Neurological emergencies: acute stroke

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Abstract

Stroke causes a vast amount of death and disability throughout the world, yet for many healthcare professionals it remains an area of therapeutic nihilism, and thus uninteresting. This negative perception is shared by the general public, who often have a poor understanding of the early symptoms and significance of a stroke. Yet within the past few years there have been many important developments in the approach to caring for stroke patients, for both the acute management and secondary prevention. After the completion of numerous clinical trials, there is now robust evidence to either support or discredit various interventions. Even more exciting is the prospect of yet more data becoming available in the near future, testing a whole array of treatments, as clinical interest in stroke expands exponentially. In this review an evidence based approach to the management of acute stroke within the first few days is presented, including ischaemic and haemorrhagic events, but not subarachnoid haemorrhage. It is explained why stroke is regarded as a medical emergency, and the importance of a rational, methodic approach to the initial assessment, which is the key to accurate diagnosis and subsequent management, is emphasised. The potential early problems associated with stroke are identified and specific interventions for different stroke types are discussed. The review ends with a brief discussion of the implications that the evolving treatments have for the organisation of modern stroke services.

Keywords: acute stroke

In the western world, stroke is the third commonest cause of death (after heart disease and all cancers), is probably the commonest cause of severe disability, and accounts for a large proportion of healthcare resources. Its impact on individual patients, their families, and society as a whole is immense. About 200 people per 100 000 population will have a first ever stroke every year. Their mean age is about 72 years, and men and women are affected in roughly equal numbers. Despite the uncertainty over whether stroke incidence is rising, falling, or remaining static, the absolute number of patients is likely to increase, as incidence increases with age and most populations are aging.

Stroke has in the past held a low priority for many professional groups; in 1988 the King’s Fund Forum concluded that hospital, primary care and community services for stroke in the United Kingdom were “haphazard, fragmented, and poorly tailored to the patients’ needs”. However, increasing awareness of the impact of stroke has led to it being identified as a priority for improving services and research. Numerous well conducted randomised trials and systematic reviews of medical and surgical treatments for acute stroke, as well as primary and secondary prevention strategies, mean that we are now considerably better equipped to know which treatments work, and which should be abandoned, and what sort of services we should be providing for our patients.

In this review, we will describe our approach to the management of acute stroke, focusing mainly on the first few days. Although the World Health Organisation (WHO) definition of stroke (see below) includes subarachnoid haemorrhage, we will not consider this distinct syndrome further, as it is covered in detail elsewhere in this series. Our approach is based on our interpretation of the available evidence, with particular emphasis on randomised controlled trials and systematic reviews, as we think that these provide the most reliable data on the risks and benefits of treatments. However, where such evidence is either absent or insufficient (and despite the many welcome advances, there is still much we do not know for sure), we will describe what we do in routine practice. We accept that other interpretations of the evidence are possible and are likely to be influenced by the context in which one works. For example, in the United Kingdom, unlike some other countries, our patients tend not to demand “treatment” (whether of proved benefit or not), “fee for service” is rare, and the resources available for health care are restricted so that only cost effective treatments will be advocated for widespread use. Lastly, although rehabilitation and secondary prevention are not the focus of this review, it is important to emphasise that for many patients, their management after the
acute stage currently has the greater impact on their lives; with the advent of new acute strategies, this balance may change.

Is acute stroke an emergency?
Medical emergencies can be defined by certain criteria including rapidity of onset, poor prognosis, and requirement for prompt intervention. Although stroke has traditionally been treated as less of an emergency than, for instance, acute myocardial infarction or meningitis, we illustrate that this conservative approach is no longer tenable, and that stroke should now be regarded as a medical emergency.

STROKE COMES ON RAPIDLY
The WHO has defined stroke as a clinical syndrome characterised by rapidly developing symptoms and/or signs of focal, and at times global (for patients in coma), loss of cerebral function, with symptoms lasting more than 24 hours or leading to death, with no apparent cause other than that of vascular origin.1

STROKE HAS A POOR PROGNOSIS
The outcome after stroke is crucially dependent on the extent and site of the brain damage, as well as the patient’s age and prestroke health status.8 The case fatality rates after a first ever stroke (all types combined) are 12% at 7 days, 19% at 30 days, and 31% at 1 year; haemorrhagic stroke carries a higher risk of death than ischaemic stroke.9 Deaths occurring within the first week after stroke are mostly due to the direct effects of cerebral damage; later on, the complications of immobility (for example, bronchopneumonia, venous thromboembolism) and cardiac events become increasingly common.10 About 20% of those with first ever stroke will be dependent on another person for everyday activities (for example, washing, dressing, mobility) at 12 months, and 50% will be independent.11 The risk of a recurrent stroke in survivors is about 10% to 16% within the first year, thereafter falling to about 5% per year.12 The relative risk of death in stroke survivors is about twice the risk of people in the general population, and this risk persists for several years; many of these deaths are due to other vascular problems (for example, ischaemic heart disease or peripheral arterial disease). This emphasises the importance of targeting these areas as well as stroke when we consider secondary prevention strategies.

STROKE PATIENTS MAY REQUIRE IMMEDIATE TREATMENT
There are several reasons why many patients require urgent inpatient care after an acute stroke. Firstly, stroke may lead to various potentially life threatening complications such as airway obstruction and respiratory failure, swallowing problems with the risk of aspiration, dehydration and malnutrition, venous thromboembolic complications, seizures, and infections.14–17 These may arise within hours of stroke onset and require early assessment and intervention so that they can be anticipated, prevented, and treated. Furthermore, although stroke has represented an area of therapeutic nihilism for many years, various acute and potentially effective treatments (medical and surgical) are now becoming available.

