Uncomplicated moderate coronary artery dissections after balloon angioplasty: good outcome without stenting

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Abstract

Objective—To study the relation between moderate coronary dissections, coronary flow velocity reserve (CFVR), and long term outcome.

Methods—523 patients undergoing balloon angioplasty and sequential intracoronary Doppler measurements were examined as part of the DEBATE II trial (Doppler endpoints balloon angioplasty trial Europe). After successful balloon angioplasty, patients were randomised to stenting or no further treatment. Dissections were graded at the core laboratory by two observers and divided into four categories: none, mild (type A-B), moderate (type C), severe (types D to F). Patients with severe dissections (n = 128) or without available reference vessel CFVR (n = 139) were excluded. The remaining 256 patients were divided into two groups according to the presence (group A, n = 45) or absence (group B, n = 211) of moderate dissection.

Results—Following balloon angioplasty, there was no difference in CFVR between the two groups. At 12 months follow up, a higher rate of major adverse cardiac events was observed overall in group A than in group B (10 (22%) vs 23 (11%), p = 0.041). However, the risk of major adverse events was similar in the subgroups receiving balloon angioplasty (group A, 6 (19%) v group B, 16 (16%), NS). Among group A patients, the adverse events risk was greater in those randomised to stenting (odds ratios 6.603 v 1.197, p = 0.046), whereas there was no difference in risk if the group was analysed according to whether the CFVR was < 2.5 or ≥ 2.5 after balloon angioplasty.

Conclusions—Moderate dissections left untreated result in no increased risk of major adverse cardiac events. Additional stenting does not improve the long term outcome.

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Keywords: coronary dissection; intracoronary Doppler; angioplasty

Coronary artery dissection is observed angiographically in up to 50% of cases after balloon angioplasty. Although coronary stenting has greatly curbed the need for urgent surgical revascularisation for dissections that impair distal perfusion, it remains to be established whether moderate dissections with unimpaired flow and a good epicardial lumen would benefit from additional stenting. Previous studies have shown that mild to moderate angiographic dissections do not increase the risk of major adverse cardiac events or restenosis rate at a six month follow up after balloon angioplasty. However, limited data are available on the impact of stenting on the short and long term clinical outcome after the development of moderate dissections.

We investigated the relation between dissection after balloon angioplasty and coronary flow velocity reserve, and the impact of stenting on the subsequent clinical outcome in patients treated by balloon angioplasty or stenting in the DEBATE II trial (Doppler endpoints balloon angioplasty trial Europe).

Methods

PATIENTS

Patients scheduled to undergo angioplasty because of stable or unstable angina pectoris or documented myocardial ischaemia, caused by a single de novo coronary stenosis less than 25 mm long and potentially amenable to stent implantation, were eligible for the DEBATE II trial. Those with total coronary occlusions, ostial lesions, bifurcated lesions, lesions in a previously bypassed vessel, lesions in an extremely tortuous vessel, or lesions containing thrombus were excluded from the study, as were patients with previous Q wave infarction, or patients with previous Q wave infarction in the myocardial territory supplied by the target vessel, or evolving myocardial infarction in the previous week. The study was carried out according to the principles of the Declaration of Helsinki, and all patients provided written informed consent.

STUDY OBJECTIVES AND DESIGN

The primary objective of the DEBATE II trial was to compare the cost-effectiveness of elective stent implantation (primary stenting) with balloon angioplasty guided by quantitative coronary angiography and Doppler flow velocity measurements. Stent implantation was permitted for bail out situations or whenever an “optimal result” could not be achieved. The secondary objective was to evaluate differences in benefits of additional stenting in patients with and without an optimal result. Thus a double randomisation was required. The first randomisation (1:5) allocated 620 patients to...
either primary stenting \((n = 97)\) or guided balloon angioplasty \((n = 523)\). All patients in the guided balloon angioplasty group who did not require bail out stenting \((n = 395)\) underwent a second randomisation to additional stenting or termination of the procedure. For the purpose of our analysis, we selected from the latter 395 patients all those \((n = 256)\) in whom a reference vessel coronary flow velocity reserve \((\text{CFVR})\) measurement was available, and divided them according to the presence or absence of moderate type C dissections (fig 1).

