Is transluminal coronary angioplasty mandatory after successful thrombolysis?

Quantitative coronary angiographic study

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SUMMARY Percutaneous transluminal coronary angioplasty has been advocated as a mandatory procedure to prevent reocclusion after successful thrombolysis in acute myocardial infarction. This study describes our experience with both procedures over a 12 month period. Out of 105 patients catheterised in the acute phase of myocardial infarction, 64 were recanalised with 250 000 units of streptokinase, while in 25 patients recanalisation could not be achieved. In the remaining 16, the infarct related vessel was patent at the time of the procedure. Eighteen of the 78 patients who had a patent infarct related vessel at the end of the recanalisation procedure underwent transluminal angioplasty immediately afterwards. Post lysis angiograms were analysed quantitatively with a computerised measurement system. The contours of the relevant arterial segments were detected automatically. Reference diameter, minimal obstruction diameter, length of the lesions, and percentage diameter stenosis were averaged from multiple views.

In 31% of our patients a diameter stenosis of less than 50% was found whereas one of 70% or more was seen in only 19%. Eleven stenotic lesions, recanalised at the acute stage, reoccluded in the short term, and in the long term eight other patients sustained a reinfarction in the same myocardial territory. Seventeen of these 19 recanalised lesions had a diameter stenosis of 58% or more. In view of these results, we felt justified in combining recanalisation and angioplasty in 18 patients selected from the most recent admissions. In these patients, the mean diameter stenosis decreased from 59% to 30% and mean pressure gradient from 41 to 8 mmHg. Late follow up showed reocclusion in one case. Though percutaneous transluminal coronary angioplasty does not seem to be mandatory at the acute stage in the majority of patients, it is feasible to undertake in one sitting and seems to prevent reocclusion in patients selected on the basis of quantitative angiographic criteria.

Salvage of ischaemic but still viable myocardium around areas of myocardial infarction is currently a topic of much clinical interest. The removal of obstruction of the nutrient artery by intracoronary thrombolysis in the first hours after the onset of myocardial infarction has proved a new approach which is undergoing randomised trials in our own and other institutions. Since there is often residual stenosis, additional transluminal angioplasty and/or coronary artery bypass grafting have been advocated as a mandatory procedure after successful recanalisation. It has been argued that a severe residual stenosis in the area of the previous occlusion might cause reocclusion over the ensuing days. In order to eluci-

date this question, quantitative angiographic analysis was applied to recanalised vessels of 78 patients who had an open infarct related vessel at the end of the procedure. Tentative answers were formulated on three questions: How severe are the residual lesions after ‘successful’ thrombolysis? Is it possible to identify those lesions that are liable to reocclude in the short term? In order to prevent reocclusion after initial successful recanalisation, is transluminal angioplasty a mandatory procedure?

Patients and methods

Between September 1980 and December 1982, coronary recanalisation was attempted in 105 patients. Different procedures were used since our first experi-
ence with intracoronary thrombolysis in September 1980. They have been described elsewhere. In the current randomised trial, patients below 65 years of age were selected, without a history of haemorrhagic diathesis or previous cerebrovascular accident. On admission all patients suffered from chest pain lasting less than four hours. The electrocardiogram showed typical myocardial infarction with ST elevation. The combination of hypotension (systolic pressure below 90 mmHg) and sinus tachycardia (heart rate over 100 beats/minute) led to temporary exclusion, but if the haemodynamic condition of the patient returned to normal quickly, he was still able to be included in the study. Informed consent was obtained from all patients assigned to thrombolytic treatment. Immediately after admission, an infusion of glyceryl trinitrate was started and as soon as possible the patient was transferred to the catheterisation laboratory. Prophylactic lignocaine was given intravenously in a dose of 2 mg/min.

