Reactivity of the cerebral circulation in patients with carotid occlusion

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SUMMARY Cerebral blood flow (CBF) and the response to hypercapnia (cerebral reactivity) have been measured in 41 patients with unilateral or bilateral internal carotid artery occlusion in an attempt to identify those with limited collateral reserve. Normocapnic CBF was within normal limits in the majority of subjects. The response to hypercapnia varied from normal to absent, with impaired reactivity becoming increasingly likely when more than one artery was diseased. In 19 patients with unilateral carotid occlusion, hemisphere reactivity was well preserved in the majority, but was significantly lower on the side of the occlusion (mean 2.9%/mm Hg) compared to the normal side (mean 3.4%/mm Hg). Reactivity on the side of the occlusion was further reduced in 15 patients with occlusion and contralateral internal carotid artery stenosis (mean 1.7%/mm Hg) and was even lower in seven patients with bilateral occlusion (mean 1.1%/mm Hg). There was no difference in reactivity between asymptomatic hemispheres in the 41 patients (mean 2.7%/mm Hg) and hemispheres in which a previous stroke had occurred (mean 2.8%/mm Hg). In contrast the response in hemispheres subject to continuing transient ischaemic attacks was significantly impaired (mean 1.6%/mm Hg), suggesting that the cerebral symptoms in some of these patients may have had a haemodynamic origin more often than suspected from the clinical history.

Some patients with occlusion of the internal carotid artery suffer a major stroke, but others present with a minor stroke or a transient ischaemic attack (TIA); occasionally occlusion of one or both carotid arteries occurs without any symptoms. It might be thought that once a carotid artery has occluded completely the risk of further stroke or TIA on the side of the occlusion is over. However, some patients continue to experience delayed vascular events ipsilateral to established carotid occlusion and controversy persists as to the cause of these episodes. Barnett drew attention to coincidental proximal external carotid and common carotid disease and suggested that this could act as a potential source of emboli to the retina or hemisphere via the collateral circulation. Indeed, haemodynamic factors appeared clinically important in only two of the 27 patients described. Two subsequent studies have emphasised the importance of coincidental external carotid artery disease. A further potential source of emboli is thrombus within the stump of the proximal internal carotid with embolisation via anastomotic channels. On the other hand, some patients undoubtedly have symptoms and signs which point to haemodynamic insufficiency and Sundt and Whisnant have suggested that ischaemic symptoms in the region of an occluded artery are usually haemodynamic in origin and therefore an indication for bypass surgery.

Although the brain is supplied by four major arteries and has efficient routes for collateral blood flow, occlusion of one internal carotid artery produces a fall in perfusion pressure in middle cerebral artery branches of the ipsilateral hemisphere. The initial response is vasodilatation of the small resistance arteries with a consequent fall in vascular resistance and the maintenance of perfusion rates (the phenomenon of autoregulation). The degree of vasodilation required to preserve normal blood flow depends on the resistance of the patent cerebral arteries, the circle of Willis and the collateral vessels. When the vascular bed is maximally dilated, any further reduction in mean pressure will lead to a passive fall in blood flow.

Carbon dioxide (CO₂) is a potent vasodilator of the normal cerebral circulation. However, if vaso-
dilatation is already maximal the capacity to increase flow in response to hypercapnia is limited.\(^{18}\) The degree of cerebral CO\(_2\) reactivity can therefore provide an index of the capacity of the cerebral circulation for further vasodilatation (cerebrovascular reserve). We have measured CBF and CO\(_2\) reactivity in 41 patients with occlusive carotid artery disease in an attempt to identify those with limited cerebrovascular reserve and have related the results to the extent of large vessel disease and to the presence or absence of continuing ischaemic symptoms. Because of the wide range of CO\(_2\) reactivity in normal subjects and in those with cerebrovascular disease we have used patients as their own controls by comparing reactivity in the hemisphere on the side of the occluded carotid artery with that of the contralateral hemisphere.

Patients

Forty-one patients found to have occlusion of one or both internal carotid arteries were studied. The majority of the subjects were ambulant outpatients and none had had a recent major completed stroke. Most subjects were referred by vascular surgeons during the consideration of carotid artery or bypass surgery. They were divided into the following three groups according to the results of recent angiography (digital subtraction venous imaging in the majority). All had patent vertebral arteries.

