Subarachnoid haemorrhage and myths about saccular aneurysms

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Introduction
The incidence of subarachnoid haemorrhage is about 10:100000 per year in Western Europe and USA and comprises some 10% of all cases of cerebrovascular disease.1 Forty five per cent of patients die within the first 30 days following subarachnoid haemorrhage; nearly two thirds die as a result of the initial haemorrhage and most within the first two days.2 Despite vigorous therapeutic efforts, the mortality rate from subarachnoid haemorrhage remains high. A greater understanding of the basic mechanisms and pathology of subarachnoid haemorrhage may lead to more effective prevention and therapeutic strategies.

This leader reviews the causes and results of subarachnoid haemorrhage and examines how myths surrounding subarachnoid haemorrhage and saccular aneurysms have been dispelled by the introduction of new techniques and more reliable clinical and epidemiological data.

Outcome of subarachnoid haemorrhage
Subarachnoid haemorrhage occurs mainly between the ages of 40 and 60 years and is more common in women than men by a proportion of 2–3:1.1 Characteristically, there is a sudden onset of severe headache, frequently concentrated in the occipital region; patients may or may not lose consciousness. Diagnosis is made by detecting blood in the subarachnoid space by computed tomography (CT) or by the presence of fresh or altered blood in the cerebrospinal fluid (CSF). The mortality rate in the first two days is 28%, mostly from the initial bleed; 45% of patients are dead by 30 days, either from the initial bleed or from re-bleeding.5

Most deaths from subarachnoid haemorrhage occur rapidly and the mortality rate is proportional to the amount of blood released. Those who have a haemorrhage of 15 cm³ or less have a better prognosis.6 A proportion of patients suffer massive intraventricular haemorrhage, usually from a ruptured anterior communicating artery aneurysm or intracerebral bleeding from a middle cerebral artery aneurysm.7 The exact cause of death when blood is confined to the subarachnoid space is unclear. Death may be associated with a rapid rise in intracranial pressure due to a sudden rush of blood at arterial pressure into the subarachnoid space.

Vasospasm of major cerebral arteries is seen mainly between two and seven days of subarachnoid haemorrhage or later. It is detectable by angiography7 and may significantly reduce blood flow, as measured by Doppler techniques.8 Vasospasm may complicate surgical clipping of an aneurysm and may result in extensive cerebral infarction; it is a major cause of death or disability in 13–5% of patients in hospital trials for the treatment of subarachnoid haemorrhage.7 However, arterial vasospasm and infarction contribute to only about 5% of all deaths during the first 30 days after subarachnoid haemorrhage.

Hydrocephalus may be seen in the acute stages of subarachnoid haemorrhage or may occur as a late complication. A small proportion of patients show ventricular dilatation on their first CT scan; hydrocephalus at this stage may be caused by impedance of CSF flow through the blood filled subarachnoid space. At a later stage, often years after the subarachnoid haemorrhage, patients may develop intermittently raised pressure (normal pressure) hydrocephalus with the insidious onset of dementia, ataxia and incontinence.9 Hydrocephalus at this stage is probably caused by organisation of the subarachnoid blood clot, resulting in fibrous adherence of the arachnoid mater to the pia and to the surface of the brain, possibly impeding the flow of CSF through the subarachnoid space. There is little evidence that the arachnoid granulations become occluded; although red blood cells may enter channels within arachnoid granulations following a subarachnoid haemorrhage,10 the granulations do not seem to become occluded by fibrous scar tissue.

Pathology of subarachnoid haemorrhage
Rupture of a saccular cerebral artery aneurysm accounts for 70–90% of all subarachnoid haemorrhages in subjects over 30 years of age.3 Other causes, such as ruptured arteriovenous malformations, intracranial artery dissections, mycotic aneurysms, trauma, bleeding disorders, and a cervical origin for haemorrhage should be considered in the 15–20% of patients in whom no aneurysm is discovered on initial angiogram.11 Although saccular aneurysms are
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Willis may reveal the aneurysm or may facilitate dissection of the aneurysm in the fixed brain. If the blood clot around the aneurysm is allowed to harden by fixation, the aneurysm may be difficult to locate or it may be damaged during the removal of the solidified blood clot.

