in fact been the trigger for significant numbers of patients re-presenting with DES restenosis in contemporary clinical practice.

The treatment of patients with ISR continues to remain a challenge, and currently available options include angioplasty alone, repeat stenting with DES or drug-coated balloons (DCB). Recent meta-analyses have performed comparisons of these available options in an attempt to identify the preferred therapeutic modality. Therefore, this review will discuss the currently available therapeutic strategies for the management of patients with ISR and the evidence supporting their use.
DEFINITION AND CLASSIFICATION OF ISR

ISR is defined as the gradual re-narrowing of a stented coronary artery lesion due to arterial damage with subsequent neointimal tissue proliferation. The angiographic definition of ISR remains a binary event defined as a stenosis within the stented segment or its edge (5-mm segments adjacent to the stent) of >50% of the vessel diameter as determined by coronary angiography. The clinical definition of ISR requires the presence of >50% diameter in-stent stenosis and one of the following: clinical symptoms of recurrent angina, objective signs of ischemia (electrocardiography changes), positive coronary hemodynamic assessment with fractional flow reserve (FFR) <0.80, intravascular ultrasound (IVUS) minimum cross-sectional area <4 mm² (6 mm² for left main), or restenosis with ≥70% reduction in lumen diameter even in the absence of clinical symptoms or signs.

The Mehran system is a morphological classification created for BMS-ISR lesions (pattern I: focal, pattern II: diffuse, pattern III: proliferative, and pattern IV: occlusive) which can help to predict the need for repeat revascularization (19%, 35%, 50%, and 98%, respectively). This classification scheme has also been shown to have prognostic value in DES-ISR. Additionally, the American College of Cardiology/American Heart Association classification has been validated in patients with ISR: lesions B2 and C are more frequently associated with suboptimal acute results; a higher restenosis rate; and poorer long-term clinical outcomes.

CHARACTERIZATION AND PRESENTATION OF ISR

Mounting evidence strongly suggests that there are significant differences between ISR in BMS and DES, with the main disparities, time of presentation, morphological patterns, underlying substrate, and the response to interventions. The time course of neointimal accumulation differs considerably between DES and BMS, which is a manifestation of the delayed arterial healing that appears to characterize the vascular response to DES implantation. Moreover, compared with BMS-ISR, DES-ISR tends to be focal, particularly at the stent edge or in areas of stent fracture. Lack of diffuse neointimal hyperplasia in DES may be due to the high overall suppression of neointimal growth by DES unless there is mechanical stent failure. In addition, focal neoatherosclerosis occurs not only more frequently, but also significantly earlier in DES-ISR compared with BMS-ISR.

Assessing the underlying etiology for ISR is critical for guiding and optimizing repeat interventions to prevent repeated ISR. The most well recognized and preventable cause for ISR is stent under-expansion, and this is considered a major factor triggering ISR after either BMS or DES implantation. This problem may be due to stent under-sizing, low deployment pressures, or extensive vessel calcification leading to stent under-deployment or under-expansion. Stent misplacement or stents not fully covering the underlying lesion are other important risk factors for ISR. Geographic miss leaves a characteristic “candy-wrapper” angiographic appearance at the edge of a stented segment and is thought to lead to ISR because of stent-related edge dissection, poor endothelialization and subsequent proliferation of the atherosclerotic plaque burden. Stent fractures may also trigger focal ISR as they cause similar problems to edge disease except within the stent. Finally, the adoption of DES has led to the recognition of drug resistance and local hypersensitivity reactions as another possible cause of ISR.
Intracoronary imaging, which can be performed with either IVUS or optical coherence tomography (OCT), plays an important role in evaluating the potential mechanism of ISR.\(^\text{17,20}^\) IVUS can detect the presence of neointimal hyperplasia within the stent, stent under-expansion, stent fracture or edge restenosis, and the borders of the external elastic lamina for vessel sizing enabling optimization of stent expansion.\(^\text{27}\) However, due to its superior axial resolution (15 µm), OCT provides better detailed images of the vessel-lumen interface, the neointimal tissue, and the strut distribution.\(^\text{11,18}\) It has enabled more detailed evaluation of the ISR etiology and has highlighted the morphologic differences between BMS- and DES-ISR. OCT in BMS-ISR typically shows a homogeneous high-signal tissue band, which is characteristic of neointimal hyperplasia rich in smooth muscle cells.\(^\text{11}\) In contrast, DES-ISR is typically characterized by a focal, heterogeneous and layered intrastent tissue band, which represents hypocellular neointima with high proteoglycan or fibrin content which likely occurs in the setting of neoatherosclerosis. Specific findings that are also suggestive of neoatherosclerosis include neointimal rupture, thin-cap fibroatheroma, lipid pools, macrophage accumulation, and evidence of non-occlusive thrombosis.\(^\text{20}\)