Unilateral occlusion Nineteen patients with a total occlusion of one internal carotid artery in whom the contralateral internal carotid artery was normal or had minor irregularities only. This group had a mean age of 61.1 (SD 6.6) with a range of 51–75 years.

Oclusion with contralateral stenosis Fifteen patients in whom total occlusion of one internal carotid artery was associated with 50% or greater stenosis of the internal carotid artery on the contralateral side. This group had a mean age of 64.1 (SD 8.1) with a range of 48–76 years.

Bilateral carotid occlusion Seven patients with total occlusion of both internal carotid arteries. This group had a mean age of 65.1 (SD 11.7) with a range of 41–74 years.

Neurological symptoms

All the subjects were questioned at the time of the CBF study about neurological symptoms. Seven subjects had no history of previous cerebral or ocular ischaemia. Most of those had been found to have carotid artery occlusion during the course of investigation for cardiac or peripheral vascular disease. The remainder were divided into the following two groups:

(a) Previous stroke Fourteen patients had a history of previous stroke (including reversible ischaemic neurological deficits) and five a history of previous TIA's, but all had been free of hemisphere symptoms for at least 2 months. The mean age of this group was 60.1 (SD 8.0) years.

(b) Continuing symptoms Twenty patients were classified as having continuing neurological symptoms if they had had symptoms appropriate to the carotid circulation on the side of the occlusion within the previous 2 months. Eight had amaurosis fugax and 12 hemiplegic or hemisensory TIA's. Many patients had frequent attacks; in two the symptoms were clearly associated with change of posture, in three others the association was less certain and in the remainder the attacks occurred at random. The mean age of this group was 63.6 (SD 7.9) years.

Methods

CBF was measured by the non-invasive intravenous xenon\(^{133}\) (Xe\(^{133}\)) technique described by Thomas et al.\(^{19}\) In brief, a bolus of about 7 mCi of Xe\(^{133}\) dissolved in saline was injected into an arm vein of the recumbent subject and the clearance of the isotope from the cerebral hemispheres monitored by six external 25 mm diameter scintillation detectors. Expired concentrations of Xe\(^{133}\) were monitored by a 7th detector and the end tidal levels used to estimate the concentration of recirculating arterial Xe\(^{133}\). Regional CBF was calculated for each detector from a one minute initial slope analysis of the clearance curves. Results presented in this paper for each hemisphere are means of the initial flow values calculated from three detectors placed over the frontal, fronto-parietal and temporoparietal regions.

Blood pressure was recorded from the left arm with a standard mercury sphygmomanometer.

Arterial partial pressure of carbon dioxide was estimated from the mean end tidal expired concentration of CO\(_2\), monitored throughout the recordings by a Datex CD 300 infrared capnograph.

Each patient had two measurements of CBF within 2 hours separated by at least 45 minutes to allow for complete elimination of the first dose of Xe\(^{133}\) between measurements. The first CBF was determined with the patient breathing room air (normocapnia) and the second with the patient breathing 5% CO\(_2\) in air (hypercapnia). CO\(_2\) was included in the breathing circuit for at least 2 minutes before the CBF measurement.

![Fig 1 Mean normocapnic hemisphere cerebral blood flow (CBF) ± SD in three groups of patients related to the state of the ipsilateral carotid artery (N = normal, O = occluded, S = stenosed, R = right, L = left). *p < 0.01 (paired t test).](http://jnnp.bmj.com/)

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measurement to allow for stabilisation of end-tidal CO₂ concentrations, and was continued throughout the CBF measurement. This technique produced an average sustained rise of 11 mm Hg (1.5 kPa) in arterial pCO₂ between measurements.

Hemisphere CO₂ reactivity was calculated as the percentage rise in CBF above normocapnic values per mm Hg increase in pCO₂ (1 mm Hg = 0.13 kPa).

