DANUBIO - a new drug-eluting balloon for the treatment of side branches in bifurcation lesions: six-month angiographic follow-up results of the DEBSIDE trial

Jacques Berland1*, MD; Thierry Lefèvre2, MD, FESC, FSCAI; Philippe Brenot3, MD; Jean Fajadet4, MD; Pascal Motreff5, MD, PhD; Patrice Guerin6, MD, PhD; Patrick Dupouy7, MD; Christian Schandrin8, MD; DEBSIDE trial investigators

1. Clinique Saint-Hilaire, Rouen, France; 2. Institut Hospitalier Jacques Cartier, Massy, France; 3. Centre Cardiologique d’Evequeumont, Evequeumont, France; 4. Clinique Pasteur, Toulouse, France; 5. Hôpital Gabriel Montpied, Clermont-Ferrand, France; 6. Hôpital Guillaume et René Laennec, Nantes, France; 7. Clinique les Fontaines, Melun, France; 8. Pôle Santé République, Clermont-Ferrand, France

Abstract

Aims: We aimed to evaluate the role of drug-eluting balloon SB inflation, using the novel DANUBIO balloon, after placement of a drug-eluting stent in the main branch in patients with bifurcation lesions.

Methods and results: Fifty-two patients with bifurcation lesions suitable for stenting were enrolled in the DEBSIDE trial at eight French centres between May 2012 and July 2013. Two patients were excluded from the trial because of significant protocol deviations. Systematic Nile PAX stent placement was followed by final drug-eluting balloon inflation, using the DANUBIO balloon, according to the size of the side branch. Clinical follow-up was scheduled at one, six, and twelve months and an angiographic control at six months. The primary endpoint was six-month late lumen loss (LLL) at the ostium of the side branch. Secondary endpoints were main branch (MB) LLL, binary restenosis of the SB and MB, and clinically driven revascularisation rates for both branches. The procedural success rate was 100%. Angiographic control at six months post-procedure was performed in 48 patients (96%). Two patients with no reported clinical events refused the angiographic control. At six-month follow-up the primary endpoint of side branch LLL was -0.04±0.34 mm and the secondary endpoint of MB LLL was 0.54±0.60 mm. There was only one myocardial infarction (2%) and no reported cardiac deaths. Only one patient (2%) had a non-clinically driven target lesion revascularisation (TLR) at the level of the side branch combined with a main branch revascularisation.

Conclusions: Systematic final inflation of a DANUBIO balloon in the side branch after placement of a Nile PAX stent in the main branch for the treatment of a bifurcation lesion is safe and effective and results in very low LLL and a low restenosis rate at the side branch ostium. The DEBSIDE clinical trial was registered at the United States National Institute of Health website (NCT01485081).

*Corresponding author: Cardiology Department, Clinique Saint-Hilaire, 2 place Saint-Hilaire, 76000 Rouen, France. E-mail: jberland@clinique-sainthilaire.fr
Abbreviations

BTHC butyryl tri-n-hexyl citrate
CEC clinical events committee
DEB drug-eluting balloon
DES drug-eluting stent
DM distal main branch
DS diameter stenosis
LLL late lumen loss
MACE major adverse cardiac events
MB main branch
MBC main branch catheter
MI myocardial infarction
MLD minimum lumen diameter
PCI percutaneous coronary intervention
PM proximal main branch
POBA plain old balloon angioplasty
QCA quantitative coronary angiography
RVD reference vessel diameter
SB side branch
SBC side branch catheter
TLR target lesion revascularisation
TVR target vessel revascularisation

Introduction

Coronary bifurcation lesions continue to be considered complex coronary lesions to treat. The introduction of drug-eluting stents (DES) has lowered the rate of restenosis in these lesions 1. Even so, in the recent consensus document of the European Bifurcation Club, the complexity of treatment was again underlined, calling it "one of the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events" 2, and the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events 2, and the complexity of treatment was again underlined, calling it "one of the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events" 2, and the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events" 2, and the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events" 2, and the most challenging lesions in interventional cardiology in terms of procedural success rate as well as long-term cardiac events. The balloon surface coating is an L605 cobalt-chromium (Co-Cr) stent and is designed to be used in a final kissing post-dilatation to optimise deployment. The Nile PAX allows an optimal provisional T-stenting approach. The device has been evaluated in the BIPAX trial 11 (NCT01308229). Further to these preclinical trial results, the DANUBIO obtained CE mark approval in October 2011.

