Original Studies

Head-to-Head Comparison of Sirolimus-Eluting Stent Versus Bare Metal Stent Evaluation of the Coronary Endothelial Dysfunction in the Same Patient Presenting with Multiple Coronary Artery Lesions: The CREDENTIAL study

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Objectives: To assess the endothelial dysfunction (ED) after bare metal stents (BMS) and sirolimus eluting stents (SES) implantation in the same patient, overcoming the confounding role of individual variables. Background: SES reduce restenosis rate compared to BMS but causes more ED. ED is a potentially unsafe phenomenon, since it is the first step in the cascade of atherosclerosis. Studies showing more pronounced ED with drug eluting stents than BMS involved different series of patients, making the comparison difficult because endothelial function (EF) is responsive to many risk factors. Methods: we designed a prospective comparison of 6 months post-deployment EF of SES versus BMS implanted in the same patient, but in different coronary segments. Forty-eight lesions were randomly assigned on a 1:1 allocation using block sizing of 4 according to a computer-generated sequence (SAS System, Version 9.1) basis to treatment with SES or BMS. The EF was evaluated by measuring vessel diameter variation in the stented segment, before and after selective intracoronary infusion of acetylcholine (iiACh). Results: In eligible patients, the relative magnitudes of major vasoconstriction were 2.6, 2.9, 4.6, and 3.1 at 5 mm proximal and 5, 10 and 20 mm distal to the stent edge. Overall, a 3.5-fold major distal vasoconstriction after iiACh of SES vs. BMS was calculated. Conclusions: in the same patients, but treating different coronary segments, SES implantation induces a higher rate of vasoconstriction compared to BMS. The increased vasoconstriction after iiACh is an indicator of ED. © 2013 Wiley Periodicals, Inc.

Key words: endothelial function; acetylcholine; endothelium

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INTRODUCTION

Sirolimus eluting stents (SES) reduce restenosis rate compared to bare metal stents (BMS) [1], but cause more pronounced endothelial dysfunction (ED) due to lack or delay of endothelization. The endothelium plays a critical role in vascular homeostasis by secreting substances and influencing vascular inflammation and cell migration: lack or changes of this natural barrier for blood lipids and lipid deposition could trigger the atherogenesis process and could be associated with thrombogenicity. This potentially unsafe phenomenon [2], is the first step in the cascade of atherosclerosis [3,4], as well as a reliable predictor of future coronary events [5–7]. The studies showing more pronounced ED with SES than BMS involved different series of patients implanted with different types of stent, making the comparisons difficult since endothelial function (EF) is responsive to many risk factors. To overcome the confounding role of individual variables, we designed a prospective randomized comparison of 6-months post-deployment EF of SES versus BMS implanted in different coronary segments of the same patient.

MATERIALS AND METHODS

Study Design

The study was conducted at the Interventional Cardiology Unit of the San Camillo-Forlanini Hospital, in Rome-Italy. Patients with stable angina pectoris, positive stress test and indication to percutaneous coronary angioplasty (PCI) for at least 2 de novo >70% coronary stenoses were considered eligible. The two target lesions should have lengths >10 and <30 mm and comparable angiographic characteristics: differences measuring no more than 0.5 mm in reference vessel diameter and <50% in lesion length. Stents were implanted in different vessels or in the same vessels but in different ramifications. The following variables were recorded for each patient: age, sex, body mass index, family history of heart disease, prior myocardial infarction, angina class, heart failure class, stroke or transient ischemic attack, chronic renal failure (defined as serum creatinine >2 mg/dl), diabetes mellitus, hypertension, smoking, dyslipidemia, c-reactive protein and homocysteine levels. PCI was performed according to standard guidelines. The most severe of two lesions was randomly assigned to receive a SES (Cypher, Cordis Corporation, Miami Lakes, Florida) or a BMS (Coroflex Blue, B. Braun Melsungen, Germany) with allocation 1:1 using block sizing of 4 according to a computer-generated sequence (SAS System, Version 9.1). Consequently, a different stent was implanted on the second lesion. The treatment of further stenosis, if present, was left to the operator’s discretion and these stenoses were excluded from the analysis (Fig. 1—Study Protocol). Particular attention was paid to obtain the best angiographic result and to implant only one stent for each lesion. All patients received optimal medical therapy, including aspirin 75 mg/day and clopidogrel 75 mg/day for at least 6 months. All patients were asked to return for invasive evaluation at 6 month of follow-up (FU). Medications with potential effects on vasomotor responses were discontinued 72 hr before the procedure, but short-acting nitrates were permitted 6 hr prior to the procedure. Baseline coronary angiograms were taken and patients without intra-stent or peri-stent restenosis were included in the pharmacological protocol study. The protocol included intracoronary infusion of 0.9% normal saline (2 ml for 1 min), followed by baseline coronary angiography. The endothelium-dependent vasomotor response was estimated after a 2 min super-selective intracoronary infusion of acetylcholine $10^{-5}$ mol/l (iiAch) using a pump (Perfusor Compact B. Braun Melsungen, Germany) and a coronary micro-catheter (FINECROSS, Terumo Corporation, Tokyo, Japan) positioned 3 cm above the proximal edge of the stent; after iiAch infusion, the micro-catheter was withdrawn and coronary angiograms were collected every 60 sec for 3 min (Fig. 2). The endothelial independent vasomotion was assessed 1 min after an intracoronary 200 µg bolus of nitroglycerin. Clinical status, heart rate, arterial pressure, and electrocardiographic leads were continuously monitored.

