Feasibility of percutaneous transluminal angioplasty for carotid artery stenosis

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Abstract
Percutaneous transluminal balloon angioplasty was attempted in seven patients with internal carotid artery stenosis, including one patient who had two procedures. All had recurrent, carotid territory, neurological symptoms considered haemodynamic in origin. Six had occlusion of the contralateral internal carotid artery. Cerebral blood flow studies confirmed diminished cerebrovascular reserve in six patients studied. In five patients (six procedures) angioplasty of the stenosed internal carotid artery was carried out successfully. With two patients technical difficulty in crossing the stenosis prevented angioplasty and in one patient with bilateral stenosis the procedure was not attempted on the second side because of the severity of the stenosis. In two patients transient aphasia developed during cannulation of the carotid artery and in another a transient monoparesis developed during the procedure. Both these haemodynamic complications recovered within ten minutes. No other complications occurred. Our experience suggests that balloon angioplasty is technically feasible in the management of stenotic carotid disease associated with haemodynamic stroke. It is a technically simple procedure requiring only a brief admission to hospital. However, its general application to patients with thromboembolic carotid-territory stroke will depend on the risk/benefit ratio compared to carotid endarterectomy or to conventional medical treatment.

The treatment of stenotic cerebrovascular disease is controversial. The value of carotid endarterectomy is disputed and the indications for surgery are uncertain.1,2 The usual aim of endarterectomy is to remove atheromatous sources of thromboembolism but, in some patients, particularly those with high-grade stenosis or contralateral internal carotid artery occlusion, the aim may be to improve cerebral perfusion pressure. However, carotid endarterectomy carries significant risks. The reported risk of death or permanent stroke following carotid endarterectomy varies from 21% to 2%, in different series.3,4 The average combined mortality and stroke risk between different centres has been estimated at around 6%.5 The major risk is of stroke in the territory of the operated artery but there is also significant morbidity from stroke in vascular territories not directly related to the operated artery and from myocardial infarction, particularly in patients with ischaemic heart disease or severe hypertension.6 There are also local risks, such as cranial nerve palsy and anaesthetic risks, such as pulmonary embolism.7 Because of these problems we have investigated the feasibility of percutaneous transluminal angioplasty (PTA) in the management of selected patients with symptomatic stenosis of the internal carotid artery. This procedure is brief, and does not require a surgical incision, or general anaesthesia. Since the development of appropriate balloon dilation catheters, PTA has become established in the treatment of peripheral, renal and coronary vascular disease. Its success rates in these situations approaches 90%, complications occurring in less than 5% of procedures.8,9

In contrast, angioplasty of vessels supplying the intracranial circulation has not been recommended because of anxiety about the risks of cerebral embolism following the procedure.10,11 However, web-like stenosis of the common carotid artery has been successfully dilated with a catheter passed retrogradely through an arteriotomy, performed for endarterectomy at the carotid bifurcation, and by PTA.12,13 Fibromuscular dysplasia of the internal carotid artery14,15 and of brachial vessels, including the subclavian, vertebral and external carotid arteries,16-18 has also been successfully treated with PTA. With increasing experience of PTA at these and other sites, a few reports of PTA performed successfully for atherosclerotic stenosis of the internal carotid artery have appeared suggesting that PTA might be used in place of carotid endarterectomy in selected cases, in contrast to earlier opinion.19-23 In this report we describe our initial experience of PTA in haemodynamic stroke due to internal carotid artery stenosis.

Subjects and methods
Patients were offered internal carotid PTA as an alternative to carotid endarterectomy if they
had a history of recent appropriate cerebrovascular events and if initial investigations showed smooth stenotic lesions of the internal carotid artery without evident ulceration or thrombosis. Patients were only considered for PTA if they were felt to be at high risk of recurrent stroke and if they would otherwise have been offered open carotid endarterectomy. Patients with angiographic or clinical features which increased the risk of carotid endarterectomy were also included.

The seven patients (table 1) who fulfilled these criteria gave fully informed consent to the procedure as required by the Ethical Committee of The London Hospital. Their ages ranged from 46-64 years (mean 55 years). Four were female and three male. Four had significant athrosclerotic stenosis at the origin of one internal carotid artery associated with complete occlusion of the contralateral internal carotid artery (cases 1, 3, 4 and 5). One had severe internal carotid artery stenosis associated with only minor stenosis of the contralateral internal carotid artery (case 6). Another patient had an irregular, short, 95% segmental stenosis of the upper cervical internal carotid artery secondary to fibromuscular dysplasia, associated with complete occlusion of the contralateral internal carotid artery due to dissection (case 2), whilst the seventh patient had bilateral severe stenoses of the internal carotid arteries. Two patients presented with minor strokes (cases 1 and 6) and five presented with recurrent transient ischaemic attacks (TIAs). In the latter, TIAs had continued despite aspirin or anticoagulant therapy. Three patients gave a history consistent with a haemodynamic origin for their symptoms in that TIAs occurred in association with assumption of the erect posture, or during a hot shower. One patient, with recurrent TIAs despite anticoagulant therapy, that occurred nine months after a technically successful angioplasty, had repeat angioplasty on the same artery (case 5).