The DANUBIO study device was used in conjunction with the Nile PAX, a polymer-free abluminal-coated paclitaxel-eluting coronary stent dedicated to bifurcation lesions. The stent platform is an L605 cobalt-chromium (Co-Cr) stent and is designed with a single link at the level of the carina to prevent side branch obstruction. The delivery system is 6 Fr compatible and is composed of two parallel rapid exchange (RX) balloon catheters, the main branch catheter (MBC) and side branch catheter (SBC), which can be used in a final kissing post-dilatation to optimise deployment. The Nile PAX allows an optimal provisional T-stenting approach. The device has been evaluated in the BIPAX trial 11 (NCT01308229).

Methods

STUDY DESIGN

The DEBSIDE clinical trial was a prospective, non-randomised, multicentre, interventional study evaluating the investigational DANUBIO paclitaxel-eluting coronary balloon for the treatment of side branches of de novo bifurcation lesions with a side branch reference vessel diameter ≥2.0 mm and ≤3.0 mm.

STUDY DEVICE

The DANUBIO paclitaxel-eluting PTCA balloon catheter (Minvasys) is a new technology developed in order to treat coronary artery disease and especially restenosis. The balloon surface coating is an L605 cobalt-chromium (Co-Cr) stent and is designed with a single link at the level of the carina to prevent side branch obstruction. The delivery system is 6 Fr compatible and is composed of two parallel rapid exchange (RX) balloon catheters, the main branch catheter (MBC) and side branch catheter (SBC), which can be used in a final kissing post-dilatation to optimise deployment. The Nile PAX allows an optimal provisional T-stenting approach. The device has been evaluated in the BIPAX trial 11 (NCT01308229).

PATIENT POPULATION

Between May 2012 and July 2013, 52 patients with de novo bifurcation lesions (except Medina class 0,0,1 type lesions) were treated with the DANUBIO drug-eluting balloon in eight clinical centres in France. Inclusion criteria for enrolment in the trial included male and non-pregnant female patients over 18 years of age with clinical evidence of ischaemic heart disease, stable or unstable angina with signs of ischaemia, silent ischaemia, or a positive functional test. Lesion length had to be ≤6 mm in the side branch and the reference vessel diameter had to be between 2.5 and 3.5 mm for the MB and 2.0 and 3.0 mm for the SB. Exclusion criteria included previous percutaneous coronary intervention (PCI) of the target vessel within six months prior to the procedure, left main bifurcation and acute myocardial
infarction within 72 hours. Two patients were excluded post-procedure because of significant protocol deviations: one patient had a previous PCI at the target lesion and the other had a side branch lesion longer than 10 mm (Figure 1).

All patients provided written informed consent prior to enrolment and procedure. The study complied with the Declaration of Helsinki regarding investigation in humans, and was approved by the local ethics committee.

PROCEDURE
First, a sequential main branch and side branch predilatation was performed with a conventional balloon. If the angiographic result was good after predilatation, a dedicated drug-eluting stent (Nile PAX) was deployed in the main branch of the bifurcation site. The stent diameter was selected based on the diameter of the distal MB. A recommended kissing balloon procedure was then performed with the MBC and SBC of the Nile PAX delivery system. After that, the guidewire in the main branch was removed in order to prevent any drug loss when the DEB was placed into the side branch. Indeed, as already reported by Afari et al, the active coating is subject to erosion during tracking to the target lesion, notably in the guiding catheter. The use of a second wire is a potential cause of increased friction resulting in loss of coating.

If there was no dissection impeding the flow in the side branch, then the DEB (DANUBIO balloon catheter) was inflated only once in the side branch for 30 seconds at low pressure (8-10 bar). The procedure was performed based on the recommendations of the German consensus group13. As defined in the protocol, the DEB diameter could not exceed in any way the SB diameter and had to match the Nile PAX SBC diameter. In addition, the balloon length had to extend at least 2 mm beyond the side branch lesion distally in order to avoid geographical mismatch.

Patients then received dual antiplatelet therapy (DAPT) for at least 12 months.

