Causes of ischaemic stroke in the young

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Stroke is the third most common cause of mortality in Westernised countries and accounts for 12% of all deaths in the UK. The economic cost of stroke is enormous – approximately 4–5% of the annual National Health Service budget.1 Twelve per cent of first strokes occur in patients under 45 years of age,2 of which approximately 50% are ischaemic in nature (cerebral haemorrhage is relatively over-represented in this age group compared to the elderly).

Reported incidence rates of ischaemic stroke in the young vary according to study design and population structure. The annual incidence of young stroke in the UK has been estimated at approximately 10 per 100 000 (female:male, 1.6:1)3 in a prospective, community-based study. Other Western European studies have provided crude incidence rates up to three times higher.4-6

Most reports of young stroke apply an age limit of 45 years to the study population. Over this age, incidence rises sharply and the spectrum of underlying causes narrows as atherosclerosis becomes increasingly prevalent. Whereas degenerative atherosclerotic disease and cardioembolism account for most ischaemic strokes in the elderly, cerebral infarction in younger age groups may be the presenting feature of a diverse range of local and systemic diseases. Full evaluation of the young patient is likely to elucidate an underlying cause, many of which are treatable. The management of young stroke therefore requires a modified approach, encompassing initial investigation and treatment, as well as advice on prognosis and counselling for the devastating psychological consequences.

The clinical encounter

No matter which classification of stroke syndrome the clinician chooses to use, the clinical picture will reflect the anatomical distribution of brain damage and will, with occasional exceptions, mirror the stroke syndromes seen in the elderly. In history taking, it is important to enquire specifically about previous deep vein or pulmonary thromboses (coagulopathies), arthralgia (systemic lupus), skin rashes (vasculitis, antiphospholipid syndrome, Fabry’s disease), miscarriages (antiphospholipid syndrome), a family history of thromboses (inherited thrombophilies), or drug abuse. The clinician must also be clear about the nature of onset of the neurological deficit. It is not uncommon for multiple sclerosis to cause an isolated hemiparesis which may be wrongly attributed to vascular disease. However a hemiparesis due to demyelination usually develops over at least 24–48 hours rather than abruptly, and is often partial rather than complete as in most vascular events.

The clinical examination will focus on the nervous and cardiovascular systems. The presence and equality of all peripheral pulses must be sought (coarctation, subclavian stenosis, Takayasu’s arteritis). Auscultation of the heart should be followed by a search for carotid and subclavian bruits although the presence of the former shows poor accuracy in predicting a significant carotid stenosis.1 The cutaneous stigmata of hyperlipidaemias are usually readily apparent. Other features such as a Horner’s syndrome (carotid dissection), Marfanoid habitus (Marfan’s syndrome, homocystinuria), skin laxity and joint hypermobility (Ehlers-Danlos, pseudo oxanthoma elasticum), livedo reticularis (Sneddon’s syndrome), vasculitic rash, splinter haemorrhages, oral and genital ulcers (Behcet’s) and venulpuncture marks should not be overlooked.

Fundoscopy can provide important clues: papilloedema (cerebral venous thrombosis), Roth spots (subacute bacterial endocarditis), optic atrophy and retinitis pigmentosa (mitochondrial cytopathy), cholesterol emboli (cerebral stenosis) and signs of vasculitis (attenuated vessels, retinal haemorrhages, cotton wool spots, etc).

Investigations

Is the stroke ‘arterial’ or ‘venous’? Venous infarction arises from thrombosis of the dural sinuses and cortical veins. It is usually haemorrhagic with symptoms evolving over several days; headache, papilloedema, seizures and a fluctuating...
neurological deficit with depression of consciousness are common. Cerebral venous thrombosis has been reviewed previously. Which and how many arterial territories are involved? Infarction in multiple arterial territories often indicates a cardioembolic source, or the presence of a more diffuse process such as an arteritis. If there has been only one event, it is impossible to differentiate a peripheral from a cardiac source unless there are other clinical clues. However, multiple events within the same arterial territory should focus initial attention on the supplying vessel.

As technology progresses, so the range of diagnostic tests available increases (Box 1). Rather than subject each patient to investigations as part of a routine work-up it would seem better practice to tailor the investigations according to clinical pointers. Thus a patient with splinter haemorrhages and a cardiac murmur requires blood cultures and echocardiography whereas a patient with hemicranial pain and a Horner’s syndrome requires assessment of the carotid arteries to exclude dissection.