In six patients there was diminished cerebrovascular reserve in the symptomatic hemisphere as determined by a significantly reduced cerebral blood flow response to hypercapnia (table), measured by the non-invasive Xenon131 technique.25 Five patients had medical conditions which were felt to increase the risks of surgery. Two of these had severe ischaemic heart disease, one had poorly controlled hypertension, one had rheumatoid arthritis and one had such severe ankylosing spondylitis of her cervical spine that endarterectomy was impossible. In one patient with fibromuscular dysplasia the site of the stenosis was surgically inaccessible.

Protocol

All the procedures were undertaken by one of us (PB). Before angioplasty, each patient was fully anticoagulated with warfarin for between two and four weeks. The procedure was performed under sedation with droperidol 5 mg iv, and fentanyl 0·1 mg iv. The anticholinergic action of droperidol was valuable in countering...
any reflex bradycardia resulting from manipulation of the carotid bifurcation. Heart rate, blood pressure and the electrocardiogram were monitored throughout the procedure.

A “Check-Flo” 7.5 French gauge introducer sheath (Wm Cook, Europe A/S) was inserted via the right femoral artery. Preformed 5 French gauge “Headhunter” (HNI) or “Sidewinder” (Simmons II) balloon dilation catheters were used (Wm Cook, Europe A/S). Each balloon was 1.5 cm long and could be inflated to a diameter of 5 mm at a maximum pressure of 3 atmospheres. The catheter tip extended for 1 cm beyond the balloon. Catheter placement was assisted by a 0.028” diameter Teflon-coated straight guidewire with a 5 cm floppy tip. Just before balloon inflation, arterial blood was withdrawn into a heparinised 50 ml syringe from the sidearm of the introducer sheath. Whilst the balloon was inflated this blood was injected at a rate of approximately 100 ml/min through the catheter lumen to maintain some perfusion of the hemisphere rostral to the temporarily occluded internal carotid artery.

Up to three dilations, each of 40 seconds duration and at least two minutes apart, were performed using an “Indeflator Plus” (Advanced Cardiovascular Systems Inc) to monitor the exact inflation pressure. During angioplasty each patient was asked to report the development of any symptoms to a neurologist in attendance. In addition, the neurologist conducted a limited neurological examination (speech, facial and extremity movement and plantar responses) at frequent intervals. Any change in the neurological examination was an indication for immediate withdrawal of the catheter.

**Results**

Internal carotid PTA was carried out successfully in five of the seven patients (table). In three of these the procedure was carried out entirely without complications. Two patients developed transient contralateral symptoms during balloon inflation which resolved fully after ten minutes. One of these patients had a repeat angioplasty performed on the same artery nine months later without complication. There was no undue haemorrhage at the
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Discussion

Our experience shows that percutaneous angioplasty (PTA) of the internal carotid arteries is a feasible alternative to endarterectomy and can be successfully carried out without major complications in suitable patients. PTA has major advantages over endarterectomy in terms of resource allocation and patient comfort, and in avoiding general anaesthesia. The procedure itself is technically simple and there is immediate ascertainment of any neurological complications since the patient remains conscious throughout. No specialised nursing care is required and the patient can be discharged 24 hours after angioplasty.

Our seven patients were carefully selected. All had recurrent TIAs or strokes with continued symptoms despite medication with aspirin or anticoagulants. They were considered to be at high risk for further stroke. Three had a history which strongly suggested a haemodynamic origin for their symptoms. Six had high internal carotid artery stenosis associated with occlusion of the contralateral internal carotid artery and the six who had blood flow studies had evidence of impaired cerebrovascular reserve suggesting a haemodynamic origin for their symptoms. Improvement in perfusion pressure by dilating the internal carotid artery was therefore likely to be beneficial. All had relatively smooth stenotic lesions of the internal carotid artery of a type for which we would usually recommend endarterectomy. Six had at least one feature thought to increase the risk of carotid endarterectomy, for example, contralateral carotid occlusion, ischaemic heart disease or severe hypertension. In one patient the stenosis was surgically approachable. Repeat PTA was performed on the same artery in one patient without complication.