ANGIOGRAPHIC ANALYSIS
Serial coronary angiographic studies were obtained at baseline (pre-procedure) and post-procedure after intracoronary administration of nitrate (recommended 100-200 μg, unless clinically contraindicated) in two orthogonal matching views. For the study, the dedicated bifurcation algorithm from the CAAS 5.10 system (Pie Medical Imaging BV, Maastricht, The Netherlands) was used. All angiographic analyses were performed by an independent angiographic core laboratory (European Cardiovascular Research Center [CERC], Massy, France). A single analysis was performed after placing boundaries for the proximal and distal main branch and side branch segments. The worst analysable view was selected. Automatic contour detection was used; manual editing was restricted to cases where the contour detection was not following the vessel. Both main and side branch reference diameters were automatically interpolated by the software using the automatic segmentation. Both six-segment and 11-segment models were analysed; however, due to the limitations related to the role of the point of convergence (segment 7) in the 11-segment models, data are presented according to the six-segment model as proposed by Ramcharitar et al14 and Girasis et al15. The minimum lumen diameter (MLD) at the ostium of the side branch was evaluated at the level of the ostium in the first 5 mm of the proximal portion of the side branch. The late lumen loss (LLL) was calculated as usual: LLL = MLD post-procedure - MLD at six months.

STUDY ENDPOINTS
The primary endpoint of the study was the evaluation of LLL at the ostium of the side branch at six months by quantitative coronary angiography (QCA) analysis. Secondary endpoints were angiographic success, LLL in the main branch, angiographic binary restenosis (ABR) rate (%) in the side and main branches, and major adverse cardiac events (MACE: cardiac death, myocardial infarction [MI] and target lesion revascularisation [TLR] in the MB and SB). The procedure success was also measured (success defined as achieving a stenosis of less than 50% with no clinical or ischaemic events during the initial hospitalisation). Clinical follow-up was scheduled at one, six and twelve months, and patients had systematic six-month angiographic analysis. A clinical events committee (CEC) adjudicated all clinical events.

STATISTICAL ANALYSIS
Statistical analysis was performed by an independent core laboratory (CERC). All patients/lesions were analysed on a per protocol basis. The continuous variables are described as mean±standard deviation, and discrete variables with frequencies and percentages. All analyses were carried out using SAS software, version 9.4 (SAS Institute Inc., Cary, NC, USA).

The Student’s t-test was used for the comparisons analysis between results obtained before and after the procedure and at six-month follow-up. A p-value <0.05 was considered statistically significant.

Figure 1. Patient CONSORT-type flow chart.
Results

BASELINE CHARACTERISTICS

Patients’ clinical demographics are presented in Table 1.

The clinical profile of the 50 patients included in the protocol analysis was typical of a population suffering from coronary artery disease with an average age of 66 years, 78% male, 26% rate of diabetes and a previous history of ischaemic heart disease in 40% of patients. Fifty percent of patients were clinically stable, either with stable angina or with silent ischaemia, whereas 50% of patients were unstable, with 28% STEMI or NSTEMI. The most frequently treated bifurcation involved the LAD and a diagonal artery (diagonal 1 in 30 patients [60%], diagonal 2 in five patients [10%], followed by marginal circumflex artery 1 and 2 in nine patients [18%], and the right posterior lateral artery/right descending posterior artery (RPL/RDP) bifurcation in the right coronary artery (RCA) in four patients [8%]).

The bifurcation lesions, described according to the Medina classification based on visual estimation by the investigators (Figure 2), comprised 16 type 1,1,1 lesions, 13 type 1,0,1 lesions, and three type 0,1,1 lesions.

In total, 32 patients (64%) presented with a visually significant lesion of the side branch. For the 18 other patients, the distribution of lesions was as follows: 12 type 1,1,0 lesions, five type 1,0,0 lesions, and one type 0,1,0 lesion.

PROCEDURAL CHARACTERISTICS

The main branch was predilated in 50 patients and the side branch in 41 patients. All patients with good angiographic results were then treated with the Nile PAX drug-eluting stent in the main branch.

The distal diameter of the dedicated stent used was 3.5 mm for 10 patients, 3.0 mm for 33 patients and 2.50 mm for seven patients. Then, a kissing inflation with the balloon dedicated to the side branch of the Nile PAX system was performed on 40 patients (diameter 2.0 mm for six patients, 2.5 mm for 26 patients and 3.0 mm for eight patients). For the remaining patients, the opening of the SB post-stenting was performed with SB inflation only.

