Temporary stent as a bail-out device during percutaneous transluminal coronary angioplasty: preliminary clinical experience

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Abstract

Objective—To evaluate the safety and efficacy of a prototype temporary stent (RX Flow Support Catheter, Advanced Cardiovascular Systems) in maintaining coronary perfusion and improving vessel patency in the event of acutely compromised flow complicating percutaneous transluminal coronary angioplasty.

Design—Prospective clinical study as part of a multicentre trial.

Setting—Regional cardiac centre catheterisation laboratory.

Patients—Eight patients undergoing routine percutaneous transluminal balloon coronary angioplasty in whom coronary artery dissection resulted in impaired coronary artery flow with angina or electrocardiographic ST segment shift, needing bail-out treatment at the time of the procedure.

Results—The RX Flow Support Catheter was successfully used and improved coronary flow in all cases, with a reduction in luminal stenosis and resolution of symptoms. The temporary stent was expanded for an average of 85 (range 30-209) minutes. In six patients it was used as a bridge to further treatment (permanent stent in four and coronary artery surgery in two) and two patients did not need further treatment.

Conclusion—The temporary stent was safe and effective in the acute management of coronary dissection. The main advantages are its ease and speed of use, and successful restoration of coronary flow both to the distal artery and to affected side branches pending definitive treatment.

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Coronary artery dissection at the site of balloon angioplasty results in abrupt vessel closure in between 3% and 8% of coronary angioplasties. Various bail-out strategies have been devised to manage these problems. These include prolonged inflation with either an angioplasty balloon or an autoperfusion balloon to stabilise the dissection and thus limit ischaemia, permanent stents to hold the vessel open, laser balloon angioplasty to tack up the dissection flap, and directional atherectomy to excise the dissection flap.

These strategies have reduced the need for emergency coronary artery surgery in the management of abrupt closure. The results of emergency coronary artery surgery are compromised by the presence of acute myocardial ischaemia on arrival in the operating theatre. This problem has been partially improved by autoperfusion catheters and the use of permanent stents for bail-out.

There remains a need for a bail-out device that ensures high coronary blood flow both to the distal artery and affected side branches and that is simple to use for operators with limited or no experience with permanent stents. As part of an international multicentre study, we have investigated a prototype temporary stenting device, (RX Flow Support Catheter, Advanced Cardiovascular Systems, USA). We evaluated the safety and effectiveness of this temporary stent in maintaining coronary perfusion and improving vessel patency when flow is compromised after coronary angioplasty.

Patients and methods

RX FLOW SUPPORT CATHETER

This consists of a temporary stenting catheter that provides an internal scaffold to the artery wall (fig 1). The catheter is used over an angioplasty guidewire 0.014 inches in diameter by a rapid exchange technique. The tip of the catheter is composed of a radio-opaque braided wire mesh cage (30 mm long) and is permanently attached to a 3.7 French catheter. There is a radio-opaque marker at each end of the cage.

A screw action manipulator on the proximal end of the catheter is used to expand the cage to the desired size and collapse it as required. As the cage expands it shortens. The cage can be expanded to any size up to a maximum diameter of 4 mm and exerts a low radial force (maximum 300 mm Hg) on the vessel wall. Because blood flows through the mesh cage, side branches that have been covered by the cage remain perfused. A side port is provided to maintain slow continuous flushing with heparinised saline during expansion of the cage. During the procedure the
PATIENTS

Eight patients (mean age 58.7 (range 41–69) years; six men) underwent the insertion of the RX Flow Support Catheter. Four patients had single vessel disease and four had multivessel disease. They were selected from those patients undergoing routine coronary angioplasty in whom there was acute impairment of coronary flow to TIMI Thrombolysis in Myocardial Infarction grade 2 or less after balloon inflation or a poor angiographic result needing intervention to prevent a flow-impeading condition and in whom symptoms or electrocardiographic changes were present. Four patients had chronic stable angina and four had unstable angina. All patients were pretreated with aspirin, and received 15 000 IU heparin and 500 ml 10% dextrose at the time of angioplasty. Patients were candidates for coronary artery surgery. Patients were excluded if they were undergoing angioplasty to the left main stem or an ostial stenosis or if the vessel was larger than the maximum size of the cage.

Patients gave informed consent to the use of the temporary stent which was approved by the ethics committee of the Royal Brompton National Heart and Lung Hospital.

STUDY PROTOCOL

Before the temporary stent was used, the angioplasty balloon was withdrawn leaving the angioplasty guidewire across the lesion. The temporary stent was then introduced over the angioplasty guidewire. The cage was positioned across the stenosis and then expanded to the diameter of the artery. A control angiogram confirmed the position of the cage: the cage was collapsed and re-expanded at a different site if the first position was not optimal.