THE BRAIN
All young patients with stroke require at least a cranial computed tomography (CT) scan in order to exclude primary intracerebral or subarachnoid haemorrhage. CT should confirm the site of the infarct, although it may be normal within the first 24 hours of stroke onset. After the clinical encounter, CT, or preferably magnetic resonance imaging (MRI), is the next guide in determining whether future tests should concentrate on a given arterial territory, a cardiac cause or a systemic cause. Haemorrhagic transformation occurs commonly in cardiac embolism and venous infarcts, and multiple deep white matter infarcts are more likely to represent a systemic disorder.

The greater availability of CT compared to MRI dictates that the former will be performed first in most cases of young stroke. Although MRI better delineates areas of infarction in the acute phase, it is relatively poor at demonstrating acute parenchymal or intraventricular haemorrhage. If there is doubt about whether the CT lesion is due to infarction, MRI will often distinguish other pathologies. However, interpretation of small high signal lesions seen on T2 and proton density weighted scans requires care since these appearances can result from small vessel ischaemia (hypertension, arteritis) or demyelination. The latter pathology is suggested by a characteristic periventricular distribution, or the presence of similar lesions in the brainstem and cerebellar peduncles. MRI is the modality of choice for imaging lacunar, brainstem and posterior fossa infarcts. It is more likely than CT to show small subclinical infarcts.

THE HEART
Standard 12-lead electrocardiography and chest radiography are required. Unless it is certain that the cause of stroke lies outside the heart, transthoracic echocardiography (TTE) is indicated, although its yield in the absence of clinical signs is low. TTE provides an indication of left atrial and ventricular size and function plus a guide to the morphology and function of the mitral and aortic valves. Valvular vegetations may be imaged as may left ventricular mural thrombus. However, left atrial and a significant proportion of left ventricular thrombi will be missed - although clues such as dyskinetic segments or left atrial enlargement may suffice for the clinical decision to anticoagulate. TTE, like all ultrasound techniques, is operator and patient dependent, and because pathology is not demonstrable does not mean that it does not exist.

Although large atrial septal defects may be imaged by TTE, small defects such as patent foramen ovale are easily missed. The introduction of bubble contrast (5–10 ml of agitated saline or water) via a peripheral vein aids the detection of patent foramen ovale since bubbles appear to travel from the right to left atrium during the Valsalva manoeuvre in the presence of a right to left shunt. A similar technique can be used during transoesophageal echocardiography (TOE) which is the imaging modality of choice for imaging the aortic root, atria, and inter-atrial septum. Atrial septal aneurysms, patent foramen ovale, left atrial appendage thrombi and valvular vegetations are all more clearly visualised using TOE. In deciding to use TOE, the clinician must weigh the expected diagnostic yield against the availability and semi-invasive nature of the technique. It seems reasonable to proceed to TOE if TTE were technically unsatisfactory in demonstrating the cause of an abnormal clinical finding, if TTE shows a lesion which requires further characterisation, or if a careful search elsewhere has failed to elucidate an embolic source.
THE EXTRACRANIAL VESSELS
Ultrasound scanning is the screening modality of choice for the detection of extracranial carotid or vertebral artery disease. Modern systems provide high resolution grey-scale images of the carotid bifurcation, show the column of flowing blood by colour-coded ultrasound and estimate blood flow velocities using the Doppler principle. Examination is entirely noninvasive but is highly operator dependent. The technique shows high accuracy in the detection of moderate and severe stenosis of the carotid arteries but the differentiation of very high grade stenosis from complete occlusion can be difficult. Carotid ultrasound can demonstrate carotid artery dissection - either directly by B-mode imaging of a dissection flap, or by demonstrating the typical to and fro signal of blood oscillating within the residual internal carotid artery stump.11

It is our policy to confirm ultrasound detected abnormalities using MRI and magnetic resonance angiography (MRA). MRA shows high sensitivity and specificity for the detection of carotid stenosis when compared against intra-arterial angiography.12 Standard T1 weighted axial MRI sequences through the neck demonstrate the presence of fresh thrombus (appearing as high intensity signal) within the arterial wall following dissection of the carotid or vertebral arteries (figure 1).11 In carotid dissection, MRA demonstrates the typical tapering internal carotid artery stump (figure 2) and may show a 'string sign' indicating a narrow residual lumen. This sign is also seen with very high grade stenosis or pseudo-occlusion. The physiological asymmetry of the vertebral arteries makes noninvasive diagnosis of vertebral dissection more difficult on ultrasound and MRA but the demonstration of high signal within the vessel wall on axial MRI sequences enables confident diagnosis.