After checking that there was no dissection of the ostium of the side branch and after removal of the guidewire in the main branch to avoid any friction between the drug-eluting balloon and guidewire, the DANUBIO balloon was successfully inflated in the side branch of all 50 patients for a mean duration of a 41±25 seconds. The diameter of the DANUBIO balloon was chosen by the investigator according to visual estimation: a 2 mm balloon was chosen for 11 patients, a 2.5 mm balloon for 26 patients and a 3 mm balloon for 13 patients. The mean balloon inflation pressure was 9.4±2.2 atmospheres. One Amazonia PAX (Minvasys) stent was positioned distally in the main branch for a residual dissection.

In total, taking this additional stent into account, the procedure was successful in 100% of cases.

ANGIOGRAPHIC RESULTS

Angiographic analysis was possible for all patients before and immediately after the procedure, whereas angiographic follow-up at six months was performed on 48 patients (96%). The results of this analysis are given in Table 2.

Before the procedure, the reference diameter of the proximal main branch was 2.93±0.43 mm, which did not change after the procedure (3.03±0.38 mm). The reference diameter of the side branch was 1.84±0.34 mm, without modification after the procedure. Mean lesion length was 12.0±3.5 mm in the main branch and 5.50±0.97 mm in the secondary branch. The percentage stenosis in the main branch before the procedure was 62±14%, decreasing to 26±11% after placement of the stent. The mean percentage of the side branch diameter stenosis was 30.0±14.5% before the procedure, decreasing to 23.9±11.3% after the procedure.

Representations of the minimum lumen diameter (MLD) in the side branch including cumulative frequency curves for all patients pre-procedure, post-procedure and at follow-up are presented in Figure 3A and Figure 3B.

The MLD of the side branch increased from 1.36±0.38 mm to 1.52±0.31 mm after the procedure (p=0.01), and remained
identical at 1.55±0.35 mm at six months. The LLL at the ostium was, on average, slightly negative (–0.04±0.34 mm) and homogeneous in the population, with no difference between the group of patients with significant lesions of the side branch before the procedure (n=32) and the group without side branch lesions (n=18). The MLD of the proximal and distal main branch provided in Table 2 increased from 1.39±0.62 mm and 1.44±0.50 mm before the procedure to 2.62±0.31 mm and 2.18±0.33 mm after the procedure (p<0.0001), respectively, and it decreased to 2.02±0.54 mm and 1.52±0.50 mm at six months (p<0.0001).

Table 2. Angiographic acute and 6-month follow-up results.

<table>
<thead>
<tr>
<th></th>
<th>MB proximal</th>
<th>MB distal</th>
<th>SB</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pre-procedure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RVD, mm</td>
<td>2.93±0.43</td>
<td>2.16±0.39</td>
<td>1.84±0.34</td>
</tr>
<tr>
<td>MLD, mm</td>
<td>1.39±0.62</td>
<td>1.44±0.50</td>
<td>1.36±0.38</td>
</tr>
<tr>
<td>DS, %</td>
<td>52.6±20.0</td>
<td>33.3±19.5</td>
<td>29.8±14.5</td>
</tr>
<tr>
<td><strong>Post-procedure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RVD, mm</td>
<td>3.03±0.38</td>
<td>2.48±0.32</td>
<td>1.83±0.33</td>
</tr>
<tr>
<td>MLD, mm</td>
<td>2.62±0.31</td>
<td>2.18±0.33</td>
<td>1.52±0.31</td>
</tr>
<tr>
<td>DS, %</td>
<td>13.1±6.5</td>
<td>11.9±6.7</td>
<td>23.9±11.3</td>
</tr>
<tr>
<td>Acute gain</td>
<td>1.18±0.68</td>
<td>0.74±0.45</td>
<td>0.18±0.34</td>
</tr>
<tr>
<td><strong>6-month follow-up</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Angiographic binary restenosis, %</td>
<td>4.0</td>
<td>14.0</td>
<td>2.0</td>
</tr>
<tr>
<td>RVD, mm</td>
<td>2.91±0.47</td>
<td>2.22±0.42</td>
<td>1.96±0.34</td>
</tr>
<tr>
<td>MLD, mm</td>
<td>2.02±0.54</td>
<td>1.52±0.50</td>
<td>1.55±0.35</td>
</tr>
<tr>
<td>DS, %</td>
<td>30.1±16.5</td>
<td>31.9±18.8</td>
<td>25.7±12.6</td>
</tr>
<tr>
<td>In-stent LLL, mm</td>
<td>0.61±0.53</td>
<td>0.69±0.46</td>
<td>-0.04±0.34</td>
</tr>
<tr>
<td>MB in-stent LLL, mm</td>
<td>0.54±0.60</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