### GUIDED BALLOON ANGIOPLASTY

**Doppler flow measurements**

Target vessel Doppler measurements were performed before and after balloon angioplasty and again following additional stent implantation. It was also a requirement to perform a Doppler assessment of the CFVR of an adjacent angiographically non-diseased reference vessel \((< 30\% \text{ diameter stenosis})\). A 0.014 inch \((0.36 \text{ mm})\) Doppler guide wire \((\text{CardioMetrics FloWire}; \text{EndoSonics, Rancho Cordova, California, USA})\) was advanced distal to the lesion, and velocity recordings were obtained under basal and hyperaemic conditions. Maximum hyperaemia was induced by adenosine, either by an intracoronary bolus injection \((12 \mu g \text{ for the right coronary artery and } 18 \mu g \text{ for the left coronary artery})\) or by intravenous infusion \((140 \mu g/\text{kg/min})\). Absolute CFVR was calculated as the ratio of hyperaemic to baseline time averaged peak velocity. Relative CFVR was calculated as the ratio of the absolute CFVR to the non-diseased reference vessel CFVR.

### Quantitative coronary angiography

Intracoronary glyceryl trinitrate \(0.1–0.3 \text{ mg}\) or isosorbide dinitrate \(1–3 \text{ mg}\) was given to achieve maximum coronary vasodilatation. At least two cineangiograms were performed before the angioplasty or stenting procedure and were repeated in the same projections afterwards. Quantitative angiography was performed using a standardised protocol described previously.5

#### Definition of an “optimal” result

An “optimal” result (on quantitative angiography and coronary flow reserve determination) was defined as a diameter stenosis of < 35% and a coronary flow reserve of > 2.5,7 and was achieved by upsizing the balloon or increasing the inflation pressure, or both, if necessary.

#### Bail out stenting

Bail out stenting was allowed in the following situations: a residual stenosis of more than 50%; dissection of type D, E, or F; persistent myocardial ischaemia along with a dissection type C; a fall in thrombolysis in myocardial infarction (TIMI) flow grade of at least one grade; or TIMI grade 0 or 1.

### SECOND RANDOMISATION

After an optimal result was achieved or when further attempts to improve the result were deemed unsafe by the operator, the final diameter stenosis and coronary flow velocity reserve were assessed. Thereafter, and irrespective of these measurements, the second randomisation was performed.

### DISSECTION EVALUATION

Intimal dissection incidence and grading was determined by an independent core laboratory \((\text{Cardialysis BV})\) classification,1 blinded to Doppler flow results and clinical outcome, according to the National Heart, Lung, and Blood Institute \((\text{NHLBI})\) classification:

- type A: small radiolucent area within the vessel;
- type B: no persisting extravasations of contrast;
- type C: persisting contrast medium extravasations;
- type D: spiral filling defect with delayed but complete distal flow;
- type E: persistent filling defect with delayed antegrade flow;
- type F: filling defect with total occlusion.

Dissections were clinically divided into “mild” dissections (type A or B), “moderate” dissections (type C without signs or symptoms of ischaemia), or “severe” dissections (type C with symptoms or signs of ischaemia plus types D to F). The patient population was analysed on the basis of the presence \((\text{group A})\) or absence \((\text{group B})\) of uncomplicated “moderate” dissections.
EFFICACY END POINTS

For the DEBATE II study, the efficacy end point was a composite of major adverse cardiac events within 12 months after the procedure and included the following: death from any cause, non-fatal myocardial infarction, and percutaneous or surgical target lesion revascularisation. After hospital discharge, patients were seen at the outpatient clinic at one, six, and 12 months. No follow up angiogram was performed unless clinically indicated.

STATISTICAL ANALYSIS

Continuous variables are expressed as mean (SD), and differences between groups of patients were studied using the unpaired Student’s t test or one way analysis of variance, as appropriate. Categorical variables are presented as percentages, and differences between groups were evaluated using the χ² test or Fisher’s exact test. Multivariate logistic regression analysis was used to study the value of the clinical, angiographic, and Doppler derived data to predict major adverse cardiac events at the 12 months follow up. Odds ratios and 95% confidence intervals (CI) are presented. The Breslow-Day test was used to assess the homogeneity of odds ratios between subgroups. The log rank test was applied to study differences in event-free survival between subgroups at the 12 months follow up. All statistical tests were two tailed, and significance was assumed at p < 0.05.

RESULTS

From 523 patients randomised to the guided angioplasty group in the DEBATE II trial, 128 underwent bail out stenting because of severe dissections. All the remaining patients (n = 395), irrespective of the presence or absence of a moderate dissection, underwent a second randomisation to additional stenting or to halting the procedure. Of these 395 patients, 139 were excluded from our analysis because no reference CFVR measurements were available. The remaining 256 patients were divided into two groups according the presence (group A, n = 45) or absence (group B, n = 211) of uncomplicated moderate dissections (fig 1).