Exclusion Criteria

Exclusion criteria were (a) clinical data: acute coronary syndrome in the last 3 months, severe risk factors for ED [uncontrolled diabetes mellitus (defined as HbA1C>9%), uncontrolled hypertension (systolic blood pressure >180 mm Hg), hypercholesterolemia (total cholesterol >240 mg/dl), persistent smoking]; any contraindication/intolerance to the use of aspirin, heparin, and/or clopidogrel; chronic renal failure requiring dialysis; severe left ventricular dysfunction (defined as an ejection fraction <35% by echocardiography); survival expectancy <1 year; (b) basal coronary angiographic findings: reference vessel diameter <2.5 mm, vasospasm, fresh thrombus, dissection, bifurcation/ostial lesions; (c) FU angiographic findings: restenosis (vessel diameter reduction >50%) or development of de novo significant stenosis (>70%). The study end-point was maximal coronary vasomotor response to iiAch. It was determined as a drug-induced percentage change in vessel diameter using baseline angiogram as reference. Eight points in the stent and peri-stent site and the proximal (10 mm proximal to the proximal stent edge) and distal (distal stent edge to 20 mm distally) average segment...
540 patients screened
Not meeting inclusion criteria (n=483)
Declined to participate (n=12)
Other reasons (n=21)

Randomized 24 patients
(48 coronary segments)

Allocated to BMS (n=24 lesions)
Allocated to SES (n=24 lesions)

Lost to FU: 3 patients refused, 1 patient suffered AMI in non stented vessel
20 patients performed 6 months angiographic FU

Excluded to final analysis: 4 patients in the BMS group for intra-stent restenosis, 2 for progression of lesion in non target lesion

Analysed 28 coronary segments (14 patients)

Fig. 1. Study Protocol. AMI: acute myocardial infarction; BMS: bare metal stents; SES: sirolimus eluting stents; FU: follow-up.

Diameters were analyzed (Fig. 2). The percent changes in vessel diameter after iiAch and nitrates were calculated and compared between SES and BMS treated segments and baseline. Two orthogonal views with less foreshortening or without overlapping of side branches were selected and averaged for biplane assessment by two experts. End-diastolic images for each segment were chosen and quantitative coronary angiography (QCA) was performed using the CAAS II system (Pie Medical Imaging, Maastricht, The Netherlands). The contrast-filled tip catheter was used for calibration. Independent, masked reviewers performed the QCA measurements at baseline, after iiAch and nitrates. The independent predictors of ED were also investigated.

Conduct of the Clinical Study
A masked, independent committee collected the endpoints; an independent study monitor verified all data from the reported cases. The Local Institutional Review Board approved the Protocol; all participants were provided with written, informed consent forms. The operators were aware of the assigned stent during PCI, but at angiographic FU and EF evaluation, staff was blinded to the allocation of stent type. A sample size has been calculated on the basis of previous reports [8–11] and we anticipated the occurrence of maximal vasoconstriction in response to iiAch measured at 5 mm segments proximal and distal to the stent would be 40–60% and 0–20% respectively for SES versus BMS. Assuming a 0.05% alpha type error and 0.95 power, a total of 20 patients needed to be enrolled for a paired data study. To take into account potential losses to FU, we randomized 24 patients.

Statistical analysis: baseline characteristics are expressed as mean ± SD and categorical variables as a number (n) and percentage (%). The univariate comparisons between the continuous variables were performed...
using the Student’s t-test for paired and unpaired data. To identify the potential variables to enter into a multivariate predictive model, we tested the correlation between clinical and angiographic variables. Multiple linear regression analysis was used to determine independent predictors of ED. Variables considered were: age, gender, diabetes, hypertension, current smoker, statin, and angiotensin converting enzyme inhibitor/angiotensin-receptor blocker used. A P value of <0.05 was considered statistically significant. SPSS statistical program (Chicago, IL) was used.