Assessment
Early assessment allows the formulation of an accurate and early diagnosis (as stroke is primarily a clinical diagnosis, the sooner a physician can elicit a history, the more likely it is to be reliable), the organisation of relevant and cost effective investigations, and the initiation of appropriate secondary prevention (which is likely to be most effective early on, when the risk of recurrence is highest). However, as with any medical emergency, the first priority in assessing a patient after a suspected stroke is to identify and treat any immediately life threatening complications. For stroke, this will usually be an obstructed airway, respiratory failure in a comatose patient, or an acute circulatory disturbance. Once the patient is stable, we apply a systematic, staged approach to making the diagnosis and formulating a management plan. This initial assessment should consider the following questions:

(1) IS THIS A VASCULAR EVENT?
The diagnosis depends crucially on an accurate history, taken from the patient or carer. We ask ourselves the following questions to help decide whether it was a vascular event.
• Are the neurological symptoms focal rather than non-focal?
• Are the focal neurological symptoms negative (loss of function) rather than positive (for example pins and needles rather than numbness)?
• Was the onset of the focal symptoms sudden?
• Were the focal symptoms maximal at onset (coming on over minutes to hours) rather than progressive (evolving over hours to days)?

If the answer to all these questions is yes, then a vascular cause (either cerebral ischaemia or haemorrhage) is very likely. Of course presentations vary. Occasional patients have symptoms or signs which are not easily localised (for example, memory impairment, confusion, or a reduced conscious level), symptoms may be positive (for example, movement disorders), and many patients describe symptoms evolving over hours or even days. These exceptions simply make the clinical diagnosis less certain and should lead to early investigation to exclude alternative diagnoses which require different urgent treatment (for example, hypoglycaemia, non-convulsive seizures, cerebral infection, or subdural haematoma).

In addition, we consider the context in which the event has occurred. Strokes are uncommon in the young, and as about 80% of stroke patients have at least one vascular risk factor at presentation,18 the absence of risk factors should lead one to be slightly more sceptical about a diagnosis of stroke. Accurate diagnosis in the hyperacute phase (less than 6 hours from onset) is often difficult because symptoms and signs may be changing rapidly. The introduction of acute therapies which need to be...
administered within this time window suggests that early and accurate diagnoses will become increasingly important.

(2) WHICH PART OF THE BRAIN IS AFFECTED?
In reaching a diagnosis of stroke one inevitably makes some assessment of where in the brain the lesion might be. However, it may be useful to further subclassify the stroke as this may give clues to the likely underlying cause, allow more cost effective investigations, and help in predicting both the risk of recurrence and functional outcome. Although there are many subclassification systems available, we use the system developed from the Oxfordshire Community Stroke Project (table 1), which works well at the bedside.

(3) IS IT A HAEMORRHAGIC OR ISCHAEMIC STROKE?
Distinguishing between a haemorrhagic and ischaemic stroke is important in terms of acute management, prognosis, and secondary prevention. In white people, about 80% of first ever strokes are ischaemic.

Although various scoring systems have been devised to help differentiate between infarction and haemorrhage, none provide sufficient accuracy to guide treatment. The only reliable method of differentiating is early brain imaging. In many countries, this is best performed by CT. Lumbar puncture may be useful in confirming subarachnoid haemorrhage if the brain imaging is equivocal, but it has no place in differentiating ischaemic and haemorrhagic stroke.

Computed tomography
Intracerebral blood immediately appears as an area of high density on CT, but thereafter decreases so that haemorrhagic lesions will eventually appear either isodense or hypodense, and thus be indistinguishable from an infarct. Smaller haemorrhages may become isodense within days although usually this process takes weeks. Computed tomography in the hyperacute stage of an ischaemic stroke is often normal although there may be subtle changes which are easily overlooked by the inexperienced observer. Infarcts are most easily seen on CT after a few days or in the chronic phase, when they may become markedly hypodense and well defined, although up to 50% of patients with a clinically definite stroke never have an appropriate lesion identified on CT. Although early CT will reliably identify intracerebral haemorrhage, the distinction between a primary intracerebral haemorrhage (ICH) and haemorrhagic transformation of an infarct (HTI) is unreliable and difficult. The frequency and clinical relevance of HTI is uncertain, and radiologically ranges from small petechial haemorrhages to frank haematoma, which may or may not be accompanied by clinical deterioration. An HTI can occur very early, and the only definitive way of diagnosing HTI is to have an earlier scan excluding haemorrhage, so we recommend scanning as early as possible, ideally at the time of initial assessment.

Magnetic resonance imaging
Magnetic resonance imaging is probably more sensitive than CT for detecting stroke, particularly lacunar strokes and those occurring in the posterior fossa. However, even MRI can be normal in clinically definite stroke.

Certain MRI techniques, such as diffusion weighted imaging, are very sensitive at highlighting the “culprit” lesion, which may be useful when several areas of abnormality are shown. The differentiation between an ischaemic and haemorrhagic stroke on MRI in the first few days is less easy for the non-expert than with CT but MRI can help diagnose intracerebral haemorrhage months or even years after the event when CT shows only a hypodense area indistinguishable from an infarct. However, physicians in many countries do not have urgent access to MRI, and it is currently a difficult technique to use safely and satisfactorily in many acutely ill patients; consequently, CT is likely to remain the principal imaging technique for stroke patients for the foreseeable future. Where available, MRI, including the use of specific sequences such as diffusion weighted imaging and MR angiography, may add significantly to the understanding of stroke mechanisms.