**Results**

Figure 1 illustrates mean hemisphere CBF during normocapnia according to the patency of the ipsilateral carotid artery in each group. Mean hemisphere CBF was similar in all three groups. There was a tendency for CBF to be lower on the side of an occlusion compared with the contralateral hemisphere, but the difference was small and only significant in the group with contralateral stenosis (mean difference 1.6 ml/100 g/min, paired t = 2.88 p < 0.02). Mean CBF was also slightly lower in the patients with bilateral occlusion compared with the other two groups but the difference was not significant (mean 3.7 ml/100 g/min lower than patent hemispheres, Student's t = 1.47).

Figure 2 shows the mean response to hypercapnia in each group and fig 3 shows the resulting values for mean hemisphere CO₂ reactivity. In patients with unilateral occlusion hemisphere reactivity was normal on the side of the patent carotid artery (mean 3.4%/mm Hg), but was significantly reduced on the side of the occlusion (mean 2.9%/mm Hg, paired t = 2.41, p < 0.05). In the patients with occlusion and contralateral stenosis mean reactivity was lower in both hemispheres and again was significantly reduced on the side of the occlusion (mean 1.7%/mm Hg) compared with the contralateral hemisphere (mean 2.6%/mm Hg. Student's t = 3.18 p < 0.01). This impaired response to hypercapnia on the side of the occlusion in patients with contralateral stenosis was significantly lower than that of the normal hemisphere (Student's t = 3.33, p < 0.01) and occluded hemispheres of the patients with unilateral occlusion (Student's t = 2.56, p < 0.02). The patients with
bilateral carotid artery occlusion had even greater impairment of the response to hypercapnia with mean hemisphere reactivity of only 1.1%/mm Hg, which was also highly significantly lower than that of the normal hemispheres (Student's t = 8.94, p < 0.001) and occluded hemispheres of the patients with unilateral occlusion (Student's t = 8.08, p < 0.001). Figure 2 illustrates that the rise in pCO2 was similar in each group and the results could not be explained by changes in blood pressure (BP) which rose during hypercapnia in the majority of subjects (mean normocapnic BP 167/91; mean hypercapnic BP 183/99).

In all groups there was a wide range in hemisphere reactivity from normal to absent responses, but impaired reactivity on the side of an occlusion became increasingly likely with increasing severity of contralateral carotid artery disease. For example, figures of less than 1.5%/mm Hg were found on the side of the occlusion in 11% of patients with unilateral occlusion, 60% of patients with occlusion and contralateral stenosis and in at least one hemisphere in 86% of the patients with bilateral occlusion.

Figures 4 and 5 illustrate the relationships between mean normocapnic CBF, hemisphere reactivity and the patients' symptoms. There was no significant difference between normocapnic CBF in asymptomatic hemispheres, hemispheres with a history of previous ipsilateral stroke or hemispheres with continuing ipsilateral symptoms. There was also no difference in mean CO2 reactivity between asymptomatic hemispheres and hemispheres with a previous stroke. However, reactivity in hemispheres subject to continuing symptoms was significantly impaired compared with both asymptomatic hemispheres (Student's t = 2.76, p < 0.01) and hemispheres with a previous stroke (Student's t = 2.59, p < 0.02). This contrast was most striking in the patients with bilateral occlusion, among whom patients with continuing symptoms had a mean cerebral reactivity of only 0.4%/mm Hg compared with a mean of 2%/mm Hg in those not currently symptomatic. The individual reactivity values on the side of an occlusion are illustrated in fig 6 for each group according to whether the patient had continuing ipsilateral symptoms or not. Fifty-five per cent of patients with continuing symptoms had hemisphere reactivities of less than 1.5%/mm Hg, compared with only 19% of asymptomatic or previous stroke patients. Of the two
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patients in whom TIAs were clearly precipitated by changes in posture, one had the lowest reactivity encountered (CBF actually falling by 0.8%/mm Hg during hypercapnia) and the other had a reactivity of only 1.2%/mm Hg. Comparing individual hemispheres, it was found that an asymmetrical reduction in the symptomatic hemisphere compared to the contralateral hemisphere was even more closely associated with continuing symptoms. Reactivity on the ipsilateral side was less than 75% of the contralateral hemisphere in 65% of patients with continuing symptoms but in only 3 (14%) patients who were asymptomatic or had previous stroke.

