Transcatheter occlusion of coronary to bronchial anastomosis by detachable balloon combined with coronary angioplasty at same procedure

J F REIDY, E SOWTON, D N ROSS
From the Departments of Radiology and Cardiology, Guy's Hospital, London; and Harley Street Clinic, London

SUMMARY The first case of non-operative occlusion of a large coronary to bronchial anastomosis is described. The patient who had severe angina had also a critical stenosis of the anterior descending coronary artery which was successfully dilated by an angioplasty procedure. An occluding balloon was detached in the large distal circumflex coronary artery, beyond all the normal branches. At a repeat catheterisation study seven months later the balloon was intact and in position, and the anastomosis remained occluded.

There have been many descriptions of coronary artery to bronchial artery anastomosis published.1-3 Small anastomoses, presumably congenital in origin, have been noted to occur in about 20% of normal subjects in the absence of other abnormalities.1 Larger anastomoses have been seen in one of two situations: when there is significant coronary artery disease the bronchial arteries may communicate via collaterals with the coronary artery distal to the site of obstruction.1 The opposite situation occurs when there is severe right ventricular outflow tract obstruction and then the coronary to bronchial anastomoses communicate with the lower pressure pulmonary circulation. In both these situations the coronary to bronchial anastomoses are of secondary significance compared with the primary disease and no specific treatment is usually needed. In the case described here there was a large haemodynamically significant coronary to bronchial anastomosis that occurred as an isolated finding and not secondary to any other abnormality.

Much has been written on congenital coronary artery fistulae and in the majority of cases surgical correction has been advocated.4 We present a case of a patient with severe angina who was successfully treated by the procedure of combined coronary angioplasty and transcatheter occlusion of a large haemodynamically significant coronary artery to bronchial artery anastomosis. To our knowledge this is the first case of its kind to be described and this now offers a new form of treatment for these uncommon but significant anastomoses.

Case report

The patient is a man of 67 who was referred by Dr Palencia Perez in Valencia for possible surgical treatment. His history was of admission to hospital with acute coronary insufficiency including unstable ST segment elevation on the electrocardiogram, widespread T wave changes suggesting anterior subendocardial infarction, and repeated episodes of pain. He was treated with bed rest, isosorbide, nifedipine and metoprolol, and in between the episodes of pain his electrocardiogram always returned to normal. The symptoms settled without any evidence of actual infarction and coronary arteriography was subsequently performed in Spain. This showed a severe stenosis of the proximal left anterior descending coronary artery and a large circumflex coronary artery to bronchial anastomosis. There was no clinical evidence of respiratory disease.

Clinical examination was normal. There were no detectable murmurs; electrocardiogram and chest x-ray film were normal.

In view of the patient's age and the type of lesions present it was decided to attempt a coronary angioplasty and embolisation of the circumflex anastomosis at the same procedure with full surgical standby in case of complications.

In May 1981 cardiac catheterisation was performed via the right percutaneous femoral approach. This again showed severe proximal stenosis of the anterior descending artery (>90%) but with no other occlusive...
Transcatheter occlusion of coronary to bronchial anastomosis

There is a severe proximal stenosis of the anterior descending branch (white arrow). Note the very large calibre of the circumflex artery, the last obtuse marginal branch (black arrowhead), and the distal artery just before it runs into the bronchial anastomoses (black arrow).

disease. The circumflex artery was of very large calibre. After it gave off a distal obtuse marginal branch it continued as a large and tortuous artery behind the heart (Fig. 1). Passing in a superior direction it terminated in a leash of small vessels situated in the region of the left hilum and the bronchial arteries (Fig. 2). There was no evidence of any communication with pulmonary arteries. Left ventricular angiography and pressure studies were all normal.

Both the angioplasty and the embolisation procedures were performed via a 9F non-tapered coronary guide catheter (Schneider Medintag Ag). The angioplasty was performed first and a gradient of 105 mmHg was abolished with a 3-7 mm diameter dilatation catheter.