(4) WHAT CAUSED THIS STROKE?
The list of potential causes is long, and obviously differs for ischaemic and haemor-

Table 1 The Oxfordshire Community stroke subclassification system

<table>
<thead>
<tr>
<th>Total anterior circulation syndrome (TACS)</th>
<th>implies a large cortical stroke in middle cerebral, or middle and anterior cerebral artery territories</th>
</tr>
</thead>
<tbody>
<tr>
<td>A combination of:</td>
<td></td>
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<tr>
<td>• New higher cerebral dysfunction (e.g. dysphasia, dyscalculia, visuospatial disorder) AND</td>
<td></td>
</tr>
<tr>
<td>• Homonymous visual field defect AND</td>
<td></td>
</tr>
<tr>
<td>• An ipsilateral motor and/or sensory deficit involving at least two out of three areas of the face, arm or leg</td>
<td></td>
</tr>
<tr>
<td>Partial anterior circulation syndrome (PACS)</td>
<td>implies cortical stroke in middle or anterior cerebral artery territory</td>
</tr>
<tr>
<td>Patients with two out of the three components of the TACS OR new higher cerebral dysfunction alone OR a motor/sensory deficit more restricted than those classified as a LACS (eg isolated hand involvement).</td>
<td></td>
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</tbody>
</table>

Lacunar syndrome (LACS): implies a subcortical stroke due to small vessel disease

<table>
<thead>
<tr>
<th>Posterior circulation syndrome (POCS)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Ipsilateral cranial nerve palsy with contralateral motor and/or sensory deficit</td>
<td></td>
</tr>
<tr>
<td>• Bilateral motor and/or sensory deficit</td>
<td></td>
</tr>
<tr>
<td>• Disorder of conjugate eye movement</td>
<td></td>
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<tr>
<td>• Cerebellar dysfunction without ipsilateral log-tract involvement</td>
<td></td>
</tr>
<tr>
<td>• Isolated homonymous visual field defect</td>
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</tbody>
</table>

**nb:** evidence of higher cortical involvement or disturbance of consciousness excludes a lacunar syndrome
rhagic stroke. In individual patients, even after extensive investigation it may be difficult to establish the cause: many will have competing causes (for example, AF and carotid disease). Thus in practice, the precise cause of stroke is often uncertain. Accepting this, we estimate that about 50% of ischaemic strokes are due to atherothromboembolism, 25% due to intracranial small vessel disease, and 20% due to cardiac embolism, with only 5% due to rarer causes. Most haemorrhagic strokes are thought to be due to small vessel disease (often associated with hypertension), although amyloid angiopathy commonly underlies lobar haemorrhages; vascular abnormalities such as aneurysms and arteriovenous malformations may also underlie haemorrhage, and the risk of haemorrhage with anticoagulant drugs increases with the international normalised ratio. The history and examination may provide important aetiological clues (for example, the use of oral anticoagulants, the presence of an irregular pulse, or heart murmur). Unusual causes are considerably more likely in younger patients (for example, evidence of drug misuse, or recent cervical trauma precipitating arterial dissection). Our approach to investigation aims to be reasonably cost effective. We perform some simple investigations (full blood count, erythrocyte sedimentation rate, plasma glucose, urea and electrolytes, random plasma cholesterol, urinalysis, 12-lead ECG, and brain CT) in all patients in whom we are considering active management, even if our clinical assessment strongly suggests a common cause. These tests may identify important modifiable risk factors as well as highlighting the possibility of a rarer, unexpected cause of stroke (for example, infective endocarditis, giant cell arteritis, or thrombocythaemia) which may coexist with either cardiac or degenerative vascular disease.

We reserve other more specialised tests for patients in whom the cause of stroke is not clear (for example, young patients (less than 50 years), or those without risk factors), for those with clinical features of a rare cause, or where simple investigations show an abnormality (table 2).

(5) WHAT ARE THIS PARTICULAR PATIENT’S PROBLEMS?

A full assessment by the various members of the multidisciplinary team should be able to identify existing problems and anticipate future ones so that a problem and goal orientated management plan can be constructed. As well as assessing individual impairments and disabilities which may lead to specific interventions (for example, positioning and physiotherapy for hemiparesis), it is important not to ignore but to treat the less specific but unpleasant symptoms such as headache, vomiting, hic-cups, vertigo, constipation, and the aches and pains which so often accompany prolonged immobility. Here, we will briefly discuss some of the most common early problems which account for significant mortality and morbidity, and which are most relevant to the physician. The list is not exhaustive and does not cover the more chronic and debilitating conditions which may affect any patient with stroke.