During the study blood pressure and the electrocardiogram were continuously monitored. Anticoagulation was monitored by measuring the activated clotting time immediately before insertion of the stent and at 30 minute intervals thereafter. Fluoroscopy, and when necessary angiography, was used to monitor the position of the stent every five minutes.

After an initial expansion for 30 minutes the wire cage was collapsed and withdrawn into the guide catheter and the state of the vessel was reassessed by angiography. The operator then decided whether to re-expand the cage for longer or abandon the procedure. After the second period of expansion, which lasted up to 30 minutes, the cage was again withdrawn into the guide catheter and the angiography repeated. If the result was satisfactory no further treatment was undertaken. If the result was unsatisfactory then the operator decided either to insert a permanent stent or send the patient for coronary artery surgery. In those patients sent for surgery the cage was re-expanded to maintain coronary flow until after cardiopulmonary bypass had been established in the operating theatre.

Creatine kinase (CK) and its MB fraction

cage can be expanded and collapsed as required to alter its position. At the end of the procedure the cage is collapsed and the catheter withdrawn leaving the guidewire in place as appropriate.
Individual patients who underwent the insertion of a flow support catheter (FSC)

<table>
<thead>
<tr>
<th>Vessel</th>
<th>Diameter (mm)</th>
<th>Flow before FSC (TIMI grade)</th>
<th>Duration of FSC expansion (min)</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>LAD 2-75</td>
<td>3</td>
<td>0</td>
<td>50</td>
<td>No more treatment</td>
</tr>
<tr>
<td>RCA</td>
<td>3</td>
<td>2</td>
<td>33</td>
<td>Stent</td>
</tr>
<tr>
<td>LAD</td>
<td>2-5</td>
<td>0</td>
<td>209</td>
<td>Coronary artery surgery</td>
</tr>
<tr>
<td>LAD</td>
<td>3</td>
<td>2</td>
<td>30</td>
<td>Stent</td>
</tr>
<tr>
<td>LAD</td>
<td>3</td>
<td>2</td>
<td>60</td>
<td>Double stent</td>
</tr>
<tr>
<td>LAD</td>
<td>3</td>
<td>3</td>
<td>60</td>
<td>Stent</td>
</tr>
<tr>
<td>LAD</td>
<td>2-5</td>
<td>2</td>
<td>60</td>
<td>No more treatment</td>
</tr>
<tr>
<td>LAD</td>
<td>3</td>
<td>1</td>
<td>180</td>
<td>Coronary artery surgery</td>
</tr>
</tbody>
</table>

FSC, RX Flow Support Catheter; LAD, left anterior descending coronary artery; RCA, right coronary artery; Stent, permanent stenting with a Palmaz-Schatz Stent (Johnson and Johnson, USA); Double stent, indicates the use of tandem permanent stents.

were measured in blood drawn before angioplasty, eight hours and 24 hours after the operation and before discharge from hospital. Twelve lead electrocardiograms were recorded at the same times.

All patients were reviewed three months after the procedure. If patients had not undergone further treatment after the use of the RX Flow Support Catheter they underwent repeat coronary angiography.

Results
Seven patients had dissections in the proximal or mid-left anterior descending coronary artery and one patient in the mid-right coronary artery. The table shows the details of the vessels treated. Four patients had a coronary artery dissection partially impeding anterograde coronary artery blood flow, three patients had a dissection causing abrupt closure, and one patient without impairment of flow had a large coronary artery dissection that was considered likely to cause occlusion. In seven of eight cases the dissection involved a side branch of the artery. Six patients had angina associated with the impaired perfusion and four patients had ST segment changes on their electrocardiogram. The RX Flow Support Catheter was used for an average (range) of 85 (30–209) min.

USE OF CATHETER
The RX Flow Support Catheter was prepared and used in all patients in less than six minutes. Preparation of the catheter took less than three minutes. The cage was expanded to the natural diameter of the artery in four patients and to 0.5 mm greater than the natural diameter in four patients. Angiogram showed that the catheter was not optimally positioned in seven of eight patients and the cage was repositioned. In one patient impaired flow developed after 10 minutes. This was corrected by minor repositioning without any adverse consequences.

SAFETY
During the studies the patients remained haemodynamically stable. Once coronary perfusion had been restored. Anticoagulation as measured by the activated clotting time was maintained in the range 300–750 s, additional boluses of 5000 IU heparin were given as required.

IMMEDIATE RESULTS
The result of expansion of the cage of the flow support catheter produced TIMI grade 3 flow to the distal artery with reduction in the luminal stenosis to less than 10% in all patients. Also, four patients in whom the dissection sites involved side branches had TIMI grade 3 flow both down the main artery and the affected side branch. There was relief of angina and resolution of electrocardiographic changes in all patients while the temporary catheter was in place.