Discussion

These studies demonstrate that at normal levels of arterial pCO₂ autoregulatory mechanisms are able to maintain nearly normal resting CBF in the presence of unilateral carotid occlusion, which is in accord with previous studies.20–25 There was a tendency for hemisphere CBF to be slightly lower on the side of the occlusion compared with the other hemisphere but the difference was only significant if the contralateral artery was stenosed. In contrast, when the collateral reserve was tested by CO₂ inhalation an impaired CBF response was revealed on the side of the occlusion in many patients. Such impaired CO₂ reactivity became increasingly likely when more than one carotid artery was diseased.

The response to hypercapnia on the side of the normal carotid artery in patients with unilateral occlusion was preserved with a mean reactivity of 3.4%/mm Hg, which is close to the figure of 3.1%/mm Hg found in a group of patients with cerebrovascular symptoms but normal carotid arteries by Bullock et al.25 using a similar method. In comparison with these normal hemispheres CO₂ reactivity was significantly lower on the side of the occlusion (mean 2.9%/mm Hg) in the patients with unilateral occlusion, but was still adequately preserved in many patients. Impaired reactivity was more common in the patients in whom carotid occlusion was associated with contralateral carotid stenosis and the mean reactivity was more significantly reduced on the occluded side at 1.7%/mm Hg. The most striking abnormalities were found in the patients with bilateral occlusion. Although this group had a normocapnic CBF which was only slightly lower than normal, CO₂ reactivity was grossly impaired bilaterally in all but one patient and the mean reactivity of 1.1%/mm Hg was lower than in the other two groups. These results confirm findings from earlier studies where compromised reactivity has been related to the extent of contralateral arterial disease.22 23 25

A failure to respond to hypercapnia implies either insensitivity to the metabolic stimulus or that the capacity to increase CBF is limited because vaso-dilatation is already maximum. The latter explanation has been suggested by previous studies.20–22 24–27 It has also been invoked as the mechanism behind the loss of CO₂ reactivity which occurs under conditions of severe hypotension18 and may also explain loss of reactivity in the vicinity of acute infarction.28 We suggest that our finding of a graduated reduction in reactivity with increasing severity of carotid artery disease reflects increasing distal vasodilation. This conclusion agrees well with the studies of Gibbs et al.29 who reported an increase in cerebral blood volume distal to carotid occlusion, reflecting focal dilatation, which was greatest in the patients with severest arterial disease. It is also consistent with the finding of Norrving et al23 that impaired reactivity in patients with carotid occlusion correlated with angiographic evidence of inadequate collateral supply.

More intriguing than the morphological correlate is our finding that there was a close association between continuing TIAs and a loss of responsiveness to CO₂. The present data suggest that continuing cerebral symptoms after carotid occlusion may have a haemodynamic origin more often than suspected from the history since TIAs were clearly related to change of posture in only two of our 20 subjects with recurrent symptoms. However, our study was based on a highly selected group of patients and it remains uncertain how often haemodynamic factors are important in the majority of patients with delayed ischaemic events.

On superficial analysis, surgical revascularisation would appear to be a logical treatment for patients with haemodynamic symptoms or impaired cerebrovascular reserve. Baron et al.11 in a detailed study of one patient with posturally induced hemispheric symptoms, reported a dramatic clinical improvement with concomitant normalisation of measurements of critical perfusion, following a successful extracranial/ intracranial (EC/IC) bypass. On the other hand, even if a subgroup of patients with haemodynamic TIA can be identified it does not necessarily follow that surgical revascularisation is indicated because the symptoms may resolve with time,30 presumably as collaterals enlarge. Furthermore, there is recent evidence that in general EC/IC bypass does not protect against further stroke in patients with internal carotid occlusion (EC/IC Bypass Study Group31—preliminary results). However, the protocol of the EC/IC study group32 did not request measurements of cerebral blood flow, nor was there any other systematic attempt to differentiate between embolic and haemodynamic TIA. The present study has demonstrated that cerebrovascular reserve is likely to

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be reduced in patients with recurrent TIAs and multiple vessel disease. It should prove possible to use the EC/IC data base to determine if surgery reduces the chance of stroke in this particular sub-group.

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