Table 2  Second line investigations in selected stroke patients

<table>
<thead>
<tr>
<th>Investigation</th>
<th>Indications</th>
<th>Disorders suggested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver function tests</td>
<td>Fever, malaise, raised erythrocyte, ESR, malignancy</td>
<td>Giant cell arteritis, infective and non-bacterial thrombotic endocarditis</td>
</tr>
<tr>
<td>Calcium</td>
<td>Hypercalcaemia may rarely cause recurrent focal symptoms</td>
<td>Hypercalcaemia</td>
</tr>
<tr>
<td>Activated partial thromboplastin time, dilute Russell’s antigen, antinuclear and other antibodies</td>
<td>Young patient, previous or family history of venous thrombosis, recurrent miscarriage, thrombocytopenia, cardiac valve vegetations, livedo reticularis, raised ESR, malaise, positive VDRL</td>
<td>Antiphospholipid antibody syndrome, systemic vasculitis, systemic lupus erythematosus</td>
</tr>
<tr>
<td>Protein C and S, antithrombin III, activated protein C resistance, thrombin time</td>
<td>Previous or family history of thrombosis (usually venous) of young onset</td>
<td>Deficiency states</td>
</tr>
<tr>
<td>Serum proteins and electrophoresis, plasma viscosity</td>
<td>Raised ESR</td>
<td>Paraproteinemia, nephrotic syndrome, cardiac myxoma</td>
</tr>
<tr>
<td>Haemoglobin electrophoresis</td>
<td>AfroCaribbean patients</td>
<td>Sickle cell trait or disease, other haemoglobinopathies</td>
</tr>
<tr>
<td>Blood cultures</td>
<td>Fever, cardiac murmur, haematuria, deranged LFTs, raised ESR, malaise</td>
<td>Infective endocarditis</td>
</tr>
<tr>
<td>VDRL, HIV serology</td>
<td>Young, unexplained or “at risk”</td>
<td>Neurophilis, AIDS</td>
</tr>
<tr>
<td>Serum homocysteine, urinary amino acids</td>
<td>Marfanoid habitus, high myopia, dislocated lenses, osteoporosis, mental retardation, young</td>
<td>Homocystinuria</td>
</tr>
<tr>
<td>Leucocyte α-galactosidase A</td>
<td>Corneal opacities, cutaneous angiokeratomas, paraesthesias and pain, renal failure</td>
<td>Fabry’s disease</td>
</tr>
<tr>
<td>Blood/CSF lactate, mitochondrial DNA analysis</td>
<td>Young, basal ganglia calcification, epilepsy, parieto-occipital ischaemia, migraine</td>
<td>MELAS/mitochondrial cytopathy</td>
</tr>
<tr>
<td>Drug screen (blood or urine)</td>
<td>“At risk” patient, no other cause</td>
<td>Drug induced stroke (amphetamine, cocaine, etc)</td>
</tr>
<tr>
<td>Chest radiography</td>
<td>Hypertension, finger clubbing, cardiac murmur or abnormal ECG, young</td>
<td>Calciumised valves, enlarged heart, pulmonary AVM</td>
</tr>
<tr>
<td>Carotid ultrasound/MR angiography</td>
<td>Carotid distribution stroke in patient suitable for surgery</td>
<td>Cervical internal carotid stenosis</td>
</tr>
<tr>
<td>Cerebral angiography (intra-arterial digital subtraction or MR)</td>
<td>Young explained stroke, especially associated with pain or trauma, suspected arteritis, AVM or aneurysm</td>
<td>Arterial dissection, vascular abnormality</td>
</tr>
<tr>
<td>Transthoracic echocardiography</td>
<td>Suspected cardioembolism</td>
<td>Cardiembolism</td>
</tr>
<tr>
<td>Transoesophageal echocardiography</td>
<td>Suspected cardioembolism when TTE negative (eg endocarditis, atrial septal aneurysm), aortic dissection or atheroma, patent foramen ovale</td>
<td>Cardiembolism, aortic dissection or atheroma, paradoxical embolism</td>
</tr>
<tr>
<td>24 hour ECG</td>
<td>Palpitations, suspicious resting ECG, clinical suspension</td>
<td>Intermittent AF, heart block</td>
</tr>
<tr>
<td>Temporal artery biopsy</td>
<td>Older (&gt;60), jaw claudication, headache, polymyalgia, malaise, anaemia, raised ESR</td>
<td>Giant cell arteritis</td>
</tr>
</tbody>
</table>

ESR=erythrocyte sedimentation rate; VDRL=venereal disease research laboratory test; AVM=arteriovenous malfunction; LFT=liver function test; TTE=transthoracic echocardiography; AF=atrial fibrillation.
not include important problems which may arise later on (for example, painful shoulders or depression).35

**Airway and breathing**

Although stroke may cause various abnormal breathing patterns35 36 (for example, periodic respiration or hyperventilation), an intermittently obstructed airway in patients with a decreased level of consciousness may mimic this and should be excluded. The presence of significant hypoxia should stimulate a search for possible causes (for example, pulmonary oedema, pulmonary embolism, or infection). It seems reasonable to attempt to correct this with supplemental oxygen, but we do not advocate routine supplemental oxygen for all patients. We increasingly use pulse oximetry in the acute phase to alert us to significant oxygen desaturation.

**Circulation**

Hypotension is relatively uncommon in stroke patients; if it does occur, it is usually secondary to coexistent heart disease (arrhythmias, heart failure or acute myocardial infarction), dehydration, or sepsis. As cerebral autoregulation is disturbed after stroke, with the result that cerebral blood flow becomes directly dependent on systemic blood pressure, urgent correction is required. By contrast, hypertension is extremely common after stroke, even in patients without pre-existing hypertension.37–38 Although some authorities recommend early pharmacological lowering of raised blood pressure, given the current absence of any convincing evidence of the effectiveness of such a policy,39–42 and the recognised potential dangers of hypotension,41 we only give hypotensive drugs early (within the first 72 hours) to patients with features of accelerated hypertension or hypertensive encephalopathy, or acute aortic dissection. We usually continue any previous antihypertensive medication a patient may have been taking, provided they are not hypotensive and can safely swallow the tablets. There is uncertainty about when hypotensive drugs should be started after the acute phase; in many cases a raised blood pressure falls spontaneously in the days after an acute stroke. We usually delay consideration of long term drug therapy for at least a week, although we acknowledge that some physicians would start therapy earlier.