**TECHNIQUE OF INTRACORONARY STREPTOLYSIS**

After puncturing the femoral vein and artery a pacemaker catheter was positioned in the right ventricle. Heparin, 50 mg, was administered intravenously as well as 250 mg acetylsalicylic acid and 100 mg diadresone F. A non-ionic contrast agent (Amipaque) was used as a contrast medium for coronary angiography of the artery suspected to be thrombosed; subsequently 0.2 mg nifedipine was injected into the occluded artery over a period of three minutes, while the aortic pressure was monitored. Coronary angiography was then repeated to evaluate the spasmolytic effect on the coronary occlusion. Intracoronary perfusion with streptokinase was carried out at a rate of 4000 units a minute to a maximum of 250 000 units of streptokinase, diluted in 500 ml physiological solution at a flow rate of 8 ml per minute. Coronary angiograms were repeated every 15 minutes until the chest pain disappeared. The appearance of ventricular extrasystoles or any conduction disturbance was an additional reason to visualise the artery. If there were no signs of recanalisation, an attempt was made to administer streptokinase locally in a higher concentration by passing a thin catheter (French 2 or 3) with a radiopaque tip through the Judkins catheter (French 8). After the procedure, selective coronary angiography in multiple projections was performed with an ionic contrast medium (urografin 76%).

All arteriograms were recorded on Kodak 35 mm cinefilm at the rate of 25 frames/s. The stenotic areas were filmed in two different projections in stenoses of the right and left circumflex coronary artery and in at least three projections, including one craniocaudal, in stenoses of the left anterior descending artery.

**TECHNIQUE OF PERCUTANEOUS TRANSLUMINAL CORONARY ANGIOPLASTY**

In 18 patients an attempt was made to dilate the residual stenosis. In 16 patients, percutaneous transluminal coronary angioplasty was performed in the same session, 20 to 60 minutes after the end of streptokinase infusion. In two other patients, percutaneous transluminal coronary angioplasty was performed in a second session, respectively eight hours and 12 days later. Via a 9F, 16 cm long introducing sheath, a guiding catheter was directed into the stenotic area under fluoroscopic and pressure control. In four of the 18 successfully treated cases, we used a balloon catheter with an outer diameter of 3.7 mm, in the 14 other patients its outer diameter was 3 mm. The mean pressure gradient across the stenotic lesion was computed on line after 20 seconds of data acquisition. Two to nine (mean 4.5; SD 2.3) balloon inflations were performed for a period of 10 to 70 seconds (mean 49; SD 12) at a pressure of 4 to 8 bar (mean 6; SD 1). After the dilatation procedure the sheath was left in place for the next 24 hours, while the patients were monitored in the coronary care unit.

All patients received heparin (20 000 U/24h) followed by oral coumarin until discharge from the hospital. Anticoagulants were continued after discharge in patients with left ventricular aneurysm or recognised mural thrombus in the left ventricle. In addition, nifedipine, 10 mg every four hours, was given for two days in patients treated with streptokinase.

**QUANTITATIVE ANGIOGRAPHIC ANALYSIS**

The quantitative analysis of selected coronary segments was carried out with the help of a computer based coronary angiography analysis system, which has been described extensively elsewhere. To analyse a selected 35 mm cineframe, the film was placed on a specially constructed cine video converter. An optically magnified portion of the cineframe was converted into video format with a high resolution video camera and displayed on a video monitor; routinely, an optical magnification factor of 2 is used. The computerised analysis of a selected coronary segment required the manual definition of a number of centre positions within the segment by means of a writing tablet. Regions of interest encompassing the arterial segment were digitised and stored in the PDP 11/44 minicomputer. Subsequently, the contour positions of the segment were detected automatically on the basis of first and second derivative criteria. From these contour positions the diameter function of the segment was computed. The diameter values are given in absolute values (mm), since the contrast catheter was used as a scaling device (Fig. 1a).

As a next step the computer algorithm determined
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Fig. 1  (a) Contours for obtuse marginal branch superimposed on the original video image. From the detected contours the diameter function is determined by computing the shortest distance between left and right contour positions. The calibrated diameter values in mm are plotted along the ordinate and the centreline positions from the proximal to the distal part along the abscissa. (b) For the lesion in Fig. 1(a) the normal size of the artery has been estimated from the normal proximal and distal diameter values. The marked area is a measure of the atherosclerotic plaque. An interpolated percentage diameter stenosis of 66% results.

the position of the obstruction by searching for the minimal diameter value in the diameter function. This position could be changed interactively by the user, if more than one focal obstruction was to be processed within the analysed segment. The extent of the obstruction was computed from the diameter function on the basis of curvature analysis and expressed in mm.