Although ISR had traditionally been thought to represent a relatively benign clinical entity with predominantly stable clinical presentation, more recent studies suggest that a significant number of patients with ISR present with acute coronary syndrome.\(^\text{21,22}\) This acute clinical presentation is likely to be related to the neoatherosclerotic process described for DES, which is more likely to follow the typical atherosclerotic cascade of coronary occlusion secondary to neoatherosclerotic plaque rupture and thrombus formation.\(^\text{23}\) It is possible that late stent thrombosis is just a step in the continuum of the neoatherosclerotic process seen in DES-ISR. Conversely, the natural history of asymptomatic patients with angiographic restenosis is favorable.\(^\text{24}\) Therefore, treatment of asymptomatic patients (oculostenotic reflex) should be avoided whenever possible.\(^\text{25,27}\) Similar to de novo lesions, the functional significance of ISR should be assessed using a pressure wire. Prospective studies have validated the use of FFR for clinical decision making in ISR, and have found that deferring revascularization in patients with an FFR of >0.75 is safe and appropriate.\(^\text{28,29}\)

**TREATMENT OF ISR**

**Medical/surgical treatment**

There is little evidence to support medical treatments for ISR. Although abciximab was considered to be of particular value in patients with ISR in early studies, larger trials failed to confirm any clinical benefit.\(^\text{30,31}\) Similarly, oral sirolimus was initially considered to be of potential value in these patients, however, the lack of long-term efficacy and the higher incidence of adverse drug effects have shown it to be a poor option.\(^\text{32,33}\) Finally, coronary surgery may be considered in patients with recalcitrant ISR, particularly in those with a diffuse ISR pattern or associated significant disease in other major vessels.\(^\text{35}\)

**Plain old balloon angioplasty (POBA)**

POBA is one of the earliest treatments that has been used in patients with ISR.\(^\text{34}\) The procedure is technically straightforward and is consistently associated with satisfactory acute results and a very low incidence of complications.\(^\text{39}\) The immediate angiographic improvement following POBA results from both axial and longitudinal tissue extrusion as well as further stent expansion.\(^\text{36,37}\) Results are particularly favorable in patients with a focal pattern of ISR and when stent/native artery size mismatch has been identified with
in intravascular imaging. High-pressure balloon dilatation and the use of a non-compliant balloon is often necessary to obtain optimal results. In general, a balloon to artery ratio of 1.1 to 1 is recommended for sizing when treating ISR. One of the limitation of POBA is that sub-acute tissue re-intrusion back to the lumen tends to occur within minutes of the last balloon inflation. This explains the “early lumen loss” phenomenon detected in POBA studies in ISR, a finding also associated with subsequent recurrent restenosis. Additionally, edge-related complications should be carefully avoided during aggressive balloon dilations. Balloon slippage outside the stent (“water-melon seeding” phenomenon), which occurs more often in severe and diffuse narrowing when balloons are oversized, can lead to edge dissections and suboptimal outcomes. Progressive balloon upsizing as well as the use of short low profile balloons can help avoid this phenomenon and edge-related complications.

**Cutting and scoring balloon therapy**

The cutting balloon is an attractive and simple technique for treatment of ISR. Theoretically, the device deeply incises neointimal tissue and may favor its subsequent extrusion. The lateral blades of the device anchor the balloon within the target lesion, preventing balloon slippage-related complications. Initial observational data suggested that cutting balloons may have superior efficacy compared to POBA, a finding which was associated with a lower rate of target lesion revascularization (TLR) (12.5% vs. 40%) at follow-up. However, in the largest randomized trial (Restenosis Cutting Balloon Evaluation Trial [RESCUT]), cutting balloon angioplasty comparing POBA failed to show an improvement in angiographic restenosis or in the rate of clinical events at late follow-up. Cutting balloon angioplasty was associated however with the need to use fewer balloons, less additional stenting, and a lower rate of balloon slippage (6.5% vs. 25%).