**Rased intracranial pressure**

Although intracranial pressure may rise very rapidly after haemorrhagic stroke (due to the space occupying effects), it usually takes at least 48 hours, and often longer, to manifest after an ischaemic stroke (except in the unusual case of a cerebellar or brainstem infarct obliterating the CSF pathways and resulting in hydrocephalus). Although treatments such as mannitol,43 hyperventilation, and even decompressive craniectomy44 undoubtedly reduce intracranial pressure, it is unclear whether such aggressive interventions are associated with improved survival with acceptable quality of life. In selected patients who are deteriorating rapidly and who are judged to have some chance of a reasonable recovery we do consider transfer to an intensive care unit for aggressive management of raised intracranial pressure. However it is important to remember that early deteriorating conscious level is usually a very poor prognostic indicator, and such intensive treatment in an elderly patient with significant other neurological deficits is, in our view, rarely justified.

**Stroke in evolution**

After the onset of symptoms, some patients continue to deteriorate over several hours or days. This is variably referred to as progressing or evolving stroke.46 47 Such patients require prompt reassessment and investigation, as there is a wide range of potential causes of deterioration, some of which may be reversible (table 3). If we suspect that the cause is progressive thromboembolism, we might use intravenous heparin despite the lack of evidence supporting its effectiveness (see below).

**Swallowing, hydration, and nutrition**

Dysphagia48 and poor nutrition49–52 are common after stroke and may lead to further complications.53 All patients should have a bedside swallowing assessment54 55 as part of their initial assessment by a suitably trained member of the multidisciplinary team. The gag reflex is an unreliable indicator of swallowing ability and should not be used for this purpose.56 The bedside assessment should lead to a decision regarding whether the patient is safe to swallow or not, and should be written down and clearly communicated to the nursing staff. For patients with an unsafe swallow, fluids should be prescribed (either intravenous or nasogastric), and arrangements made for further assessment by a speech and language therapist. The role of early enteral tube feeding, its timing, and whether this is best delivered via a nasogastric tube or percutaneous endoscopic gastrostomy remains unclear, and is the subject of an ongoing multicentre trial.57 Percutaneous endoscopic gastrostomy tube feeding is clearly the best option where prolonged tube feeding is necessary. However in the early stages, where the advantages and disadvantages of early versus delayed tube feeding, and the optimal type of tube are unclear, we randomise our patients in this trial.

**Table 3  Causes of deterioration after stroke**

<table>
<thead>
<tr>
<th>Neurological:</th>
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<tbody>
<tr>
<td>• Progression/completion of stroke</td>
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<tr>
<td>• Extension/early recurrence</td>
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<tr>
<td>• Haemorrhagic transformation of an infarct</td>
<td></td>
</tr>
<tr>
<td>• Developing cerebral oedema*</td>
<td></td>
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<tr>
<td>• Obstructive hydrocephalus*</td>
<td></td>
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<tr>
<td>• Epileptic seizures*</td>
<td></td>
</tr>
<tr>
<td>• Incorrect diagnosis*</td>
<td></td>
</tr>
<tr>
<td>Non-neurological:</td>
<td></td>
</tr>
<tr>
<td>• Infection*</td>
<td></td>
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<tr>
<td>• Metabolic derangement*</td>
<td></td>
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<tr>
<td>• Drugs*</td>
<td></td>
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<tr>
<td>• Hypoxia*</td>
<td></td>
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<tr>
<td>• Hypercapnoea*</td>
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</table>

*Potentially reversible causes.
Glycaemic control

Hypoglycaemia, although unusual after a stroke, should always be excluded on admission, as it may mimic stroke perfectly, and delay in its correction can lead to permanent disability or even death. Hyperglycaemia is much more common and has been attributed to previously recognised or occult diabetes, or part of an acute stress response.\(^\text{57}\) Hyperglycaemia is associated with poor outcomes and although work in animal models suggests that this association may be causal, it may simply reflect the severity of the stroke or underlying vascular disease. Thus it is unclear how aggressively hyperglycaemia should be corrected and at least one randomised controlled trial is now in progress.\(^\text{60}\) Until further evidence is available our policy is to use a glucose, potassium, and insulin infusion to correct blood sugars persistently above 15 mmol/l, and to treat even at lower levels provided there are adequate facilities for close monitoring to minimise the risk of hypoglycaemia.

Pyrexia

This may be due to infection preceding the stroke (consider endocarditis and encephalitis), the stroke itself, or most commonly a complication such as a chest or urinary infection, or venous thromboembolism. Obviously the underlying cause should be sought and treated but it is probably sensible to try to reduce the temperature using simple means (for example, antipyretic drugs) in any case as this is likely to make the patient more comfortable and there is a possibility, based on animal models and the observation that raised temperatures are associated with poor outcomes in patients,\(^\text{56,62}\) that a raised temperature may exacerbate any ischaemic cerebral damage.\(^\text{65}\) There are no published randomised trials of cooling therapy yet, for patients with either a raised or normal temperature, but small open studies have started to explore the use of cooling.\(^\text{64}\)

Pressure areas

Decubitus ulcers or pressure sores are an entirely avoidable complication, assuming that they did not develop before medical help was sought. When they do occur, they are painful, slow the patient’s recovery, and may sometimes be fatal. Prevention relies on an early assessment of the patient’s risk, expert nursing care, and the judicious use of specialised cushions and mattresses.\(^\text{65}\)

Bladder management

Incontinence of urine is common in the first few days and a source of major distress for patients and their carers.\(^\text{68}\) Usually it can be attributed to several factors including impaired sphincter control, immobility, communication problems, constipation, pre-existing prostatic or gynaecological problems, inadequate nursing, infection, confusion, and impaired consciousness. Obviously the cause or causes should be identified and rectified if possible. Most patients can be managed using absorbent pads, external urinary devices, and regular toiletting regimes. If these are impractical, and transfers very difficult and the patient’s pressure areas are causing concern, we insert an indwelling catheter despite the risk of infection and trauma. Incontinence often resolves spontaneously within the first week or two, so it is wise to try removing the catheter if it seems likely that things will have improved. For patients with persisting incontinence further investigation with bladder ultrasound or post-micturition catheterisation may be useful to assess bladder contractility and outflow. Urinary retention, particularly in men, is common and easily missed in patients with communication problems.