The severity of a coronary obstruction was determined by the interpolated percentage diameter stenosis measurement. The basic idea behind this technique is the computer estimation of the original contour positions over the obstructive region (assuming there was no coronary disease present) based on the course of the proximal and distal centreline segments and on the diameter function. The resulting reference contours of the arterial segment of Fig. 1a are shown in Fig. 1b, with the difference in area between this boundary and the detected contours marked over the obstructive lesion; this area is a measure of the atherosclerotic plaque. The interpolated percentage diameter stenosis was then computed by comparing the minimal diameter value at the obstruction...
with the corresponding value of the reference contour at this position. For Fig. 1b an interpolated percentage diameter stenosis of 50% results. It is clear that the computed reference contour can only be a reasonable approximation of the non-diseased segment if the proximal and distal segments are essentially free of atherosclerosis. Even if this is not true this method may still be the method of choice in the majority of cases for the determination of percentage diameter stenosis, because it is minimally influenced by any observer error.

Results

PATENCY OF INFARCT RELATED VESSEL, ACUTE AND CHRONIC STAGE
The angiographic findings at the beginning and at the end of the recanalisation procedure are given in Fig. 2. In 64 patients, thrombolysis was successful and after recanalisation transluminal angioplasty was performed in 18 of these 64 patients. In 16 other patients the infarct related vessel was found to be patent at the first coronary angiogram. In two of the 16 cases, occlusion of an initially patent artery occurred during the procedure. Thus, 78 infarct related vessels were patent at the end of the catheterisation procedure. In 25 patients we did not succeed in recanalising the thrombosed arteries.

Fifty seven patients, excluding those who had undergone percutaneous transluminal coronary angioplasty in the acute phase, agreed to have further angiograms two weeks later (Fig. 3). In 41 patients, the infarct related vessel was still patent. Five lesions which had remained occluded at the acute stage were found to be recanalised at this point; in seven other patients the coronary arteries which had been successfully recanalised at the acute stage were now found to be reoccluded. This observation suggests a reocclusion rate of 17% two weeks after recanalisation.

Moreover, in the long term follow up (mean 8.3 months) eight other patients sustained a reinfarction in the same myocardial territory, which was documented with the electrocardiogram and cardiac enzymes: two of them were restudied angiographically and in one patient the infarct related vessel had reoccluded. Finally, four lesions which had remained occluded at the acute stage were still occluded at the chronic stage.

QUANTITATIVE ANGIOGRAPHIC ANALYSIS
The individual data of the quantitative analysis of 75 stenotic lesions are given in Fig. 4. Three lesions could not be analysed because of the poor quality of the angiograms. Each depicted value represents the average value of measurements obtained in different angiographic projections. The median value for the reference diameter is 2.98 mm whereas the 10th and 90th centiles are 2.22 mm and 4.20 mm respectively; the median value for the minimal obstruction diameter is 1.32 mm, and the values of the 10th and 90th centiles are 0.78 mm and 1.88 mm, respectively. As for the length of the lesion, the median value is 9 mm.
while values of the 10th and 90th centiles are 5 and 16mm, respectively. Fig. 5 is a histogram of the percentage diameter stenosis measured on 75 stenotic lesions after the successful thrombolysis. At the acute stage, the median value for the percentage diameter stenosis is 58% in this group of 75 patients; the values of the 10th and 90th centiles are 37% and 74%, respectively. A percentage diameter stenosis less than 50% is measured in 31% of the patients, whereas a diameter stenosis greater or equal to 70% is seen only in 19%. When comparing the percentage diameter stenosis in a subgroup of 26 individual lesions, analysed in identical projections in the acute and chronic stage, no significant improvement or deterioration of the recanalised lesions at the chronic stage could be demonstrated (Fig. 6).