Scoring balloons are based on the same principle as cutting balloons but are especially attractive in patients with ISR due to their superior flexibility and deliverability. The Intracoronary Stenting and Angiographic Results: Optimizing Treatment of Drug-Eluting Stent In-Stent Restenosis 4 (ISAR-DESIRE IV trial) assessed the use of scoring balloons prior to DCB treatment of DES-ISR. The results showed superior angiographic outcomes at 6 to 8 months in the scoring balloon arm, but failed to show any significant difference in clinical outcomes.

**Debulking techniques**

Debulking techniques such as directional/rotational atherectomy and excimer laser are a novel treatment for ISR through their physical removal of neointimal tissue or neoatherosclerotic plaque. It was believed that after the initial removal of excess stenotic tissue by the debulking device, just a low-pressure balloon post-dilation is required to avoid additional vessel wall injury. Early observational studies suggested that the use of laser or rotational atherectomy, followed by a POBA post-dilation, was superior to conventional POBA alone in ISR. Directional atherectomy was also assessed in early studies, but this was soon abandoned because it was not well suited for small or distal vessels, which are common locations for ISR. The excimer laser showed good results in some cases but eventually proved to have poorer ablation capability compared with rotational atherectomy.

The value of rotational atherectomy in patients with BMS-ISR was evaluated in 2 randomized trials. In Randomized Trial of Rotational Atherectomy Versus Balloon Angioplasty for Diffuse In-Stent Restenosis (ROSTER), rotational atherectomy reduced the amount of residual tissue within the stent and the rate of TLR at follow-up, compared with POBA alone. On the other hand, in the Angioplasty Versus Rotational Atherectomy for Treatment of Diffuse In-Stent...
Restenosis Trial (ARTIST), which compared rotational atherectomy with POBA alone, lower restenosis rates, an improved safety profile and superior clinical outcomes were seen in the POBA group. Recently, the value of debulking techniques in patients with DES-ISR has been re-evaluated with the latest study showing greater acute luminal gain after percutaneous coronary intervention with excimer laser atherectomy. Reassuringly, even though excimer laser atherectomy was used for DES-ISR in significantly more complex lesions, the long-term clinical outcomes were favorable. Therefore, although debulking techniques are not considered to be a routine treatment of ISR, they can be considered as a pre-treatment option for undilatable ISR lesions, especially those as a result of severely under-expanded stents or calcified intrastent neoatherosclerosis.

**Vascular brachytherapy**

Brachytherapy was one of the most promising treatment options for patients with neointimal hyperplasia related to BMS-ISR. It involved temporary intracoronary deposition of a radioactive isotope within the diseased segment, which led to significantly reduced clinical and angiographic restenosis rates. Randomized clinical trials in patients with ISR showed it to be more effective in preventing ISR progression and improving clinical outcomes than either POBA or debulking procedures with laser or atherectomy. However, the advent of DES signaled the end of brachytherapy. The 2 large randomized clinical trials which compared the efficacy of brachytherapy versus DES in patients with BMS-ISR were Sirolimus-Eluting Stents versus Vascular Brachytherapy for In-Stent Restenosis Within Bare-Metal Stents (SISR) and Paclitaxel-Eluting Stents versus Vascular Brachytherapy for In-Stent Restenosis Within Bare-Metal Stents (TAXUS V ISR). Both showed that DES were superior in decreasing restenosis rates and the need for revascularization as compared to brachytherapy at long-term follow-up. Disappointingly whilst observational studies of DES-ISR suggested a role for brachytherapy, no randomized trials comparing it to DES or DCB therapy have ever been conducted. Finally, the complexity of the procedure, as well as issues with radioprotection/radiation dosing, led to the virtual abandonment of this strategy.

**Repeat stenting with BMS**

Early studies suggested that the problem of early tissue loss, which was seen with POBA, was virtually eliminated with the use of BMS, which gave credence to the possible superiority of stenting over POBA in the treatment of ISR. In BMS-ISR, IVUS studies also demonstrated that repeat stenting was the best strategy to obtain a larger acute lumen gain and better immediately results post procedure. In the Restenosis Intra-stent Balloon Angioplasty Versus Elective Stenting (RIBS I) trial, patients with BMS-ISR, were randomized to receive either POBA or repeat BMS implantation, with acute angiographic results being significantly better after BMS placement due to a larger acute gain. However, at 6-month follow-up, significant late lumen loss in the BMS group resulted in the final angiographic appearance being similar in both groups. To date, large randomized trials assessing the value of BMS in patients with DES-ISR are lacking.