DS: diameter stenosis; MB: main branch; MLD: minimum lumen diameter; RVD: reference vessel diameter; SB: side branch

Note that the reference diameter of the side branch increased between post-procedure and six months from 1.83±0.33 mm to 1.96±0.34 mm (p=0.06). A graphical representation of the location of the different restenoses is provided in Figure 4. Six were located in the distal portion of the main branch stent, one at the bifurcation segment of the stent matched with the only ostial side branch restenosis, one in the proximal segment of the stent and two at the edges.

Figure 3. Minimum lumen diameters of the side branch: pre-procedure, post-procedure and at follow-up. A) Cumulative frequency distribution curves. B) Individual data.

Figure 4. Restenosis localisation. DM: distal main branch; PM: proximal main branch; SB: side branch

**CLINICAL OUTCOME**

There were no in-hospital complications and no adverse clinical events up to 30 days, except for a single myocardial infarction (MI: 2.0%) where the patient received emergency revascularisation for an acute occlusion in the target lesion at the level of the stent. There were no additional MI and, within the six-month follow-up, only one patient had a non-clinically driven TLR at the level of the side branch combined with a main branch revascularisation (2.0%). Non-clinically driven re-dilation was performed on three patients (6.0%) at the level of the lesion in the main vessel, one patient on the vessel beyond the lesion, and in two patients angioplasty was performed on another vessel at six months. A total of five major adverse cardiac events (hierarchical and cumulative: 10.0%) were noted during follow-up. To date, there has been no reported cardiac death (0.0%).

**Discussion**

The main results of the DEBSIDE study were the good angiographic and clinical outcomes in the side branch after treatment with a drug-eluting balloon at six months post-procedure. Although the angiographic results obtained in the main branch were not as favourable as in the side branch, the clinical results were acceptable. Indeed, only 6.0% of patients needed repeat revascularisation at the target lesion at the level of the main branch.

The treatment of bifurcated coronary artery stenosis remains challenging and the complication rate is high. PCI in bifurcation lesions is associated with lower immediate angiographic and clinical success and higher rates of restenosis, which, in addition, are more challenging to treat compared to non-bifurcated lesions."
Data from the Nordic Bifurcation Study\textsuperscript{16} and the British Bifurcation Coronary study\textsuperscript{17} (BBC ONE) showed that the method of choice to treat bifurcation lesions is stenting the main branch and only dilating the side branch, because double stenting appears not to lower the risk of restenosis, while the placement of two stents in the bifurcation makes the procedure more difficult to perform. Thus, the current practice to treat bifurcation lesions is the provisional T-stenting technique, i.e., stenting only the MB and final kissing balloon, avoiding SB stenting when possible. However, with this technique, whilst the success rate in the main branch is satisfactory, the rate of restenosis in the side branch remains problematic. At the six-month angiographic follow-up in Nordic Bifurcation I, the restenosis rate in the side branch was 19.2\% when the side branch was treated with a simple strategy without placement of a second stent, and 11.5\% when a second drug-eluting stent was positioned in the side branch\textsuperscript{16}.

In the BBC ONE clinical study\textsuperscript{17}, where there was no systematic angiographic follow-up, the restenosis rate in the side branch was 2.7\% in the simple strategy group.

The most recent study with angiographic follow-up is the DKCRUSH-II study\textsuperscript{16}. In this study, which used a complex double kissing technique with two stents, the late lumen loss in the side branch was assessed as $0.22\pm0.41$ mm and it was identical in the control group with one drug-eluting stent in the main branch and a conventional balloon in the secondary branch $(0.18\pm0.45)$ mm. All of these publications confirm that the result of treatment of the side branch remains problematic since the use of a drug-eluting stent as compared to a conventional balloon does not seem to yield significant improvements.