There are, of course spectacular individual changes in diameter stenosis, but for the whole group the percentage diameter stenosis does not change significantly: 54 ± 16% at the acute stage versus 48 ± 14% at the chronic stage.

The percentage diameter stenosis of eight recanalised lesions in patients who initially underwent a successful recanalisation and subsequently sustained a reinfarction in the same myocardial territory is shown in Fig. 7. Six of eight stenotic lesions had a percentage diameter stenosis greater than or equal to 58% (the median value) immediately after thrombolysis. The second column represents 11 stenotic lesions which reoccluded at the acute stage (n=4) or at the chronic stage (n=7) after they had been recanalised. All of them had a percentage diameter stenosis greater than or equal to 58%. The diameter stenoses of 14 stenotic lesions found to be patent at the first coronary angiogram during the acute phase of their myocardial infarction are also given. Nine of them had a diameter stenosis less than 58% (median value).

From these data it appeared that reocclusion and recurrent myocardial infarction were more frequent in patients with a stenosis greater than 58% in diameter after recanalisation. The angiographic results of the combined procedure performed in 18 patients are also shown. Though the selection and decision to perform angioplasty had been based on a visual evaluation of the severity of the lesion at the time of the procedure, it appears retrospectively that 13 of the 18 lesions had a stenosis greater than 58% in diameter after thrombolysis. In these 18 patients angioplasty decreased the average percentage diameter stenosis from 59% (SD: 9-9%) to 30% (SD: 9-9%) and the minimal obstruction diameter increased from 1-3 mm (SD: 0-4 mm) to 2-2 mm (SD: 0-3 mm). This reduction in diameter stenosis was highly significant (p<0-0001) and was associated with a significant decrease in mean pressure gradient from 41 mmHg (SD: 17 mmHg) to 8 mmHg (SD: 5 mmHg) (Fig. 8). On the basis of the changes in pressure gradient, dilatation had not been necessary in two of these patients.

Fifteen out of the 18 patients who underwent percutaneous transluminal coronary angioplasty were
restudied angiographically after a mean follow up of four months (range from 10 days to 11 months). In all but one the dilated vessel was patent. One patient developed exertional angina three months later. A restenosis at the site of the previous occlusion was successfully dilated a second time.

**Discussion**

In this paper it is shown that intracoronary streptokinase infusion and percutaneous transluminal coronary angioplasty can be carried out in the same session in an effort to enhance reperfusion. The issue, however, is not whether these combined interventions can be performed safely in the setting of acute infarction, but rather whether one is justified in doing so.

The present work is a tentative answer to this question and its conclusions are based on the results of quantitative angiographic analysis of recanalised vessels.

Since at the end of the procedure, 31% of our patients had a recanalised vessel with percentage diameter stenosis below 50%, whereas a diameter stenosis greater than 50% was seen in the remainder, we had no evidence of a major residual thrombus at the site of the stenosis. No retention of contrast medium or staining of a distal thrombus was seen, so residual clot, reported by others, appears unlikely as well. Furthermore it has been the custom to continue the streptokinase treatment for about 30 minutes after patency has already been established.
On the other hand, severe residual stenosis of the recanalised vessel has been reported by other groups. Their reports are based on visual and therefore subjective interpretation of the stenotic lesions. Overestimation and excessive variations by intra- or interobserver error have been reported. These discrepancies led us to evaluate the exact condition of the stenotic lesions after successful lysis by an objective, computer aided, interpretation. This method has been validated not only with copper and perspex models filled with contrast medium but also from necropsy studies.