BASELINE CHARACTERISTICS

Patients’ baseline characteristics are summarised in table 1. In group A patients were older and had a smaller proportion of smokers than those in group B. Lesion characteristics are given in table 2. Group A had a greater proportion of calcified lesions than group B. Both groups had similar vessel size (mean (SD): 2.93 (0.43) mm v 3.03 (0.59) mm, group A v group B, respectively (NS)). In group B, 110 of the 211 patients (53%) had no dissection, whereas 101 (47%) showed “mild” dissections.

Stent length was similar in both groups (16.3 mm v 15.6 mm in groups A and B, respectively; p = 0.64).

CORONARY DISSECTION SEVERITY AND CORONARY FLOW

For both groups, baseline and hyperaemic averaged peak velocity values before and after the procedure are given in fig 2. The preinterventional hyperaemic response was slightly impaired in the group A patients randomised to stenting. After balloon angioplasty, both groups showed similar baseline and hyperaemic averaged peak velocity values. In patients randomised to additional stent implantation, no differences in baseline and hyperaemic averaged peak velocities were seen.

Absolute and relative CFVR values are given in table 3. Before and after balloon angioplasty, absolute CFVR was similar in the patient population as a whole. However, in the subgroup randomised to stenting, the absolute CFVR was lower in group A than in group B. As the non-diseased reference vessel CFVR was also significantly lower in group A than in group B (2.43 (0.71) v 2.91 (0.78), p < 0.001), the resulting relative CFVR after stent implantation was similar in the two groups.

CORONARY DISSECTION SEVERITY AND CLINICAL OUTCOME

Complete follow up data were obtained in all patients. Thirty three cardiac events occurred. There were five deaths (one in group A and four in group B) and six myocardial infarcts.
At 12 months, the rate of target lesion revascularisation and major adverse cardiac events was higher in group A than in group B (table 4). Event-free survival at 12 months was 78% in group A vs 89% in group B (p = 0.041).

All variables for which univariate analysis yielded a significant difference (age, smoking, degree of lesion calcification) were entered into the multivariate logistic model as potential predictors of major adverse cardiac events, along with the presence of moderate dissections. The presence of moderate dissections was the only independent predictor of major adverse cardiac events (odds ratio 2.42; 95% CI, 1.06 to 5.57; p = 0.036).

**Impact of Stenting on Moderate Dissections**

Among group A patients, those randomised to stent implantation had a higher rate of major adverse cardiac events than those randomised to stopping the procedure (31% vs 19%). The Breslow-Day test for homogeneity of odds ratios between the two patient subgroups was significant, showing a higher risk of major adverse cardiac events at 12 months of follow up in the stent arm (odds ratios 6.603 vs 1.197, p = 0.046). The CFVR (< 2.5 vs > 2.5) before second randomisation did not affect the long term outcome (odds ratios 3.617 vs 1.410, p = 0.287).

**Discussion**

The main findings of our study are that uncomplicated moderate dissections after balloon angioplasty left untreated had a good long term clinical outcome. In agreement with these results, several angioplasty studies have described a lack of association between the

| Table 3  Coronary flow velocity data |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                | **Group A**     |                | **Group B**     |                |
|                | Stent group     | Balloon group  | Stent group     | Balloon group  |
|                | n=13            | n=32           | n=111           | n=100          |
| DS before BA (%) | 76 (7)          | 69 (11)        | 68 (10)†        | 68 (11)        |
| DS after BA (%) | 23 (6)          | 23 (7)         | 22 (10)         | 21 (8)         |
| DS after stenting (%) | 7 (9)         | 8 (8)          |                 |                |
| CFVR before BA | 1.29 (0.32)*   | 1.55 (0.52)    | 1.64 (0.61)     | 1.64 (0.60)    |
| CFVR after BA | 2.04 (0.59)*   | 2.65 (0.95)    | 2.50 (0.74)     | 2.50 (0.73)    |
| CFVR after stenting | 2.33 (0.87) | 2.88 (0.30)†  |                 |                |
| RCFVR before BA | 0.57 (0.21)    | 0.57 (0.19)    | 0.60 (0.25)     | 0.58 (0.21)    |
| RCFVR after BA | 0.86 (0.17)    | 1.00 (0.28)    | 0.89 (0.27)     | 0.89 (0.27)†   |
| RCFVR after stenting | 0.97 (0.24) | 1.02 (0.30)    | 1.02 (0.30)     |                |
| Reference CFVR | 2.43 (0.71)*   | 2.83 (0.81)    | 2.91 (0.78)†    | 2.90 (0.75)    |

Values are mean (SD).

*p < 0.05 vs balloon group A; †p < 0.05 vs same subgroup in group A.