Intra-arterial digital subtraction angiography carries a 1% overall risk of serious morbidity or mortality. We reserve it for those patients in whom the above techniques have failed to demonstrate adequately the suspected pathology, and for patients too claustrophobic to tolerate MR scanning. We only perform angiography if it has a high chance of yielding useful diagnostic data with implications for acute management, secondary prevention, or sometimes prognosis. In general, each angiographic study should be limited to the culprit arterial territory in order to minimise potential complications.13 Angiography demonstrates occlusive disease due to atheroma or Takayasu's arteritis, carotid dissection (string-sign), fibromuscular dysplasia (beading of the vessel wall), intracranial vasculitis (segmental narrowing and tapering of medium and small arteries) in 50% of cases, and the 'puff of smoke' characteristic of moyamoya disease.

THE INTRACRANIAL VESSELS
The intracranial vessels can be imaged noninvasively using MRA. Limitations in spatial resolution dictate that smaller vessels are less well imaged. MRA can demonstrate intracranial arterial occlusion, stenosis, arteriovenous malformations and aneurysms of over 5 mm diameter.14 Diseases of smaller vessels such as arteritis can be detected, although intra-arterial angiography remains the modality of choice.15

Transcranial Doppler ultrasound is able to provide a measure of intracranial artery blood flow velocity (usually limited to the middle cerebral artery). Intracranial artery stenosis and occlusion can be detected, and the cheap and portable nature of transcranial Doppler ultrasound makes it a potentially useful bedside monitoring technique, especially for monitoring arterial recanalisation.16 Transcranial Doppler ultrasound is able to detect microembolic phenomena17 and the registration of such signals in the cerebral circulation in response to a peripheral venous injection of echo-contrast agents is an indicator of right to left shunting in patients with atrial septal defects or patent foramen ovale.18

Principal causes
DISSECTION OF THE EXTRACRANIAL ARTERIES
Dissection of the carotid or vertebral arteries is associated with disorders of connective tissue such as Ehlers-Danlos syndrome, Marfan's syndrome, pseudoxanthoma elasticum, fibromuscular dysplasia and cystic medial degeneration. There is an association with intracranial aneurysms and possibly with alpha-1-antitrypsin deficiency.19 An underlying disorder is apparent in a small minority of cases. Dissection can be either spontaneous or post-traumatic but the provoking event can be trivial, thus blurring this distinction. Reported causes include whiplash injuries, cervical manipulation by chiropractors, abrupt changes in posture, skiing, swimming and seizures.

The incidence of spontaneous carotid dissection is similar to that of subarachnoid haemorrhage—approximately 2.5–3 per 100 000 per year.20
Principal causes of ischaemic stroke in young adults

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<td>- Fabry's disease</td>
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<td>Venous infarction</td>
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Box 2

Carotid dissection accounts for 10–25% of stroke in the 15–45 year age group.\(^{21,22}\) Its recognition has important management implications. Carotid dissection presents with local and/or ischaemic symptoms. Local symptoms are neck or hemicranial pain (75% of patients), sometimes accompanied by an ipsilateral Horner's syndrome (35% of patients) due to stretching of the sympathetic fibres surrounding the internal carotid artery. Sweating of the face is usually preserved since these fibres accompany the external carotid artery. Rarely a carotid aneurysm may develop but these virtually never rupture. Lower cranial nerve palsy (XII, IX, X, in order of frequency) and rarely third nerve palsy are recognised. Isolated headache and pulsatile tinnitus are unusual.

Local symptoms are followed in 60–75% of patients by transient ischaemic attacks or completed stroke due to distal embolisation from the thrombosing false lumen.\(^{23,24}\) Dissection of the vertebral artery commonly provokes neck pain together with ischaemic symptoms referable to the posterior circulation. Unlike carotid dissections, vertebral dissections more frequently originate or extend intracranially and blood may enter the subarachnoid space causing meningism. A minority of patients (<30%) have simultaneous dissections of two or more vessels - in these patients a careful search for an underlying connective tissue disorder is necessary.