For this reason, the use of a drug-eluting balloon may be useful to provide an antiproliferative substance to a lesion without the use of a polymer-covered stent.

Over the past few years, drug-eluting balloons have demonstrated their clinical safety and efficacy for the treatment of small coronary vessels\textsuperscript{19} and in-stent restenosis\textsuperscript{20,21}. Drug-eluting balloons have now received a guideline\textsuperscript{22} class I level of evidence A recommendation in the treatment of in-stent restenosis after bare metal stent (BMS) or drug-eluting stent implantation.

To date, there have been very few clinical trials on the treatment of bifurcation lesions with drug-eluting balloons and all of these included only a small cohort of patients.

The DEBIUT registry\textsuperscript{2} enrolled 20 patients with coronary artery bifurcation lesions. A DEB (the first generation of the DIOR\textsuperscript{8} balloon catheter; Eurocor GmbH, Bonn, Germany) was used during the predilatation step in both main and side branches before provisional main branch stenting with a BMS. Final kissing inflation with regular balloons was performed according to routine practice. The procedure was successful in all patients. The use of sequential predilatation with a DEB was considered safe and well tolerated. No stents were placed in the SB. No MACE or reintervention occurred during the four-month clinical follow-up period, and there was no subacute vessel closure. This registry provided encouraging results for the use of a DEB in bifurcation treatment to improve long-term side branch patency. However, the larger randomised DEBIUT study\textsuperscript{6,7}, initiated after the registry, did not confirm these initial promising results. Indeed, the outcome of the primary endpoint was not reached. The proximal MB late lumen loss was better in the group treated with paclitaxel DES+plain old balloon angioplasty (POBA) $(0.18 \text{ mm})$ than in the group treated with sequential MB/SB dilatation with the DEB then BMS implantation in the MB $(0.42 \text{ mm})$. Similarly, the SB LLL was better in the DES+POBA group (DEB+BMS $0.19\pm0.66$ mm vs. DES $0.11\pm0.43$). These "poor" results obtained with the DIOR I led to the development of the DIOR II (second generation) with a coating involving a new excipient (shellac)\textsuperscript{7}. Preclinical results obtained with the DIOR II showed much higher drug concentrations into the vessel wall than the DIOR I\textsuperscript{1}. This emphasises the importance of the excipient in DEB as discussed by Afari et al\textsuperscript{12}.

The PEPCAD V trial\textsuperscript{3} was a feasibility and safety first-in-man series evaluating a DEB (SeQuent\textsuperscript{9} Please balloon catheter; B. Braun Melsungen AG, Melsungen, Germany) in the treatment of significant de novo bifurcation stenosis. The MB and SB were predilated with the DEB followed by a bare metal stent (Coroflex\textsuperscript{8} stent; B. Braun Melsungen AG) deployment in the main branch. Twenty-eight patients were included in the study. There were no MACE up to 30 days and no TLR was performed. However, two late stent thromboses were reported (one definite and one probable at six and eight months after the index procedure, respectively). Nine-month angiographic results showed an in-stent late lumen loss of $0.38\pm0.46$ mm in the main branch and an in-lesion unsatisfactory late lumen loss of $0.21\pm0.48$ mm in the side branch.

In a more recent prospective randomised clinical study, the BABILON trial\textsuperscript{9}, sequential MB/SB dilatation with a DEB (SeQuent Please), with provisional T-stenting with a BMS in the MB (52 patients), was compared to a sequential MB/SB dilatation with conventional plain balloons followed by everolimus DES implantation in the MB (56 patients). As opposed to the previous study, both strategies showed similar good results in the SB, LLL of $-0.04\pm0.71$ mm in the DEB followed by BMS group, and $-0.03\pm0.51$ mm in the DEB followed by everolimus stent group.