Multiple aneurysms occur in 25% of cases. In the series of 109 ruptured aneurysms reported by Inagawa and Hirano, 17% were less than 4 mm in diameter, 40% were 5–9 mm, and 38% were 10 mm or larger in diameter. Giant aneurysms, greater than 25 mm in diameter, also occur. One third of aneurysms bleed and others present as mass lesions, either compressing the brain or resulting in temporal lobe epilepsy and in other focal neurological signs. Often with a thick fibrous and calcified wall, giant aneurysms may be filled with partly organised lamellated thrombus and appear as dense, rounded bodies on CT scan. Angiography may show a lumen very much smaller than the total capacity of the aneurysm.

If the results of angiography are available, they usually greatly facilitate the location of a saccular aneurysm at necropsy. Whether the location of the aneurysm is known or not, there are two major choices for examination of the brain. The circle of Willis and its major branches can be dissected free from the brain and the location of the aneurysm firmly identified on the exposed vessels. However, this procedure interferes with the relations among the aneurysm, the brain and the haemorrhage. Furthermore, if the ruptured aneurysm is very small, it may not be detectable on the dissected circle of Willis. Alternatively, therefore, it is often best to section the fixed brain coronally in 1 cm slices, find the point of haemorrhage to locate the aneurysm and then examine it macroscopically and histologically in relation to the artery and adjacent brain. In this way, the full effects of the ruptured aneurysm on the brain can be recorded more accurately.

In many cases of ruptured saccular aneurysm there is significant intraventricular or infra-cerebral haemorrhage. In a necropsy study of 133 patients with ruptured saccular aneurysms, intraventricular haemorrhage was observed in 40%, half of which were caused by rupture of an anterior communicating artery aneurysm either through the inferior medial portion of the frontal lobe or through the corpus callosum. Intracerebral haemorrhage was seen in 39% of cases, caused by middle cerebral artery aneurysms in half of these.

**Treatment**

Therapy in subarachnoid haemorrhage is directed mainly towards prevention of neuronal damage in the brain and rebleeding from the aneurysm (fig 2). Nimodipine, a calcium channel blocker, is administered within the first 96 hours in an attempt to prevent neuronal damage. Red blood cells entering the subarachnoid space following rupture of an aneurysm are prevented from passing into perivascular spaces around arteries in the brain by the pia mater. It is probable, however, that soluble factors from the blood penetrate the
perivascular spaces within the brain and result in ischaemia or direct damage to neurons in the cerebral cortex and central grey matter. Nimodipine helps to prevent such damage. The British nimodipine trial included 554 patients and has shown that if oral nimodipine, 60 mg every four hours, is started within the first 96 hours of a subarachnoid haemorrhage, there is a significant reduction in cerebral infarction and an improvement in outcome. As a result, virtually every neurosurgical centre in Britain uses this drug routinely. However, as a very significant number of deaths from subarachnoid haemorrhage occur within the first two days, there is often little opportunity for therapeutic intervention. A reduction in the mortality rate and neurological damage from subarachnoid haemorrhage may, therefore, be gained more by reducing risk factors such as smoking and hypercholesterolaemia and preventing the haemorrhage than by treating it.

Occlusion of an aneurysm as a measure to prevent rebleeding can be effected by clipping the neck of the aneurysm through a craniotomy (fig 2) or, in the case of inoperable giant aneurysms, by the occlusion or ligation of a major feeding vessel such as the carotid artery. However, there is now increasing interest in the endovascular insertion of detachable coils into aneurysms to induce thrombosis and ultimate obliteration of the aneurysm itself (fig 2). This technique was originally developed for the treatment of giant aneurysms but is increasingly used for smaller ones and is most successful when the neck of the aneurysm is less than 4 mm in diameter.

Myths
One of the common misconceptions is that vasospasm of cerebral arteries in the subarachnoid space is a major cause of death following subarachnoid haemorrhage. There is no doubt that vasospasm occurs and that it may result in decreased blood flow and infarction. The cause of vasospasm is not entirely clear and it may be due to the release of factors such as endothelin into the CSF following the haemorrhage. However, vasospasm accounts for only some 6% of deaths in the first 30 days following subarachnoid haemorrhage.

Two major myths are associated with the origin and formation of saccular aneurysms. One is that the aneurysms are congenital and the other is that hypertension is the major contributing factor.