Dissection of the extracranial vessels is confirmed by duplex ultrasound, MRI, MRA or angiography.\(^{11,25,26}\) Management depends upon the presence or absence of neurological episodes and the timing of the initial event. Ischaemic events commonly occur within the first week of the dissection, but can arise up to one month afterwards.\(^{25}\) In most patients presenting acutely who have had ischaemic episodes in the carotid territory, it is our policy to anticoagulate with intravenous heparin, followed by warfarin for six months.\(^{23,25,27}\) Anticoagulation is also used in extracranial vertebral artery dissection accompanied by ischaemic events\(^{26,26}\) but we advise prior lumbar puncture to exclude subarachnoid extension. In extensive cerebral infarction the risk of haemorrhagic transformation might preclude early anticoagulation. Patients presenting late or with local symptoms alone can be managed safely with 300 mg of aspirin daily. Patients falling between these extremes must be judged in their own right and the risks and benefits of anticoagulation considered. Unfortunately, there are no good trial data to help.

Many dissected arteries recanalise within three to six months, and no surgical procedure (eg, thrombectomy, endarterectomy) is necessary. Assuming there is no underlying connective tissue disorder, the risk of recurrence is low – approximately 1% per annum.\(^{24}\)

**CARDIOGENIC EMBOLISM**

A cardiac source of emboli is found in 20–30% of young stroke patients.\(^{22,29}\) The most common cardiac lesions are prostatic heart valves, rheumatic valve disease, bacterial endocarditis, dilated cardiomyopathy, ischaemic dysskinetic segments, atrial septal aneurysm, patent foramen ovale, and mitral valve prolapse.\(^{30}\) The causative associations of the last three conditions are unclear. Both patent foramen ovale and mitral valve prolapse occur frequently in young adult populations (in up to 30% and up to 20%, respectively).\(^{31-34}\) These lesions are frequent isolated findings (40% and 10%, respectively) in series of young stroke and occur four times as commonly in such patients as in age- and sex-matched controls.\(^{30-33,35}\) Mitral valve prolapse seems likelier to be the culprit if there is significant (>2 mm) prolapse of one or both valve leaflets and if there is accompanying myxomatous degeneration.\(^{36}\) The latter predisposes to both infective endocarditis and to the formation of noninfected thrombotic vegetations. Overall, patients with mitral valve prolapse have a risk of stroke of only 1 in 6000 per year,\(^{37}\) thus it is advisable to exclude other causes of stroke first.

Atrial septal aneurysm is a lax segment of the interatrial septum whose presence frequently indicates the coexistence of a patent foramen ovale. It is unclear whether atrial septal aneurysm is an independent cause of embolic stroke or whether it indicates the presence of more extensive septal defects likely to permit right to left intratrial shunting.\(^{38,39}\) Right to left shunting is not always demonstrable in limited patent foramen ovale defects and most strokes do not originate during manoeuvres (eg, Valsalva) which increase right atrial pressures.

Spontaneous echo contrast is a frequent finding on transoephalgeal echocardiography in young stroke patients (up to 9%).\(^{40}\) It is caused by sluggish left atrial blood flow and is more common in atrial fibrillation. The significance of this finding is unclear but it may indicate flow disturbance within the left atrium sufficient to allow thrombus formation. The growing availability of transoephalgeal echocardiography and its sensitivity in picking up subtle
Case report 1: Internal carotid artery dissection

A 35-year-old woman suffered minor left-sided neck pain triggered by sudden braking of a coach. Ten days later she had an episode of left amaurosis followed several hours later by transient dysphasia, right-sided facial weakness and right hand clumsiness. She attended the local hospital and emergency CT of the brain was normal. As her symptoms had improved she was discharged.

Two weeks later the only signs were mild drooping of the right corner of the mouth and mild clumsiness of the right hand. An urgent MRI of the brain showed three discrete left fronto-parietal infarcts, one of which showed haemorrhagic transformation. MRA of the cervical carotid arteries showed occlusion of the left internal carotid artery with a small residual stump (figure 2). Axial T1-weighted images of the neck showed high signal within the wall of the left internal carotid artery with complete obliteration of the true lumen. These appearances were consistent with a left internal carotid artery dissection with cerebral embolism from the thrombosed vessel. The dissection had presumably been precipitated by relatively trivial neck trauma. Aspirin 300 mg daily was commenced and she suffered no further neurological episodes.

abnormalities whose prevalence in asymptomatic subjects is unknown, creates a dilemma in deciding whether such abnormalities are causative.

The identification of a cardiac lesion requires appropriately targeted medical or surgical management. The risk of further cardioembolic phenomena is high immediately after the first event, thus secondary prevention should be instituted with warfarin. This can be commenced immediately if the initial deficit or ischaemic lesion is small. Larger infarcts carry a greater risk of haemorrhagic transformation but after 11 days the risk is low—therefore a delay in anticoagulation of these patients is prudent.