In the literature, only one clinical trial has studied the combination of DES in the MB followed by DEB in the SB. Indeed, in a recent study conducted by Herrador et al\textsuperscript{10}, a DES (TAXUS\textsuperscript{TM} Libertet\textsuperscript{TM}; Boston Scientific, Marlborough, MA, USA) was implanted in the main branch, which is the usual approach for the treatment of a bifurcation lesion, and the SB was treated with a DEB (SeQuent Please). Fifty patients treated with the DEB were compared to 50 patients treated with a conventional balloon in the SB. angiographic follow-up and IVUS were scheduled at 12 months post-procedure. At angiographic follow-up, there was a lower percentage of SB restenosis in the DEB group (20\% vs. 7\%, $p=0.08$), with less SB late loss $(0.40\pm0.50 \text{ mm vs. } 0.09\pm0.40 \text{ mm}, p=0.01)$ and similar MB late loss $(0.62\pm0.70 \text{ mm vs. } 0.49\pm0.60 \text{ mm}, p=0.39)$. The MACE rate at 12-month follow-up was 24\% in the group treated by a conventional balloon versus...
11% in the DEB group (p=0.11). The TLR rate was also higher in the conventional balloon group (22% vs. 12%, p=0.16).

The angiographic and clinical results obtained in our study are highly encouraging for the use of DEB as the final treatment in the side branch and are significantly better than the results obtained with the same stent in the BIPAX study involving the same stenting strategy without using a DEB in the SB (–0.04 vs. 0.33 mm LLL, respectively). Indeed, in the present study, only one single patient had restenosis and target lesion revascularisation at the level of the side branch. The association of the discrete increase in the reference diameter of the side branch and the unchanged MLD in the ostium of the side branch explains the very low restenosis rate (defined as a stenosis greater than 50% diameter) of 2% in the side branch, whereas it reached 4.0% in the proximal MB and 14.0% in the distal MB. The side branch late lumen loss measured six months after the index procedure (–0.04±0.34 mm) was similar to BABILON trial results5, slightly better than Herrador’s study16 or much better than that in previously published studies on bifurcation treatment with DEB (Table 3) or that obtained with DES17-24.

Table 3. Side branch late lumen loss at six months in DEBSIDE and previous trials.

<table>
<thead>
<tr>
<th>Trial</th>
<th>Study device</th>
<th>Technique</th>
<th>Months</th>
<th>SB LLL (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEBSIDE (2014)</td>
<td>DANUBIO</td>
<td>Predilatation: conventional balloon in MB+SB MB: DES SB: DEB</td>
<td>6</td>
<td>–0.04±0.34</td>
</tr>
<tr>
<td>PEPCAD V (2011)4</td>
<td>SeQuent Please</td>
<td>MB+SB: DEB MB: BMS</td>
<td>9</td>
<td>0.21±0.47</td>
</tr>
<tr>
<td>DEBIUT (2011)2</td>
<td>Dior I</td>
<td>Predilatation: DEB in MB+SB MB: BMS</td>
<td>6</td>
<td>0.19±0.66</td>
</tr>
<tr>
<td>Herrador et al (2013)15</td>
<td>SeQuent Please</td>
<td>Predilatation: conventional balloon in MB or SB SB: DEB MB: DES</td>
<td>12</td>
<td>0.09±0.40</td>
</tr>
<tr>
<td>BABILON (2014)3</td>
<td>SeQuent Please</td>
<td>Predilatation: conventional balloon in MB+SB MB: BMS</td>
<td>9</td>
<td>–0.04±0.64</td>
</tr>
</tbody>
</table>

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Technique Months SB LLL (mm)

Although suboptimal angiographic results obtained in the main device Techniques Months SB LLL (mm)
branch were observed, they did not impact on the good clinical and angiographic results in the side branch. Additional and complementary studies with larger populations, combining newer-generation limus-eluting stents, which could enable better results on the main branch, along with the DANUBIO drug-eluting balloon would certainly be of especial interest. In this context, a sirolimus-eluting version of the Nile dedicated stent platform is currently being developed.

**Impact on daily practice**

Restenosis following drug-eluting stent implantation in the treatment of bifurcation lesions remains a concern. The six-month angiographic results of the DEBSIDE trial show that drug-eluting balloon inflation using the DANUBIO balloon resulted in very low rates of restenosis of the side branch and, while further studies are warranted, its use may be a significant addition to the interventionalist’s armamentarium by avoiding unnecessary side branch stenting in the treatment of all types of bifurcation lesion.

**Funding**

This study was supported by a grant from Minvasys.

**Conflict of interest statement**

J. Berland reports having received lecture honoraria and consulting fees from Minvasys. The other authors have no conflicts of interest to declare.

**References**