The same introducer catheter was then manipulated into the proximal circumflex artery and the embolisation achieved using the Becton-Dickinson detachable balloon catheter.* When safely positioned in the distal circumflex coronary artery the balloon was inflated with contrast medium while a test injection was made through the coronary guide catheter. When this confirmed both a good position of the balloon and showed that it was occluding the artery, it was detached by gentle traction. The silicone balloon was inflated with iso-osmotic contrast medium. There

* Becton-Dickinson, Rutherford, New Jersey.

Fig. 1 Left coronary arteriogram in the right anterior oblique projection. There is a severe proximal stenosis of the anterior descending branch (white arrow). Note the very large calibre of the circumflex artery, the last obtuse marginal branch (black arrowhead), and the distal artery just before it runs into the bronchial anastomoses (black arrow).

Fig. 2 Left coronary arteriogram in the left anterior oblique projection showing the most distal part of the circumflex coronary artery where it runs to join the bronchial arteries. Note the most terminal part (white arrow) that corresponds with the black arrow of Fig. 1.
were no complications and the patient was discharged from hospital four days later.

The patient was seen at follow up seven months after the angioplasty and embolisation. He has been completely free of symptoms and able to take vigorous exercise. In December 1981 a follow up cardiac catheterisation was performed. The left coronary arteriogram showed a little irregularity at the site of the anterior descending angioplasty but no stenosis (Fig. 3). The balloon was still intact and completely occluding the distal circumflex artery and fistula. The patient remains symptom free in August 1982, 15 months after the procedure.

Discussion

The first case of a coronary to bronchial anastomosis demonstrated on selective coronary arteriography was in 1972 in a patient who had normal coronary arteries but had evidence of obstructive airways disease. Two patients shown at coronary arteriography to have large coronary to bronchial anastomoses were also shown to have no coronary artery disease, but both of these patients had bronchiectasis as indicated by bronchography. One of these patients had severe angina associated with electrocardiographic changes and was treated by lower lobectomy combined with ligation of the anastomotic vessel.

The other patient who had no history of angina and only mild respiratory symptoms was managed by conservative treatment.

Our patient presented with severe angina but had no evidence of respiratory disease. In addition to the large coronary to bronchial anastomosis arising from the circumflex artery, there was a severe and critical stenosis of the proximal anterior descending coronary artery. The remaining coronary arteries were normal. The bronchials filled from the left coronary injection and there was no evidence of collateral vessels between the circumflex artery and the distal anterior descending to suggest that there was a bronchial to coronary “steal”. In the absence of any evidence of respiratory disease, the coronary to bronchial anastomosis appears to be a primary and presumably congenital anomaly.

The very large size of the circumflex coronary artery and the distal branch passing to the bronchial artery territory suggest that this anastomosis was of functional significance.

Thus our case had two clear-cut abnormalities that could cause his angina. As the proximal anterior descending stenosis was very suitable for an angioplasty procedure, it was decided to occlude the large coronary to bronchial anastomosis with a detachable balloon at the same occasion and via the same catheter approach and introducing catheter. Of all the techi
Transcatheter occlusion of coronary to bronchial anastomosis

ques available for occluding vessels, the detachable balloon was the only one that was suitable for this situation.

Detachable balloons have been used in many vessels but only recently in the cardiopulmonary situation. As the large branch arose distal to the origin of all the obtuse marginal branches of the circumflex, it was essential to occlude the vessel distal to these. Passing a catheter down into the distal circumflex presents many problems but using the method whereby the small uninflated balloon could be injected out along with the forward blood flow proved an ideal method. The balloon was filled with iso-osmotic contrast media, and there is evidence that if a vessel can be occluded for a minimum of 10 days then the vessel will remain occluded. To our knowledge this is the first reported case of deliberate occlusion of a coronary artery vascular lesion via a percutaneous transcatheter approach.

We are grateful to Dr RI White for advice and encouragement that enabled us to start using the detachable balloon occlusion system.

References


Requests for reprints to Dr John F Reidy, X-Ray Department, Guy's Hospital, St Thomas Street, London SE1 9RT.