PREMATURE Atherosclerosis

When the causes of stroke in the young are broken down by age, atherosclerosis becomes increasingly prominent from the 15–30 year age group (2%) to the 30–45 year group (30–35%). In addition to the usual risk factors (hypertension, cigarette smoking, hyperlipidaemia, diabetes mellitus), premature atherosclerosis has less widely recognised associations.

Patients with homocystinuria (an autosomal recessive inability to convert homocysteine to cystathionine and methionine) develop premature large vessel atherosclerosis. Homozygotes have Marfanoid features and diagnosis is confirmed by finding elevated homocysteine levels in blood and urine and a positive nitroprusside test. Heterozygotes and patients with milder hyperhomocysteinaemia also have a tendency to premature atherosclerosis. A methionine loading test (methionine being the immediate precursor of homocysteine) may be required to demonstrate the metabolic abnormality in some cases. Deficiency of folate acid, pyridoxine and vitamin B12 (all cofactors in homocysteine metabolism) appear to exacerbate the underlying metabolic deficit. Treatment by dietary therapy, folate or pyridoxine may reduce vascular complications.

Although atheroma commonly forms at the carotid bifurcation, it can also develop along the common carotid artery. Atheroma here is particularly common in patients who have received radiotherapy for laryngeal tumours. Cranial irradiation produces a radiation vasculopathy of the cerebral vessels. Endarterectomy of proximal cervical atheroma can be technically difficult if the common carotid origin is involved, necessitating alternative revascularisation techniques such as bypass procedures or angioplasty.

MIGRAINE

The number of strokes attributed to migraine varies from as few as 4% to as many as 20%. Before diagnosing migrainous stroke it is important to exclude other coexisting conditions. The overall lifetime prevalence of migraine is 10-16% and for the majority of patients with migraine who have a stroke, migraine is not the cause. Ischaemic stroke can precipitate migraine in the elderly and a case has been made for investigating such patients for cerebrovascular disease. There are similar reports of 'symptomatic migraine' occurring in young patients with internal carotid artery dissection and it is probable that many 'migrainous strokes' in early series were, in fact, dissections.

Stroke can only be attributed to migraine if certain conditions are fulfilled:

- the ischaemic event develops in a patient who suffers migraine with aura (classical migraine)
- the provoking attack is identical to previous attacks
- the neurological deficit is not reversible after seven days
- investigations have excluded other causes.

Stroke cannot be blamed on migraine if there is a history of migraine without aura (common migraine) only.

Case-control studies reveal an excess of classical compared to common migraine (odds ratio 1.3 to 0.8) in young stroke patients, and an increased susceptibility to stroke in young women with migraine, especially those who smoke, but not those who take oral contraceptives. The overall relative risk of stroke that migraine conveys is unknown but appears to be small.

Migrainous strokes typically involve the territory of the posterior cerebral arteries but not exclusively so (figure 3), and probably arise from prolonged arteriolar constriction. Small vessel thrombosis due to platelet activation also plays a role. Uncertainty in pathogenesis is reflected in uncertainty about optimum treatment; we ensure adequate codeine-based analgesia, give aspirin, and would add steroids and a calcium channel antagonist if initial measures prove inadequate. The risk of recurrent migrainous stroke was thought to be low but a third of patients in a recent survey had recurrent events. Whatever the future risk, future migraine prophylaxis is necessary.
Case report 2: Cardioembolic stroke with atrial septal defect and aneurysm

A 33-year-old woman suddenly developed a right homonymous hemianopia. She had no vascular risk factors. Cardiac examination revealed the visual field deficit but no other signs. Aspirin was commenced. Unenhanced CT performed within 24 hours was normal. MRI revealed a left occipital infarct. ECG and chest radiography were normal. There was no evidence of vertebral artery dissection on duplex ultrasound or MRA. Thrombophilia screen was negative.

Although her field defect gradually improved she continued to have discrete episodes of more pronounced right homonymous hemianopia with sensory disturbance of the right-sided limbs. She was anticoagulated with warfarin. Transthoracic echocardiography revealed an atrial septal aneurysm. Transoesophageal echocardiography confirmed an extensive atrial septal aneurysm and an interatrial septal defect (atrial septal defect or patent foramen ovale). Given the risk of lifelong anticoagulation with warfarin, the cardiac abnormalities were surgically repaired. The redundant atrial septum was excised and a large secundum atrial septal defect was closed. Warfarin was discontinued. She had no further cerebral ischaemic episodes.