Venous thromboembolism prophylaxis

Studies using radiolabelled fibrinogen leg scanning suggest that deep venous thrombosis (DVT) occurs in over 50% of patients with hemiplegia.\(^\text{57}\) However, clinically apparent DVT probably occurs in fewer than 5%.\(^\text{14}\) Similarly, although postmortem series have identified pulmonary embolism in a large proportion,\(^\text{69}\) clinically evident pulmonary embolism occurs in less than 2%,\(^\text{14,17,18}\) although some pulmonary embolism may be unrecognised. The impact of venous thromboembolism after stroke is therefore unclear.

There are two strategies for prevention of venous thromboembolism; physical interventions (for example, early mobilisation and compression stockings), and antithrombotic drug therapy. The evidence to support the use of compression stockings comes from randomised control trials in the perioperative period which may not be generalisable to stroke because in stroke the stockings are applied after the onset of paralysis, and immobilisation is often prolonged. Also compression stockings, apart from being uncomfortable and time consuming to apply, can occasionally cause gangrene in patients with poor peripheral circulation. It therefore seems reasonable to recommend early mobilisation wherever possible and compression stockings (usually full length) for patients at high risk of DVT (those who are immobilised, or who have a history of DVT). However given the difficulties and risk we think that further trials to evaluate their effectiveness in stroke patients are justified.

There is reasonable evidence that aspirin reduces the risk of DVT in several clinical situations\(^\text{1}\) and it also has a small but beneficial effect on the long term outcome of patients with ischaemic stroke (see later), so we use this routinely. Although low dose subcutaneous heparin significantly reduces the risk of DVT and pulmonary embolism, this effect is offset by the complications of haemorrhagic transformation and extracranial bleeding, such that at 6 months the average patient with ischaemic stroke has no greater chance of surviving free of dependency if treated with heparin.\(^\text{70}\) We occasionally use heparin (standard unfractionated, at a dose of 5000 units twice daily subcutaneously) in patients we judge to be at particularly high risk of venous thromboembolism (for example, those with a history of previous
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DVT/pulmonary embolism and low risk of HTI (for example, lacunar infarction).

Epileptic seizures
Early seizures (within 2 weeks of stroke) occur in about 5% of patients. They are more common in haemorrhagic stroke and large infarcts involving the cerebral cortex. Seizures should prompt a review of the diagnosis of stroke (could the focal symptoms be secondary to postictal paralysis or encephalitis?), and a search for precipitating factors (for example, alcohol withdrawal, drugs, metabolic disturbance, or infection). After treating the seizures, we would then reappraise the severity of stroke, as this is notoriously difficult in the presence of seizures. We have occasionally misdiagnosed stroke in patients with non-convulsive seizures, which requires an EEG for definitive diagnosis. The treatment of poststroke seizures is no different from other forms of secondary epilepsy.

Specific treatments for acute ischaemic stroke
In the United Kingdom few treatments aimed specifically at the ischaemic brain lesion are routinely used. However, many treatments are used routinely in other countries, and evidence is accruing that certain treatments may improve outcome in selected patients. We therefore consider some of these further and review the available evidence to support their use. Before doing so, it may be helpful to consider briefly the main pathophysiological features of an ischaemic stroke; for a more detailed review, we refer readers elsewhere.

PATHOPHYSIOLOGY OF ISCHAEMIC STROKE
Ischaemic stroke usually occurs due to occlusion of a cerebral artery, or less often a reduction in perfusion distal to a severe stenosis. As cerebral blood flow falls, neuronal function is affected in two stages. Initially, as blood flow falls below a critical threshold of about 20 ml blood/100 g brain/min (normal being over 50 ml/100 g/min), loss of neuronal electrical function occurs. Crucially, this is a potentially reversible stage. Irreversible damage occurs within minutes as blood flow falls below a second critical threshold of 10 ml/100 g/min; below this level, aerobic mitochondrial metabolism fails, and the inefficient anaerobic metabolism of glucose takes over, rapidly leading to lactic acidosis. Consequently, the normal energy-dependent cellular ion homeostasis fails, resulting in potassium leaking out of the cell, and sodium and water entering the cell, leading to cytotoxic oedema. Calcium also enters the cell, exacerbating mitochondrial failure. This loss of cellular ion homeostasis leads to neuronal death.

The identification of these two stages of neuronal failure has led to the concept of the ischaemic penumbra—that is, an area of brain which has reached the reversible stage of electrical failure, but has not yet passed onto the second irreversible stage of cellular homeostatic failure. In theory therefore, this tissue could be “rescued”, either by early reperfusion (using agents to dissolve the acute thrombotic lesion and restore normal blood flow), or by administering agents which could protect these potentially viable neurons from further damage (neuroprotection); the combination of reperfusion and neuroprotection would seem a logical conclusion. Although there is evidence that the concept of the ischaemic penumbra is valid, it remains unclear how long ischaemic human brain might survive—in other words, the time window for intervention is unknown. It seems likely that the duration of any time window will vary between individual patients, and it will be increasingly important to identify the factors which influence it.