BA, balloon angioplasty; CFVR, coronary flow velocity reserve; DS, percentage diameter stenosis; group A, patients with moderate dissections; group B, patients with no or minimal dissection; RCFVR, relative CFVR.
presence of a moderate dissection and the long term clinical outcome. Previous three dimensional intracoronary ultrasound data from our group have shown that the presence of intimal dissection is associated with a greater total vessel volume at long term follow up, probably owing to favourable remodelling. It is conceivable that the development of a more deeply seated injury reduces vessel wall strength, thereby predisposing to favourable remodelling.

Overall, the patients with moderate dissections had a worse outcome than those with no or minimal dissections. This probably reflects the findings in the stented subgroup, which had a higher risk of major adverse cardiac events than the group with moderate dissections but without additional stenting. Although stents were placed according to the DEBATE II protocol and not to the operator’s preference, the small sample size (n = 13) prevents us from drawing any firm conclusions. While coronary stenting after the development of moderate dissections has proved to reduce the acute complication rate, there is no established long term clinical benefit of additional stenting in patients with uncomplicated moderate dissection (TIMI 3 flow and absence of signs or symptoms of angina). Recent reports have shown that coronary stent implantation causes more severe injury and a greater inflammatory response, as well as worse endothelial dysfunction, than plain balloon angioplasty. It is conceivable that the combination of a deep arterial injury associated with a moderate dissection and the implantation of a metallic body may cause a synergistic proliferative response. In patients with moderate dissections, preventing favourable remodelling might offset the beneficial stent scaffolding effect. The latter could be particularly important as these patients are expected to develop an enhanced neointimal response.

Our group has previously reported a temporary reduction in absolute coronary flow reserve in patients who developed uncomplicated moderate dissections. This reflected a transient increase in the baseline velocity. Therefore the reduced coronary flow reserve values did not translate into a greater residual stenosis and obstruction of coronary blood flow. In the present study, we used the relative CFVR—a more reliable index of persistent conduit obstruction than the absolute CFVR—and found similar values between the two groups. The latter results indicate that without signs or symptoms of ischaemia and the presence of TIMI 3 flow, no significant obstruction to coronary blood flow should be expected in patients experiencing mild to moderate dissections. Although the use of long or multiple stents has recently been reported to be associated with a greater restenosis risk, in our study similar lengths and numbers of stents per patient were used in the two groups, so this is unlikely to be a confounding factor.

LIMITATIONS

The limited number of patients with moderate dissections prevents us from drawing definitive conclusions. However, this is the largest prospective study investigating a selected population of patients who developed uncomplicated moderate dissections and underwent randomisation to additional stenting or no further treatment, and in whom post-procudural Doppler flow data and one year follow up data were available.

CONCLUSIONS

Moderate dissections left unstented do not have an adverse clinical outcome. Additional stenting does not appear to be of benefit.

Table 4. Clinical outcome at 30 days and 12 months of follow up in the group as a whole and in the two subgroups.

<table>
<thead>
<tr>
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<th>Whole group (n=256)</th>
<th>Stent group (n=124)</th>
<th>No further treatment group (n=132)</th>
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<td>(n=111)</td>
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<td>p Value</td>
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<tr>
<td>MACE (30 days)</td>
<td>5 (5%)</td>
<td>3 (7%)</td>
<td>NS</td>
</tr>
<tr>
<td>MACE (12 month)</td>
<td>23 (11%)</td>
<td>10 (22%)</td>
<td>0.041</td>
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<tr>
<td>TLR (12 month)</td>
<td>14 (7%)</td>
<td>8 (17%)</td>
<td>0.006</td>
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Values are n (%).

MACE, major adverse cardiac events; TLR, target lesion revascularisation.

Obliteration of a coronary artery aneurysm with a covered coronary stent

A 71 year old man with a history of limiting stable exertional angina who had required admission for unstable angina on two occasions in the preceding year underwent elective coronary angiography. Risk factors included hypertension, non-insulin dependent diabetes, hypercholesterolaemia, and previous smoking. Angiography showed single vessel disease of the left anterior descending artery (LAD) (top: right anterior oblique 30° caudal 10°). Just distal to the first diagonal there was a discrete coronary aneurysm with a severe stenosis proximally. After pre-inflation with a 2.5 mm balloon a 16 × 3.5 mm JoMed Jostent coronary stent graft was deployed at 18 atm, achieving excellent flow in the LAD and aneurysm obliteration (bottom: right anterior oblique 45° caudal 10°). The patient was discharged on a one month course of clopidogrel and remains asymptomatic at follow up.

The Jostent coronary stent graft is a polytetrafluoroethylene (PTFE) stent sandwich ideally suited for treatment of coronary aneurysms, perforations, and fistulae.

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