**Repeat stenting with DES**

In de novo lesions, DES produce a profound inhibition of neointimal proliferation. Therefore, the use of DES has become an attractive option in the treatment of neointimal hyperplasia in BMS-ISR. The Intracoronary Stenting or Angioplasty for Restenosis Reduction-Drug-Eluting Stents for In-Stent Restenosis (ISAR-DESIRE) trial was the first randomized study assessing the value of DES in patients with BMS-ISR. The rate of recurrent restenosis was significantly lower with sirolimus- (14.3%) and paclitaxel-DES.
(21.7%) compared with POBA alone (44.6%). Similar results were also shown in a subsequent meta-analysis comparing these 2 DES for BMS-ISR. In the Restenosis Intrastent: Balloon Angioplasty Versus Elective Sirolimus-Eluting Stenting (RIBS II) trial, which compared sirolimus-DES versus POBA in patients with BMS-ISR, patients with sirolimus-DES had a significantly lower restenosis rate (11%) and superior long-term clinical outcomes. In addition, IVUS imaging confirmed the dramatic reduction of neointimal proliferation seen after the use of sirolimus-DES. The 4-year long-term follow-up study demonstrated a sustained clinical benefit from DES placement without any significant increase in major adverse cardiac events (MACE).

Unfortunately, the treatment of DES-ISR is more challenging, and overall, the outcomes in patients requiring treatment for DES-ISR are worse compared with patients with BMS-ISR. Early observational studies suggested that DES provided superior results compared with other strategies such as POBA or cutting balloon angioplasty. Investigators have proposed that DES-ISR that results from a mechanical complication (such as stent under-sizing, edge dissection or stent fracture) can be successfully overcome by placing another DES. However, debate regarding whether to use a DES eluting the same or a similar type of drug (homo-DES approach) versus a switch to a different type of drug (hetero-DES approach) has continued. The benefits of a switch approach are based on the hypothesis that it might overcome drug resistance or polymer-related problems. In Intracoronary Stenting and Angiographic Results: Drug Eluting Stents for In-Stent Restenosis 2 (ISAR-DESIRE 2) trial of sirolimus-DES-ISR, the hetero-DES strategy using a paclitaxel-DES failed to reduce restenosis or target vessel revascularization rates compared to repeat stenting with sirolimus-DES. The Restenosis Intra-Stent: Balloon Angioplasty vs Drug-Eluting Stent (RIBS III) trial also compared the DES-switch approach to same-stent implantation. Although there was no significant difference between the hetero-DES and homo-DES approach, the study suggested that the use of second-generation DES was superior to first-generation DES, and intravascular imaging for treatment guidance had improved angiographic and clinical outcomes. Despite these benefits of repeat stenting with DES in the management of DES-ISR, current data suggests that 10–20% of these patients will go on to develop recurrent ISR.

Bioresorbable vascular scaffolds (BVS)
BVS have also been proposed as treatment for patients with ISR. The main advantages are that the device eventually disappears from the vessel wall, avoiding the presence of multiple stent layers, and prevents early lumen loss associated with tissue retraction seen in balloon angioplasty. Some small studies have established that BVS placement in the treatment of ISR is safe and feasible. In 65 patients with ISR treated with BVS, clinical outcomes at 1 year revealed a TLR rate of about 12%, and all of these patients avoided having a permanent second layer of stent struts. Nevertheless, since no randomized trial evaluating the effectiveness of BVS in management of ISR has yet been performed, the routine use of this strategy cannot be recommended.