Should the mechanism of the cerebral ischaemia influence management? Some experts think that certain specific causes of ischaemia, such as basilar artery thrombosis or arterial dissection, warrant specific interventions, most commonly anticoagulation. There is no convincing evidence to support these views, and therefore we tend to treat them the same as we would any other form of ischaemic stroke. A recent randomised trial of the use of anticoagulants in cerebral venous thrombosis indicated a non-significant favourable effect.

THROMBOLYSIS
Despite having been used sporadically for over 40 years, evidence for the effectiveness of thrombolytic therapy in acute ischaemic stroke has only recently become available. A systematic review of the results of 12 of the 14 completed randomised controlled trials in the post-CT era suggests that although thrombolytic therapy (with recombinant tissue plasminogen activator, streptokinase, or urokinase) is associated with about 70 symptomatic (about 50 fatal) intracranial bleeds per 1000 patients treated, its use is associated with perhaps 65 more patients surviving free of dependency at 3 to 6 months poststroke (fig 1). Even more compelling are the updated analyses of treatment within the first 3 hours. These demonstrate less risk of early intracranial haemorrhage and early death, and greater long term net benefit (130 extra patients alive and independent per 1000 treated). How practicable the widespread use of thrombolysis will be (particularly for a condition which has not traditionally been thought of as an emergency) remains uncertain, although some units have published impressive figures. Recombinant tissue plasminogen activator (r-TPA) is now licensed in the United States, and a European licence is likely to be granted in the near future; therefore, it seems reasonable to consider using r-TPA in patients presenting within 3 hours, and who are similar to the patients included in the trials, provided there is a stroke service which can ensure its safe administration (table 4). Our view is that further trials are required to establish the balance of risks and benefits in a broader range of patients presenting at different stages, with differing severities and types of ischaemic stroke, different risk factors, and differing scan appearances. Many of the eligibility criteria currently in place are arbitrary and are not based on any reliable evidence. If a larger proportion of patients were eligible for treat-
Thrombolytic therapy should only be administered by physicians with expertise in stroke medicine, who have access to a suitable stroke service, with facilities for identifying and managing haemorrhagic complications.

Exclusion criteria: use of oral anticoagulants, or INR greater than 1.7; use of heparin in preceding 48 hours or prolonged partial thromboplastin time; platelet count less than 100 000/mm³; stroke, or serious head injury in the previous 3 months; major surgery within previous 14 days; pretreatment systolic blood pressure greater than 185 mm Hg or diastolic greater than 110 mm Hg; rapidly improving neurological condition; mild isolated neurological deficits; previous intracranial haemorrhage; blood glucose greater than 22 mmol/l (400 mg/dl) or less than 2.8 mmol/l (50 mg/dl); seizure at stroke onset; gastrointestinal or urinary bleeding within previous 21 days; or recent myocardial infarction.

Caution is advised before giving r-TPA to patients with severe stroke (NIH stroke scale score > 22) or urinary bleeding within previous 21 days; or recent myocardial infarction.

Recommended that treatment and adverse effects discussed with patient and family before treatment.

Table 4  Suggested guidelines for the use of intravenous r-TPA in ischaemic stroke [99]

<table>
<thead>
<tr>
<th>Study</th>
<th>Experiment n/N</th>
<th>Control n/N</th>
<th>Peto OR (95% CI fixed)</th>
<th>Weight %</th>
<th>Peto OR (95% CI fixed)</th>
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<td>Treatment within 3 hours</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ASK 1996</td>
<td>14/41</td>
<td>15/29</td>
<td>2.6</td>
<td>0.49</td>
<td>(0.19–1.28)</td>
</tr>
<tr>
<td>ECASS 1995</td>
<td>28/49</td>
<td>25/38</td>
<td>3.2</td>
<td>0.70</td>
<td>(0.29–1.66)</td>
</tr>
<tr>
<td>ECASS II 1998</td>
<td>39/81</td>
<td>44/77</td>
<td>6.2</td>
<td>0.70</td>
<td>(0.37–1.30)</td>
</tr>
<tr>
<td>MAST-E 1996</td>
<td>19/26</td>
<td>14/21</td>
<td>1.6</td>
<td>1.35</td>
<td>(0.39–4.68)</td>
</tr>
<tr>
<td>MAST-E 1995</td>
<td>46/79</td>
<td>69/103</td>
<td>6.6</td>
<td>0.69</td>
<td>(0.38–1.26)</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>146/276</td>
<td>167/268</td>
<td>20.3</td>
<td>0.70</td>
<td>(0.49–0.99)</td>
</tr>
</tbody>
</table>

χ² 1.61 (df = 4) Z = 2.04

<table>
<thead>
<tr>
<th>Study</th>
<th>Experiment n/N</th>
<th>Control n/N</th>
<th>Peto OR (95% CI fixed)</th>
<th>Weight %</th>
<th>Peto OR (95% CI fixed)</th>
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<td>Treatment between three and six hours</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>ASK 1996</td>
<td>70/133</td>
<td>59/137</td>
<td>10.7</td>
<td>1.47</td>
<td>(0.91–2.36)</td>
</tr>
<tr>
<td>ECASS 1995</td>
<td>143/264</td>
<td>160/269</td>
<td>20.6</td>
<td>0.81</td>
<td>(0.57–1.13)</td>
</tr>
<tr>
<td>ECASS II 1998</td>
<td>148/328</td>
<td>167/314</td>
<td>25.3</td>
<td>0.72</td>
<td>(0.53–0.99)</td>
</tr>
<tr>
<td>MAST-E 1996</td>
<td>105/130</td>
<td>112/133</td>
<td>6.0</td>
<td>0.79</td>
<td>(0.42–1.49)</td>
</tr>
<tr>
<td>MAST-E 1995</td>
<td>150/234</td>
<td>131/223</td>
<td>17.1</td>
<td>1.25</td>
<td>(0.86–1.83)</td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>616/1089</td>
<td>629/1076</td>
<td>79.1</td>
<td>0.93</td>
<td>(0.78–1.10)</td>
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χ² 9.34 (df = 4) Z = 0.86