Saccular aneurysms are still often referred to as “congenital” aneurysms but less than 10% are encountered in subjects under 30 years of age and less than 5% in those under 20 years of age. There is some familial clustering of aneurysms but this is uncommon. The major familial association of saccular aneurysms is with autosomal dominant polycystic kidney disease with an incidence of aneurysms of 10% in asymptomatic relatives. A number of other inherited disorders have been associated with intracranial aneurysms, many of which may be associated with structural abnormalities of the arterial walls. Aneurysms on cerebral arteries can be induced in experimental animals by a combination of altered haemodynamic stress and weakening of the arterial walls. This is effected by ligating the common carotid artery on one side, making the animals hypertensive and feeding them with the lathyrogen β-aminopropionitrile.

In the great majority of patients with saccular aneurysms, no predisposing factors can be detected clinically, although smoking and hypercholesterolaemia are important risk factors associated with ruptured cerebral aneurysms; a history of hypertension is of less importance.

The assumption that saccular aneurysms are congenital arose from the observation that gaps occurred in the smooth muscle tunica media of cerebral arteries at the carina where vessels branched. Similar gaps in the media have been described in arteries in other organs of the body. It was thought, however, that the gaps presented points of weakness which led directly to the formation of saccular aneurysms.

In a survey of cerebral artery bifurcations in subjects between the ages of 10 and 80 years of age, gaps in the media at the distal carina of bifurcations were found in 60% of cases, and the gaps enlarged with increasing age. The same survey showed that intimal pads, with duplication and fragmentation of the internal elastic lamina, formed at the bifurcations and proximal to bifurcations in arteries of older patients. Such pads were seen at 22% of bifurcations by 10 years of age and at 60% of bifurcations over the age of 30 years. The increase in luminal pads proximal to and at bifurcations of cerebral arteries with age has two major effects. Firstly, the pads induce focal loss of elasticity of the arterial wall and, secondly, the shape of the carina of the artery changes to become more rounded. Theoretically, the changes at the bifurcation could alter the direction of haemodynamic forces on the carina so that haemodynamic stress may be more directed at the rounded apex of the carina. In this way, the rounded carina is gradually pushed out to form an aneurysm at the point of vessel branching. Histological examination...
shows how the smooth muscle coat and the elastic lamina of the artery stop abruptly at the neck of the saccular aneurysm. The wall of the aneurysm is thus formed entirely from fibrous tissue and endothelium with or without thrombus coating its luminal surface. As aneurysms also occur at points at which vessels curve, similar forces may be in action in these regions.

The question still remains as to why aneurysms only form in 5% of the population. Development of aneurysms may depend on a combination of haemodynamic forces and the configuration of the vessel bifurcations. There is considerable individual variation in the pattern of vessels around the circle of Willis, particularly in the calibre of vessels such as the posterior communicating and anterior cerebral arteries; this could affect the pattern of stress on points of branching. Furthermore, there is a high incidence of aneurysms on arteries feeding arteriovenous malformations and aneurysms are associated with vascular abnormalities such as Moyra Moya disease, in which there is aggressive occlusion of the internal carotid artery and its branches with the formation of "net-like" vessels. These observations suggest that blood flow patterns may play a role in the formation of aneurysms, possibly by inducing intimal pads within the arteries, altering the shape of vessel branches, and thus increasing susceptibility to the development of aneurysms.

The last important question is why do aneurysms rupture? Thinning of the fibrous wall of the aneurysm may occur and rupture frequently occurs at the apex of the aneurysm. Although it was originally suggested that hypertension is a major factor involved in the formation of aneurysms, it appears to be less important than risk factors such as smoking and increased serum cholesterol. Subarachnoid haemorrhage tends to occur in ambulatory patients and following physical exertion; a sudden rise in blood pressure may therefore play a part in rupture of aneurysms. It has also been suggested that changing weather conditions precede aneurysm rupture in men but not in women. Men show a single large peak of aneurysm rupture in the late autumn, whereas in women the annual peak occurs in late Spring.

In conclusion, pathologists have an important role to play in increasing our understanding of the multiple factors in the environment, in the structure and branching patterns of the circle of Willis, and in the genetic make-up of subjects that result in the formation and rupture of saccular aneurysms. Such an understanding will allow preventative measures to be adopted to help reduce the very high mortality and morbidity associated with saccular aneurysms and subarachnoid haemorrhage.
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