RESULTS

During the study period, 540 patients were screened, 24 were enrolled and four patients were excluded: three (12.5%) refused the angiographic FU, one (4.1%) suffered acute coronary syndrome in a nonstented vessel. Angiographic FU was performed in 20 patients (83%), four (16.6%) had intrastent restenosis of the BMS, two (8.3%) showed significant progression of lesions in non-target vessels, and 14 patients entered in the final analysis. The flow diagram of the trial is provided in Fig. 1. Baseline clinical characteristics are presented in Table I. SES versus BMS had comparable angiographic and procedural characteristics (Table II). There were no differences between SES and BMS regarding mean stent length and mean stent diameter (19.29 ± 7.29 mm versus 16.1 ± 2.9 mm, P = 0.25; 2.86 ± 0.39 mm versus 3.2 ± 0.4 mm, P = 0.38, respectively).

Among the BMS, there were ten direct stents without post-dilatation and four stent implantations with pre- and post-dilations; among the SES there were nine direct stents without post-dilatation and five stent implantations with pre and post-dilations. The mean interval from stent implantation to FU angiography was 180.4 ± 10.3 days. Blood sample results at FU are listed in Table III. The percentage variation in vessel diameters after iiAch in the eight predefined points and two segment mean diameters for SES and BMS are shown in Figs. 2 and 3.

<table>
<thead>
<tr>
<th>Percentage of vasoconstriction</th>
<th>SES (%)</th>
<th>12.1</th>
<th>25.1*</th>
<th>6.4</th>
<th>8</th>
<th>11.7</th>
<th>40.5*</th>
<th>47.9*</th>
<th>44.6*</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMS (%)</td>
<td>3.8</td>
<td>9.3</td>
<td>6.9</td>
<td>6.1</td>
<td>7.5</td>
<td>13.8</td>
<td>10.3</td>
<td>14.2</td>
<td></td>
</tr>
</tbody>
</table>

*P < 0.05.

**Table I. Baseline Characteristics**

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>70.8 ± 7.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body surface area (m²)</td>
<td>1.87 ± 0.11</td>
</tr>
<tr>
<td>Male sex</td>
<td>9 (64.2%)</td>
</tr>
<tr>
<td>Current smokers</td>
<td>4 (28.5%)</td>
</tr>
<tr>
<td>Positive family history of HD*</td>
<td>8 (57.1%)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>13 (92.8%)</td>
</tr>
<tr>
<td>Prior myocardial infarction</td>
<td>6 (42.8%)</td>
</tr>
<tr>
<td>Diabetes mellitus type II</td>
<td>5 (35.7%)</td>
</tr>
<tr>
<td>Stable angina (CCS‡)</td>
<td>7 (50%)</td>
</tr>
<tr>
<td>II</td>
<td>4 (28.5%)</td>
</tr>
<tr>
<td>III</td>
<td>3 (21.4%)</td>
</tr>
</tbody>
</table>

*HD = heart disease; ‡CCS = Canadian Cardiovascular Society. Data are presented as numeric value (± SD) and percentage (%).
The relative magnitudes of major vasoconstriction for SES versus BMS were 2.6 \((P = 0.04)\), 2.9 \((P = 0.03)\), 4.6 \((P = 0.001)\) and 3.1 \((P = 0.002)\) at 5 mm proximal and 5, 10, and 20 mm distal to the stent edge. Overall, a 3.5-fold major distal vasoconstriction after iiAch of SES versus BMS was calculated as an average of the three predefined distal diameters (coronary angiography of patient \(n\) 11 in Fig. 4 after iiAch and nitrates).

In univariate analysis, the independent significant predictors that correlated with increased ED were SES, diabetes, hypertension, low High Density Lipoprotein levels, presence of atherosclerosis distal to SES implantation site, age, increased C-Reactive Protein levels and prior myocardial infarction.

After intracoronary nitrates were administered, there were no statistically significant differences in vessel diameters between SES versus BMS for any of the evaluated segments.

No differences were recorded in the mean arterial blood pressure during the iiAch compared to the baseline. Two patients had an episode of temporary asystolia (less than 5 sec) that recovered spontaneously. Intraobserver and interobserver variability for quantitative measurements of coronary angiography in the same recordings of 15 randomly selected vessels were 0.062 \pm 0.04 mm and 0.021 \pm 0.03 mm, respectively.