DRUGS

Recreational drug abuse was not listed as a cause of young stroke in a British review in 1979 and it was not identified as a precipitant of stroke in 60 young patients in Oxfordshire between 1978 and 1982. However it accounted for 10% of young stroke in Baltimore, US, from 1988 to 1989 and appears as a leading cause of stroke in most contemporary series. Drug abuse increases the relative risk of stroke six-fold across all age groups and eleven-fold in people under 35 years.

Cerebral infarction occurs with heroin, amphetamine and cocaine abuse, and abuse of over-the-counter sympathomimetics. The mechanism of ischaemia is predominantly one of obliterative arteritis due to immune complex deposition following prolonged challenges with foreign antigens. Incomplete solution of crushed oral preparations also generates an arteritic response in the brain and the lungs. The subsequent development of pulmonary arteriovenous fistulae promotes transpulmonary passage of even larger sized particles into the systemic circulation. Stroke onset is usually between six and 24 hours following drug administration. Necrotic cerebral infarction also arises indirectly in drug abusers through concomitant bacterial endocarditis or fungal infections with Nocardia or Aspergillus.

THROMBOPHILIAS

Although thrombosis is usually triggered by an abnormal endovascular surface, primary abnormalities of the coagulation and fibrinolytic systems are also associated with venous and occasionally arterial thrombosis. The inherited thrombophilias (deficiencies of proteins C, S and antithrombin) are relatively common (1:200 to 1:2000) in the heterozygous form but symptomatic deficiencies are less widespread (perhaps 1:36 000). Symptomatic episodes are usually venous thromboses of the calf or unusual sites such as the cerebral dural sinuses. Arterial thrombosis is less common and there may be over-representation of hereditary thrombophilias in young stroke series due to assay of these factors soon after the acute event. Protein S levels in particular fall with acute complement activation thus persisting deficiency over the ensuing six months should be demonstrated for definitive diagnosis. Other prothrombotic states (polycythaemia, myeloproliferative disorders, thrombotic thrombocytopaenic purpura, hyperfibrinogenemia, etc) also require exclusion.

ANTIPHOSPHOLIPID ANTIBODIES AND LUPUS ANTICOAGULANT

Lupus anticoagulant and anticardiolipin antibodies belong to a class of antibodies with activity against protein-phospholipid complexes. They are detected in 50% of patients with lupus and occur in other autoimmune disorders such as rheumatoid arthritis and giant cell arteritis. They are found in patients without evidence of collagen vascular diseases but with a history of recurrent arterial and venous thromboses, spontaneous abortions and livedo reticularis (the primary antiphospholipid [Sneddon’s] syndrome). They are an independent risk factor for cerebral infarction and are an important cause of a false positive VDRL.

The mechanism of infarction in antiphospholipid syndromes and lupus was considered to be vasculitic. Current opinion, however, favours a primary thrombotic tendency within small arterioles. Spontaneous recanalisation of these thromboses gives rise to the multiple narrowings seen on angiography previously considered inflammatory. Additional cardioembolism (Liebmann Sachs endocarditis - which may only be diagnosed on transoesophageal echocardiography) and thrombotic thrombocytopaenia are common causes of infarction in systemic lupus erythematosus. Treatment requires antiplatelet agents and/or anticoagulation according to the clinical picture. Immunosuppression using cyclophosphamide has been advocated for patients who fail to respond. Steroids used alone seem ineffective.

SYSTEMIC AND ISOLATED VASCULITIS OF THE CENTRAL NERVOUS SYSTEM

Vasculitis can affect the central nervous system (CNS) as an isolated phenomenon (isolated angiitis of the CNS) or as part of a systemic necrotising vasculitis (eg, Wegener’s, polyarteritis nodosa). A CNS vasculitis arising from the latter may occur in a previously diagnosed patient. If not, the associated features and peripheral haematological and immunological markers aid diagnosis. In addition, the predilection of polyarteritis for peripheral nerves (mononeuritis multiplex) allows readily available tissue for histology (sural nerve biopsy).