DCB
The development of DCB enabled deliver of anti-proliferative drug to the area of ISR without leaving behind an additional layer of stent strut. Although the value of DCB in de novo lesions remains controversial, the use of DCB has been to proven to be very effective in patients with both BMS-ISR and DES-ISR (Table 1). The initial study of BMS-ISR demonstrated that DCB were superior to POBA alone. The 6-month angiographic results were significantly improved in the DCB group (late loss: 0.33 mm vs. 0.74 mm, p<0.002). The subsequent
larger randomized study compared paclitaxel-DES placement to paclitaxel-coated balloon application in BMS-ISR. At 6-month follow-up, DCB significantly reduced the primary endpoint of the study (angiographic late loss: 0.17 mm vs. 0.38 mm, p=0.03), although minimal lumen diameter and diameter stenosis were similar in both arms.73 Recently, the Restenosis Intra-stent: Drug-eluting Balloon vs. Everolimus-eluting Stent (RIBS V) trial conducted a randomized comparison of DCB with second-generation everolimus-DES in patients with BMS-ISR.74 This study showed better late angiographic findings in the DES arm (minimal lumen diameter: 2.01 mm vs. 2.36 mm, p<0.001), but showed similar rates of restenosis and clinical outcomes. Therefore, the overall non-inferior outcomes with DCB treatment as compared with DES placement in several studies seem to support the use of DCB for treatment of BMS-ISR, especially in situations where additional stent layers are undesirable or bleeding events.

The value of DCB in patients with DES-ISR has also been assessed. An initial small randomized study demonstrated that in patients with DES-ISR, DCB provided superior clinical and angiographic results compared with POBA alone (late lumen loss: 0.18 mm vs. 0.38 mm, p=0.001).22 The efficacy of DCB in patients with DES-ISR was subsequently confirmed in a multicenter, randomized trial including patients with any type of DES-ISR (late loss: 0.43 mm vs. 1.03 mm vs. 2.36 mm, p<0.001), but showed similar rates of restenosis and clinical outcomes. Therefore, the overall non-inferior outcomes with DCB treatment as compared with DES placement in several studies seem to support the use of DCB for treatment of BMS-ISR, especially in situations where additional stent layers are undesirable or bleeding events.

The value of DCB in patients with DES-ISR has also been assessed. An initial small randomized study demonstrated that in patients with DES-ISR, DCB provided superior clinical and angiographic results compared with POBA alone (late lumen loss: 0.18 mm vs. 0.72 mm, p=0.002 at 6 months)

### Table 1. Randomized clinical trials of DCB on treatment of ISR

<table>
<thead>
<tr>
<th>Author/trial</th>
<th>Previous stent</th>
<th>Treatment</th>
<th>Number</th>
<th>Angiographic follow-up</th>
<th>Clinical follow-up</th>
</tr>
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<tbody>
<tr>
<td>Scheller et al.72</td>
<td>BMS</td>
<td>DCB vs. POBA</td>
<td>52</td>
<td>LLL: 0.03±0.48 mm (DCB) vs. 0.74±0.86 mm (POBA), p=0.002 at 6 months</td>
<td>MACE: 4% (DCB) vs. 33% (POBA), p=0.01 at 12 months</td>
</tr>
<tr>
<td>Unverdorben et al.73</td>
<td>BMS</td>
<td>DCB vs. DES-P</td>
<td>131</td>
<td>LLL: 0.17±0.42 mm (DCB) vs. 0.38±0.61 mm (DES), p=0.03 Binary restenosis rate: 7% (DCB) vs. 20% (DES), p=0.06 at 6 months</td>
<td>MACE: 9% (DCB) vs. 22% (DES), p=0.08 at 12 months</td>
</tr>
<tr>
<td>RIBS V74</td>
<td>BMS</td>
<td>DCB vs. DES-E</td>
<td>189</td>
<td>MLD: 2.01±0.60 mm (DCB) vs. 2.36±0.60 mm (DES), p=0.001 at 9 months</td>
<td>MACE: 8% (DCB) vs. 6% (DES), HR: 0.76, p=0.60 at 12 months</td>
</tr>
<tr>
<td>Habara et al.22</td>
<td>DES</td>
<td>DCB vs. POBA</td>
<td>50</td>
<td>LLL: 0.18±0.45 mm (DCB) vs. 0.72±0.55 mm (POBA), p=0.001 Binary restenosis rate: 8.7% (DCB) vs. 62.5% (POBA), p=0.001 at 6 months</td>
<td>MACE: 4% (DCB) vs. 40% (POBA), p=0.005 at 12 months</td>
</tr>
<tr>
<td>PEPCAD-DES75</td>
<td>DES</td>
<td>DCB vs. POBA</td>
<td>110</td>
<td>LLL: 0.43±0.61 mm (DCB) vs. 1.03±0.77 mm (DES), p=0.001, Binary restenosis rate: 17.2% (DCB) vs. 58.1% (POBA), p=0.001 at 6 months</td>
<td>MACE: 17% (DCB) vs. 50% (POBA), p=0.001 at 12 months</td>
</tr>
<tr>
<td>PEPCAD China ISR76</td>
<td>DES</td>
<td>DCB vs. DES-P</td>
<td>220</td>
<td>LLL: 0.46±0.51 mm (DCB) vs. 0.55±0.61 mm (DES), p for noninferiority=0.001 at 9 months</td>
<td>TLF: 17% (DCB) vs. 16% (DES), p=0.52 at 12 months</td>
</tr>
<tr>
<td>ISAR-DESIRE 370</td>
<td>DES</td>
<td>DCB vs. DES-P</td>
<td>402</td>
<td>DS: 38.0±21.5% (DCB) vs. 37.4±21.7% (DES) vs. 54.1±25.0% (POBA), p for noninferiority=0.007 (DCB vs. DES), p for superiority=0.001 (other vs. POBA) at 6–8 months</td>
<td>MACE: 24% (DCB) vs. 19% (DES) vs. 46% (POBA)</td>
</tr>
<tr>
<td>RIBS IV78</td>
<td>DES</td>
<td>DCB vs. DES-E</td>
<td>309</td>
<td>MLD: 1.80±0.60 mm (DCB) vs. 2.03±0.70 mm (DES), p=0.01 at 6–9 months</td>
<td>MACE: 18% (DCB) vs. 10% (DES), HR: 0.36, p=0.04 at 12 months</td>
</tr>
</tbody>
</table>