The risks of internal carotid PTA are as yet undefined. Our series of seven patients extends the total published case reports to 23 in which internal carotid artery PTA has been attempted. Two of our patients developed transient neurological symptoms during balloon inflation lasting less than 10 minutes. One other instance has been reported in which a patient developed a transient ischaemic attack 20 minutes after the procedure but no major complications had been reported. The most serious theoretical risk of PTA is cerebral embolism due to dislodgement of thrombus or atheromatous material from the vessel wall. We therefore avoided patients with angiographic features of atheromatous ulceration or thrombus in the carotid artery and regarded impaired cerebrovascular reserve as a more appropriate indication for PTA than potential amelioration of thrombogenic pathology. To reduce the risk of embolisation further, our patients were anticoagulated with warfarin for two weeks before angioplasty. Theron et al. used temporary distal balloon occlusion of the internal carotid artery in five patients to prevent embolisation during PTA of more proximal atherosclerotic lesions. However, in our patients with haemodynamic insufficiency the duration of total occlusion of the internal carotid artery required by that technique (up to 10 minutes) would have considerably increased the risk of the procedure.

Laboratory and necropsy studies of angioplasty for atherosclerotic lesions suggest two main pathophysiological mechanisms responsible for the increase in vessel diameter. If, when distended, the balloon diameter is slightly greater than that of the stenotic section, stretching of the vessel wall occurs with superficial desquamation of intimal elements. Superficial splits may occur in atherosclerotic plaque. Healing of the intima with endothelialisation results in a larger lumen. If the balloon diameter when distended is much greater than that of the stenosis, then deep focal splits occur in the intima and plaque leaving the relatively elastic media intact. Over the subsequent few weeks the intima then retracts and the elastic media, freed from the encasing effects of the intima, distends. In some cases splits may extend through the media, as in one of our patients who developed an asymptomatic false aneurysm, but arterial rupture has not been described. The splits may lead to the angiographic appearance of intimal dissection, but this is limited to the area of the angioplasty and only rarely leads to delayed secondary occlusion. Compression or redistribution of
atheromatous material does not appear to play an important part in the response to angioplasty. Embolism of damaged intima or plaque material appears to be uncommon in peripheral PTA but splits in the intima or plaque provide obvious sites for thrombus formation and subsequent post-angioplasty thrombotic occlusion or embolisation. In our protocol, therefore, anticoagulation was continued for four to six weeks after PTA. Others have used aspirin or diprydamole or a short course of heparin following carotid PTA.

We were particularly concerned that haemodynamic cerebral ischaemia might occur in our patients during temporary occlusion of the internal carotid artery by the inflated balloon. We therefore limited the period of balloon inflation to 40 seconds and waited at least 120 seconds before repeating it. In order to minimise the fall in pressure in the cerebral arteries, the balloon was inflated to the patient’s maximum tolerated internal carotid pressure. Because of the lack of evidence that the small calibre catheters used, technical difficulties were encountered in several cases. Modifications to the catheter-guide wire systems may improve selective internal carotid cannulation. The success rate of carotid PTA is likely to improve with increasing experience.

Inflation of the balloon did not stimulate the carotid sinus since there was no associated change in pulse or blood pressure. This may reflect the site of the stenoses, which were all distal to the carotid bifurcations. Reflex bradycardia and hypertension may be a problem in patients with common carotid or bifurcation stenosis. Appropriate premedication should be used in such cases. It has been suggested that stretching the carotid artery may induce reflex local arterial spasm but we did not encounter this complication. A local infusion of papaverine or general anaesthesia may relieve this spasm.

A major advantage of carotid PTA over endarterectomy is that the procedure is carried out in an alert and cooperative patient. Apart from avoiding the risks of general anaesthesia this allows continuous monitoring of the patient's neurological state, with immediate deflation of the balloon and withdrawal of the catheter in the event of complications. Evoked potential monitoring could also be used but we have preferred to use clinical examination because it is sensitive to changes within the middle cerebral artery territory and is directly relevant to the clinical problem.

The role of PTA in the treatment of carotid stenosis is as yet undefined. We suggest that PTA of the internal carotid artery is most suitable for patients with smooth lesions in whom haemodynamic mechanisms may be responsible for ischaemic cerebral symptoms. This is particularly likely in patients with continuing symptoms following carotid occlusion. The failure of the international extracranial-intracranial arterial bypass study to show any benefit of bypass surgery even in patients with carotid occlusion has redirected interest towards endarterectomy of contralateral stenotic lesions as a means of improving perfusion pressure in both hemispheres. Our experience shows that PTA provides a feasible alternative. PTA is also suitable for treatment of surgically inaccessible high lesions of the internal carotid artery, such as fibromuscular dysplasia and for the treatment of symptomatic restenosis of the carotid artery following endarterectomy, where surgical re-exploration is hazardous. It may also be indicated for restenosis following angioplasty. Many strokes in patients with cerebrovascular disease result from thrombo-embolism, but the potential role of PTA in this situation is less clear than in haemodynamic stroke. The analogy with angioplasty for coronary artery disease, however, suggests that PTA may be appropriate. Larger, long term studies are needed to define the appropriate indications, risks and outcome of PTA for carotid stenosis. Whether, and in which circumstances, the final balance will favour angioplasty will depend on its risk/benefit ratio compared to carotid endarterectomy and to conventional medical treatment.

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