Other categories of arteritis include autoimmune disorders (rheumatoid arthritis, systemic lupus erythematosus, scleroderma, etc), infections (herpes zoster, cytomegalovirus, human immunodeficiency virus (HIV), tuberculosis,
Case report 3: Migrainous stroke

For the previous 10 years a 35-year-old man had attacks of migraine twice annually. He developed one of his typical migraine attacks. Then his vision became blurred and over the next 12 hours he started bumping into things on his right hand side. The visual disturbance fluctuated in severity and he became progressively dysphasic and forgetful. His migrainous headache persisted.

Cardiovascular examination was normal. He was dysphasic and had a right homonymous hemianopia. Over the next week he developed fluctuating right pyramidal signs. MRI of the brain revealed infarction in the left occipital lobe, and repeat MRI after seven days showed extension of the infarct into the left temporal lobe (figure 3). Cerebral angiography showed pruning of the smaller vessels in the left parieto-occipital region but no definite vasospasm. Cardiac and haematological investigations were normal.

Aspirin and verapamil were commenced. When the pyramidal signs developed, steroid therapy (prednisolone 40 mg daily) was started. He gradually improved and the prednisolone was tapered off. Verapamil was replaced by propranolol for long-term prophylaxis.

Figure 3 Axial T1-weighted MRI of the brain showing extensive infarction in the left temporal lobe and along the inferior surface of the left occipital lobe (arrows). The cause of this atypical distribution of cerebral infarction was severe migraine (Case 3)

syphilis), neoplasia (lymphomas, hairy cell leukaemia, neoplastic angioendotheliomas), and drugs (cocaïne, amphetamines, sympathomimetics). All arteritides due to infections are more severe in patients with concomitant HIV infection who also develop mycotic aneurysms and thrombotic noninfective endocarditis. Stroke accounts for 3% of the CNS features of acquired immunodeficiency syndrome. Meningovascular syphilis is characterised by headache, seizures and hemiplegia in the presence of positive serology. Zoster arteritis can result in a contralateral hemiparesis arising six to eight weeks after ophthalmic zoster.

Isolated angitis of the CNS presents a diagnostic challenge. This rare disorder affects all age groups of both sexes. It presents acutely or subacutely as a focal or diffuse encephalopathy. Confusion (63%), headache (59%), hemiparesis (44%) and drowsiness (32%) are the most common features but transient ischaemic attacks, acute stroke and multi-infarct states are also described. There are usually no symptoms outside the CNS other than vague malaise, weight loss or occasional fever (10-15%). Left untreated, the condition carries a poor prognosis (90% mortality within a year of diagnosis).

Combination treatment with prednisolone and cyclophosphamide can be curative, especially in patients presenting with focal features. Presentation with a severe diffuse encephalopathy indicates a poor outcome.

There are usually no peripheral markers of isolated angitis (normal erythrocyte sedimentation rate, negative autoantibody screens) but a mono-nuclear pleocytosis and mildly elevated protein may be found in the cerebrospinal fluid. The electroencephalogram is nonspecifically slow, and MRI shows multiple small hyperintensities on T2 images whose appearance is also nonspecific. Cerebral angiography shows multiple segmental narrowing of medium and small vessels in approximately 50%. But similar appearances can arise from carotid stenosis meningitis, radiation vasculopathy, or fibromuscular dysplasia. The diagnosis is secured by leptomeningeal and wedge cerebral biopsy (nondominant frontal lobe or temporal tip) which carries a risk of 0.5-2% of focal deficit or death. The benefit of histological diagnosis, given a potentially treatable condition, outweighs the risk of biopsy in a suspected case and there is no place for empirical treatment given that steroids and cyclophosphamide administration might last for 12 months or more. The decision to biopsy must be made on an individual basis in consultation with the patient's family. We would tend to proceed to biopsy if the clinical state is deteriorating and the angiogram is negative or unhelpful.

PREGNANCY

Historically, pregnancy has been considered a risk factor for ischaemic stroke. Early studies suggested an incidence of one stroke per 2000 pregnancies. Difficulties with population bias and incorrect diagnosis prior to the CT era account for much of the over-estimation. Recent studies have suggested an incidence of 1 in 10 000-20 000 pregnancies and cohort studies suggest little difference in the stroke rate between pregnant women and nonpregnant women of childbearing age. There appears to be a definite association of cerebral venous thrombosis with pregnancy and the puerperium.

Pregnancy-related ischaemic strokes usually occur during the third trimester and puerperium. In addition to the usual risk factors, pregnancy brings its own specific conditions. Eclampsia is associated with cerebral haemorrhage but surged in blood pressure against a background of hypertension might also precipitate infarction. Other causes include chorioiocarcoma, paradoxical embolism, hypotension and postpartum cardiomyopathy. The British Neurological Surveillance Unit aims to answer some of the unresolved dilemmas surrounding pregnancy-related stroke.