BMS = bare-metal stents; DCB = drug-coated balloons; DES = drug-eluting stents; DES-P = paclitaxel drug-eluting stents; DES-E = everolimus drug-eluting stents; POBA = plain old balloon angioplasty; TLF = target lesion failure.
Finally, whether DCB proves comparable to repeat stenting with second-generation DES in patients with DES-ISR remains controversial. In the recently published Restenosis Intra-stent of Drug-eluting Stents: Paclitaxel-Eluting Balloon vs Everolimus-Eluting Stent (RIBS-IV) trial which compared second-generation everolimus-eluting DES to DCB for treatment of DES-ISR, both angiographic and clinical outcomes favored second-generation DES over DCB (minimal lumen diameter: 2.03 mm vs. 1.80 mm, p<0.01; MACE: 10% vs. 18%, p=0.04) at 6- to 9-month follow-up. In addition, whether the efficacy of DCB can be further improved by optimal lesion preparation with scoring/cutting balloon remains unknown and the ongoing ISAR-DESIRE 4 randomized trial will address this issue.

Comparison of all treatment modalities
Two recent large meta-analyses were conducted to clarify which strategy is the best treatment modality for ISR. Siontis et al. included 27 trials with a total of 5,923 patients at 6 months to 1 year follow-up. The primary outcome of this analysis was percent diameter stenosis at follow-up, and the secondary endpoint included binary restenosis, rates of TLR, myocardial infarction or death. All modalities included POBA alone, debulking techniques, brachytherapy, BMS, DES, and DCB. Repeat stenting with everolimus-DES was found to be statistically superior to all other modalities for both the primary outcome as well as for binary restenosis rates and TLR. DCB appeared to be the second most preferable treatment but did not achieve a significant difference over sirolimus or paclitaxel-eluting stents. Giacoppo et al. included 24 trials with a total of 4,880 patients, and the primary outcomes were TLR rates and angiographic late lumen loss. Both DCB and DES were superior to other treatment modalities based on the predefined clinical outcomes. Angiographic outcomes favored DCB or DES over all other modalities, however late lumen loss appeared to be slightly lower in the DCB arm compared with DES.

CONCLUSION
Although the development of DES has reduced the incidence of ISR, treatment of ISR remains a challenging clinical problem. Current clinical data suggest that among various available therapeutic modalities, second-generation DES and DCB provide the best clinical and angiographic results in patients with ISR. Implantation of more than 2 metal stents in repeated ISR lesions is likely to have a detrimental effect on long-term outcomes, even though newer DES may improve the treatment of ISR lesions. Further studies are required to clarify the role of these current therapeutic modalities which may help improve clinical outcomes in those with ISR.

REFERENCES


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