EARLY OUTCOME
After expansion of the cage a satisfactory angiographic result was obtained in two patients who needed no further treatment. In six patients (in one of whom there was an improvement in luminal stenosis) the angiographic result was inadequate: mean TIMI grade before temporary stenting was 1.7 and after removal it was 2.3. Four of these patients had a permanent Palmaz Schatz stent (Johnson and Johnson, USA) inserted, all with a satisfactory result. Figures 2 to 6 show the angiographic sequence of one of these patients.

Two particularly long expansion times of 180 and 209 minutes were needed when the temporary stent was used as a bridge to coronary surgery after dissection of the left anterior descending coronary artery. One of these arteries was considered to be too narrow for insertion of a permanent stent and the length of the dissection precluded permanent stenting in the other. In both cases vessel patency could not be maintained without
arrived in the operating theatre without chest pain or electrocardiographic changes. Flow support was maintained until cardiopulmonary bypass had been established and the surgeon was ready to graft the affected coronary artery. Both patients received internal mammary grafts and one patient also received a vein graft to the first diagonal branch, which was affected by the coronary dissection and in which flow had been maintained by the flow support catheter.

On withdrawal of the catheter, a piece of debris about 1 mm in diameter was found in the cage in two patients. Histology of one of these cases showed that the debris was fibrin. No patient sustained a myocardial infarction as indicated by electrocardiographic changes or a rise in the CK or CK-MB fraction during their stay in hospital.

**LATE OUTCOME**

Follow up at three months showed that six of the eight patients remained symptom free with no objective evidence of active myocardial ischaemia on the resting and exercise (no data in two patients) electrocardiograms. Of the two patients with recurrent angina, one underwent angioplasty to a different vessel one month after flow support; and one patient who had a definitive result with the flow support catheter alone underwent repeat angioplasty that showed a 99% restenosis with TIMI grade 1 flow. The second patient with a definitive result from the flow support catheter refused repeat angioplasty because he felt well. The patients who underwent coronary surgery had normal left ventricular function with no regional wall motion abnormalities as assessed by echocardiography and gated blood pool scanning.

**Discussion**

Coronary artery occlusion after percutaneous transluminal coronary angioplasty is a serious complication. It is associated with a high incidence of myocardial infarction and increased mortality compared with uncomplicated coronary angioplasty. As abrupt vessel closure after coronary angioplasty is unpredictable, effective bail-out devices are necessary to minimise myocardial ischaemia.

Despite a wide range of methods and devices to overcome this problem in the catheter laboratory, about one third of patients in this situation require emergency coronary artery surgery. The results of this are less good than elective surgery, and have an incidence of Q wave myocardial infarction varying from 25% to 57% and mortality from 1-4% to 19%. The poor results seem to be associated with preoperative ischaemia that also mitigates against the use of internal mammary artery grafts. Attempts to reduce ischaemia with an autoperfusion catheter have shown a reduction in perioperative myocardial infarction and an increased use of internal mammary artery grafts. Strategies to stabilise or reverse myocardial ischaemia may improve long-term outcome.
We have used the prototype flow support catheter to restore and maintain TIMI grade 3 coronary blood flow in eight patients with either acute impairment of coronary flow or a poor angiographic result with threatened vessel closure after percutaneous transluminal coronary angioplasty.

In animal studies with prototype flow support catheters regional myocardial blood flow was not significantly different in left ventricular subepicardial and subendocardium perfused by coronary arteries with or without the catheter in place. In our study, this catheter seems to have specific advantages as a bail-out device. Compared with autoperfusion balloons, the ability to alter cage size means that one catheter size will fit any artery. The wire mesh cage is less obstructive than a balloon and there is TIMI grade 3 distal flow. Unlike perfusion balloons, side branches are also perfused. Both the lower flow and occlusion of side branches may be reasons why perfusion balloons do not completely salvage the ischaemic myocardium.

Unlike permanent stents, which are not clearly better than conventional treatment in the management of abrupt vessel closure, the flow support catheter is suitable for vessels less than 3 mm diameter. The RX Flow Support Catheter is easily seen on fluoroscopy and can be repositioned if required. When the flow support catheter is used in place of a permanent stent it avoids the problems of anticoagulation and subacute thrombosis seen with permanent stents.

In our preliminary experience, the RX Flow Support Catheter was effective at reversing ischaemia but less successful at providing definitive treatment during a one hour expansion. An unexplored use of this device might be to provide coronary perfusion for a prolonged period, a technique that has been used with catheter perfusion to avoid subsequent bypass surgery.

In conclusion, we found that the prototype RX Flow Support Catheter was safe and simple to use after an unsatisfactory result from coronary balloon angioplasty. It was successful as a bail-out device in all eight patients. It restored normal coronary flow both to the distal artery and to side branches, reduced luminal stenosis, and resolved angina and electrocardiographic changes. Its effectiveness as a bail-out device may add a new dimension of security to coronary angioplasty.

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