In 14 cases with clear clinical, electrocardiographic, and enzymatic signs of an acute infarction, the diseased vessel was patent at the time of coronary angiography. In these patients long lasting vasospasm superimposed on organic lesions may have led to transient occlusion and myocardial infarction. Another possibility is that the clot had already lysed in the meantime. In nine out of these 14 patients, the infarct related vessel had a mild stenotic lesion, a factor that might have facilitated the reopening of the vessel. This is the more likely as these patients were started on an intravenous perfusion of glyceryl trinitrate before intracoronary lysis was attempted. In patients with transmural acute myocardial infarction and coronary arterial thrombi, histological sections of coronary arteries have been shown to be narrowed by the atherosclerotic plaque alone from 33 to 98% (mean 81%) at the site of the thrombus. When histological cross-sectional areas are compared with cross-sectional areas derived from diameter measurements, our quantitative angiographic results in vivo are consistent with the histological findings. Furthermore, it is remarkable how closely the obstruction diameters after successful thrombolysis correspond with the postmortem findings reported by Fulton, as shown in Fig. 9. In one third of our patients the myocardial infarction had occurred when the relevant artery showed a stenosis of less than 50%. Gertz et al. have shown experimentally that endothelial damage and thrombus formation may occur at the site of focal arterial constriction even when the reduction in luminal diameter by itself is insufficient to alter significantly the rate of flow.

As shown by Fulton in his elegant study on the morphology of coronary thrombotic occlusions, atherosclerotic lesions are usually of a complex nature. In two thirds of his cases, a break or a tear in the luminal lining exposed blood flowing to the material composing the underlying lesion. He postulated that this was the probable cause of platelet aggregation and fibrin deposition. In half of his cases, a haemorrhagic dissection was found which resulted in an apparent reduction of the lumen. Successful thrombolysis could again expose the material of the atheromatous lesion to flowing blood.

Whether the thrombogenic activity of this material would be as active as it was in the first instance remains speculative but secondary thrombotic occlusion may occur. Accordingly, there are strong theoretical reasons for adopting antithrombotic measures after successful thrombolysis. On the other hand, instrumental dilation of the coronary artery stenosis underlying the occlusive thrombus might produce desquamation of endothelium and shearing of the superficial portion of endothelial plaque, possibly altering its thrombogenic propensity. Subsequent fibrosis and healing appear to cause further enlargement of the lumen and thereby may improve local rheology. Thus, arguably, it might be advisable to perform angioplasty even in patients with a stenosis of less than 50%.

Reported rates of reocclusion vary between less than 10% to 25%. In this study two weeks after thrombolysis, the reocclusion rate was 17%. Moreover, during follow up, eight patients sustained reinfarction in the same myocardial region, so that reocclusion and/or reinfarction might affect 25% of our patients who had undergone a successful recanalisation. Thus it seems rational at the acute stage to dilate recanalised arteries with a residual stenosis greater than 58% since these stenotic lesions are liable to early reocclusion. This concept was supported by our
observations that during streptokinase infusion, recoclusion occurred in four patients (Fig. 7) despite intravenous or intracoronary administration of glyceryl trinitrate or nifedipine. Here percutaneous transluminal coronary angioplasty might have been the only way to restore blood flow adequately and to prevent immediate reocclusion. As emphasised by Meyer et al., 11 the advantages of the combined procedure are savings in time and money. The same introducing sheath can be used, the catheterisation laboratory equipment and the personnel have to be used only once, and it appears to be less exciting for the patient. To date no reclosures have occurred during such immediate dilatation of the critical stenosis. Fifteen out of the 18 patients who underwent percutaneous transluminal coronary angioplasty were restudied angiographically after a median follow up of four months (range from 10 days to 11 months). In all but one patient, the dilated vessel has remained patent. All 17 patients have been followed at the outpatient clinic at three month intervals. The mean period of follow up has been seven months and the longest one year. None of these patients has sustained a reinfarction thus far.

These observations indicate that percutaneous transluminal coronary angioplasty immediately after thrombolysis is a safe and reasonable procedure when after lysis of the obstruction a 58% stenosis is still present. This combined approach seems to result in a lower rate of reclosure or reinfarction than thrombolysis alone. As further randomised trials are necessary to show the ultimate benefit of thrombolysis in acute myocardial infarction, it is recommended that during them the additional value of immediate percutaneous transluminal coronary angioplasty is investigated.

References

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