<table>
<thead>
<tr>
<th>Study</th>
<th>Experiment n/N</th>
<th>Control n/N</th>
<th>Peto OR (95% CI fixed)</th>
<th>Weight %</th>
<th>Peto OR (95% CI fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (95% CI)</td>
<td>762/1365</td>
<td>796/1344</td>
<td>100.0</td>
<td>0.87</td>
<td>(0.75–1.02)</td>
</tr>
</tbody>
</table>

χ² 12.99 (df = 9) Z = 0.69
cardioembolic sources, such as mitral valve disease without AF, is very difficult, with little evidence to guide the physician.

**ASPIRIN**

The pooled results of two very large randomised controlled trials comparing aspirin with placebo, concluded that medium dose aspirin (160–300 mg) started in the acute phase of an ischaemic stroke produces a small (13 fewer patients per 1000 dead or disabled) net benefit.70 Whether this benefit arose from an effect on the stroke itself or simply through earlier initiation of secondary prevention of stroke and other thrombotic complications is uncertain. We therefore start all patients on stroke and other thrombotic complications is earlier initiation of secondary prevention of disease without AF, is very difficult, with little evidence to guide the physician.

**NEUROPROTECTIVE AGENTS**

To date, no neuroprotective agent has been conclusively shown to be effective, and a Cochrane review summarising the current data is due to be published.71 Trials to evaluate the neuroprotective effects of magnesium (IMAGES), benzodiazepines (EGASIS), and other novel agents are in progress.

**OTHER TREATMENTS**

Numerous other treatments have been used for ischaemic stroke, and some have been subjected to randomised trials. However, there is currently no convincing evidence to support the routine use of any of them.

**Figure 2** Results of a systematic review of the randomised trials of anticoagulants in acute presumed ischaemic stroke. The estimate of treatment expressed as an OR (square, the size of the square indicating the statistical power of the estimate), and its 95% CI (horizontal bar); the diamond shapes provide estimates of the pooled trial results. OR=1 indicates a zero treatment effect, OR<1 indicates treatment better than control, and OR>1 indicates treatment worse than control. There was no significant effect of anticoagulant treatment on death or dependency at the end of follow up (>1 month).

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### Treatment of haemorrhagic stroke

Various specific treatments designed to reduce intracranial pressure are often used for primary intracerebral haemorrhage, including osmotic agents such as mannitol, urea or glycerol, steroids, or hyperventilation; unfortunately there is no convincing evidence that these treatments improve outcome. In view of the lack of evidence we do not routinely use any specific medical therapy in haemorrhagic stroke, nor do we employ invasive devices, such as intraventricular catheters, to directly measure intracranial pressure. We would attempt to correct or reverse any clotting abnormality, including those patients on oral anticoagulant drugs, although this depends on the original indication for the anticoagulants (for example, prosthetic heart valves).

### SURGERY FOR SUPRATENTORIAL PICH

A systematic review of open surgical drainage via a craniotomy concluded that this sort of surgery was positively harmful.72 However, safer surgical techniques are now available, in particular stereotactic aspiration, and the results of ongoing surgical trials are awaited. In a previously fit person with a large lobar intracerebral haemorrhage whose conscious level is falling we would refer to our neurosurgeons and encourage them to drain the haematoma. In this situation where the patients are expected to die unless action is taken the decision is relatively easy. More difficult are those patients with lesions deep in the hemisphere and those with severe impairments but no reduction in conscious level. These patients we
usually manage conservatively, or randomise into one of the ongoing trials of surgical treatment.

SURGERY FOR INFRATENTORIAL PICH

Although there is general agreement that surgical intervention in this situation may be life saving (so much so that a randomised controlled trial is unlikely to ever be done), there is considerable uncertainty about which patients might benefit the most, or even which procedure is optimal (haematoma evacuation versus ventriculotomy via a ventriculostomy, or both). We would always consider surgical intervention in any patient who was comatose, or whose conscious level was progressively deteriorating, and in whom other exacerbating causes had been excluded (table 3). Once brainstem reflexes have been absent for several hours however, death is inevitable.80

Organisation of stroke services

In the United Kingdom, between 40% to 70% of patients are admitted to hospital after a stroke81–83 and mostly cared for by general practitioners, or elderly patients, and those already in institutionalised care. Widespread introduction of thrombolytic therapy for ischaemic stroke, with further research. It seems likely that the sooner acute specific treatments can be given the more effective they will be (“time is brain”). It is likely that there will be increasing emphasis on systems of prehospital care which facilitate earlier transfer to an acute stroke unit. However, this must not inhibit the development of other aspects of the services (for example, rehabilitation) which have been shown to have important benefits for patients.

Therefore hospitals need to develop both inpatient and outpatient services in collaboration with primary care, which can respond rapidly. As an important adjunct to developing these services, the general public should be educated about the symptoms of stroke, and the importance of early presentation to medical services.

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Richard Davenport and Martin Dennis

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