GENETIC DISORDERS

It is clear that genetic factors influence disorders such as diabetes mellitus, hypertension and hyperlipidaemias, all of which predispose to cerebral infarction. Recently a single gene disorder directly responsible for ischaemic stroke has been described. The gene for cerebral autosomal dominant arteriopathy with subcortical infarcts and leucoencephalopathy (CADASIL) localises to chromosome 19, which also hosts the gene for familial hemiplegic migraine. CADASIL presents from the third decade with discrete stroke or transient ischaemic attack then develops a phase of progressive neurological disability, pseudobulbar palsy and dementia. MRI reveals multiple small infarcts in the deep white matter which have been confirmed at autopsy. Although only a few families have so far been described, the identification of CADASIL illustrates the existence of primary genetic disorders directly responsible for cerebral vascular disease in the young.

Stroke is seen as a manifestation of other inherited disorders. Mitochondrial
Ischaemic stroke in the young

myopathy, encephalopathy, lactic acidosis and stroke-like episodes (MELAS) is a mitochondrial cytopathy diagnosed by increased lactate levels in blood and cerebrospinal fluid, muscle biopsy (for ragged red fibres) and mitochondrial DNA analysis. As with other mitochondrial cytopathies, father to son transmission is impossible since all mitochondria are derived from the ovum.

Ehlers Danlos syndrome type IV (autosomal dominant), Marfan’s syndrome (autosomal dominant), osteogenesis imperfecta (autosomal dominant [types I, II, IV] and recessive [type III]), and pseudoxanthoma elasticum (autosomal dominant and recessive forms described) are all causes of cervical artery dissection. The latter is also associated with intracranial arterial occlusive disease, as is neurofibromatosis type 1.22 Fabry’s disease (X-linked recessive inheritance) is a lysosomal storage disorder characterised by angiokeratoma corporis diffusum - dark red papules found on the lower trunk, perineum and thighs. Deficiency of α-galactosidase leads to accumulation of trihexosyl ceramide in blood vessels. Patients die in the fourth and fifth decades from cerebrovascular, cardiovascular or renal disease.

Prognosis

Prognosis in young stroke reflects the underlying cause and the extent of initial neurological damage. The greater collateral reserve in the young adult brain limits the initial size of infarction and there is greater scope for functional recovery than in the elderly.

Initial mortality in young ischaemic stroke is approximately 2 – 7%,42,72 and occurs predominantly in those with large vessel occlusive disease. Overall, the risk of recurrent stroke is 1 – 3% per annum.73,74 Premature atherosclerosis is associated with a high chance of future morbidity (myocardial infarction, peripheral vascular disease, sudden death) and patients in this category have twice the risk of future vascular events than other young stroke survivors. This necessitates an aggressive approach to risk factor modification. In contrast, patients with stroke in whom full investigation fails to elucidate a cause have a low risk of recurrence (0.5 – 1% per annum).22,73,74

Amongst survivors of young stroke, 75% have little or no handicap, up to 55% of patients suffer significant depression, 50% report significantly impaired quality of life and only 40% return to work.75,76 Although 90% of those who receive physiotherapy, occupational therapy and speech therapy feel they benefit, only 40% of young stroke patients receive these measures.77 A recent UK survey found young stroke patients to be receiving as little as one hour of speech and physiotherapy weekly whilst in hospital.77 The role of ancillary therapists to offer psychological counselling, aid in the home and in the workplace, must not be underestimated.

Conclusion

Our perception of the principal causes of ischaemic stroke has changed considerably over the last 15 years. New imaging modalities allow us to appreciate conditions such as arterial dissection which were previously greatly under-recognised. Advances in haematology and immunology promote greater awareness of the inherited and acquired thrombophilias. Recent advances in genetics have enabled the identification of a single gene disorder which manifests as stroke and further genetic defects are likely to be identified. Social changes such as the increasingly widespread use of recreational drugs are reflected in the greater representation of drug abuse as a cause of stroke in the young.

It is no longer sufficient merely to demonstrate cerebral infarction as the cause of the neurological deficit in the young stroke patient. The chances of finding an underlying cause are rising as technology and knowledge progress. In addition, the opportunities for therapeutic intervention are many and should not be missed, given the life expectancy of this age group.

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