DISCUSSION

This study shows that in the same group of patients, the 6-month FU endothelial dependent coronary
vasomotion is significantly impaired after SES compared to BMS implantation, especially distal to the stent. Instead, no differences exist between the two types of stent in regards to endothelium independent coronary vasomotion. This data supports that SES, rather than BMS implantation is associated with coronary ED [9]. Previous studies have demonstrated more pronounced ED after SES versus BMS, but the comparisons have been assessed in different patients with different risks factors [10–12]. Because EF is a complex process influenced by a number of pathophysiological mechanisms (ethnicity, hypertension, diabetes, smoking, hypercholesterolemia etc.), a considerable variability exists in the healing process after stent implantation. In our study, both types of stent were randomly implanted in two comparable coronary lesions within the same patient. This unique study design actually adjusts the comparison for all variables and definitively assesses the impact of the type of stent on EF.

A methodological issue is the protocol of study of coronary EF. The QCA evaluation of coronary response after iiAch is currently the most utilized invasive tool of investigation of coronary vasomotion [10,11]. The drug acts as a potent vasodilator in normal coronary vessels by promoting the release of endothelial nitric oxide, but in damaged vessels it can cause abnormal vasoconstriction via receptors localized in the smooth muscle cells [13]. iiAch provocation test is a sensitive and safe test, biased by the lack of standardization because a number of different protocols—fixed versus progressive doses, low versus moderate versus high doses, manual versus pump injections, non-selective (a guiding catheter in the coronary ostium) versus superselective (a microcatheter in the target vessel)—all produce variable drug concentrations, potentially contributing to different results of EF after SES or BMS implantation. As an original finding of the study, we aimed to obtain the best control of drug

![Diagram](http://example.com/diagram.png)
concentration by infusing iiAch “super-selectively” into the target vessel. We chose a fixed, single dose (10^{-5} \text{ mol/l}) of the drug, which provides a moderate, consistent vasocostrictive stimulus and ensures a very low rate of potential, serious complications. Our results are in line with previous studies where similar dose of iiAch were used [8,10–12]. Our data addresses the open issues of potential adverse biological effects of SES: coronary endothelium-dependent vasomotion is severely impaired after SES implantation, while it is virtually unaffected after BMS implantation [9,11]. Over 20% vasoconstriction after iiAch has been considered a reliable sign of ED [14]. Many studies have documented the association between ED and serious cardiovascular events [15,16]. It is common experience to detect a higher incidence of restenosis at the edge of the SES, which could be explained by the presence of ED. This phenomenon is consistent with the results of our study, that shows major ED in the same area most affected by restenosis of the SES. Several cases of diffuse coronary spasm after SES implantation have been reported [17,18] and ED is implicated in the increased incidence of very late stent thrombosis with first generation of drug eluting stents (DES), especially after discontinuation of dual antiplatelet therapy. These adverse effects could offset the potential benefits of SES. We have to define the duration of SES-induced ED and evaluate if we can counterbalance its negative effects. In agreement with a Consensus for Preclinical stent evaluation that recommended evaluation of EF as a valuable ancillary tool for differentiating the long-term performance of DES [19], questions raised by this work highlight the need for additional investigations.

Compared to extensive use of DES in the current treatment of ischemic heart disease, EF after SES-PCI focuses on a lingering issue. For the millions of patients in whom SES have already been deployed, aggressive efforts to improve general EF is a “gray” area which is not given sufficient consideration. A potential clinical use of our results highlights the biological effects of SES in terms of EF identifying the patients with increased endothelial sensitivity that require special targeted medical or interventional treatment.

Study Limitations

The number of enrolled patients was limited due to the high rate of drop-out at the angiographic FU, so the final power analysis of the study changed from 0.95 to 0.85 (maintaining an 0.05% alpha type error). At the FU angiogram, the stent type was incompletely masked because a BMS is clearly thinner than the SES. The multiple linear regression analysis was not attempted due to the small sample size. A longer FU with a second invasive evaluation of EF at 1 year would have allowed an evaluation of the recovery timeframe, if present. SES were chosen because, at the time, they were the most studied and most implanted stents in the real world, and, despite the fact that they are no longer used today, they have been implanted in millions of patients. Finally, a similar evaluation of EF of second-gen-DES would be equally interesting.

CONCLUSIONS

In the same patients, but treating different coronary segments, SES implantation induces a higher rate of ED compared to BMS. The increased vasoconstriction after iiAch is an indicator of ED. We calculated: a 3.5-fold vasoconstriction of SES vs. BMS for the distal segment average diameter. ClinicalTriail.gov number NCT01242306.

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