PATHOPHYSIOLOGICAL MECHANISMS IN SUBARACHNOID HAEMORRHAGE

A Study of the Neuropharmacological, Physiological and Morphological Changes that Occur in a Model of Subarachnoid Haemorrhage Developed in the Laboratory Rat

by

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ABSTRACT

This thesis reviews current knowledge regarding the problems created by, possible causal factors, and present management of the disordered pathophysiology that may arise following subarachnoid haemorrhage (SAH) in man. Mechanisms responsible for the normal regulation of the cerebral vasculature are also reviewed. The present investigation explores how various neuropharmacological, physiological and morphological aspects of the major cerebral arteries are affected by such haemorrhage in a small animal model of experimental SAH developed in the laboratory rat. Preliminary immunohistochemical studies revealed that contrary to earlier reports, the extraparenchymal cerebral arteries do not receive a significant serotonergic innervation. Serotonin was only rarely present under normal circumstances in the neural plexus, and when found was invariably contained within nerves also identified as catecholaminergic in nature. Following SAH, major alterations in the neurotransmitter content of the cerebrovasculature occurred. The perivascular sympathetic nerves of major cerebral vessels rapidly accumulated serotonin, while a coincident depletion of neuropeptide Y took place. Using cortically implanted platinum-wire microelectrodes, with measurement of cerebral blood flow (CBF) by hydrogen-clearance, the timecourse of the global reduction in CBF that develops acutely in this model was documented. A 50% reduction in blood flow persisted for up to 3 hours posthaemorrhage, at 24 hours this was restored
to 85% of normal, and recovered fully by 48 hours. The fall in CBF developed independently of concommitant changes in intracranial and cerebral perfusion pressure, and it would appear likely that early vasospasm secondary to released blood products, rather than pressure changes per-se, is responsible for the acute cerebral ischaemia that develops.

Electron microscopic studies, demonstrated the delayed development of a mild cerebral vasculopathy, comprising focal areas of subintimal medial necrosis. Transformation of cells derived from the pia-arachnoid into macrophages, occurred on the second day post haemorrhage. These cells were then largely responsible for a rapid phagocytic removal of the subarachnoid blood clot. Previous findings obtained by workers using experimental models of SAH are reviewed, and suggestions as to the nature and aetiology of delayed cerebral ischaemia and vasospasm arising after SAH deduced.
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1. INTRODUCTION

1.1 THE CLINICAL PROBLEM

Epidemiology and Aetiology:

Subarachnoid haemorrhage (SAH) constitutes a major health problem. In England and Wales alone, an average of 3,000 deaths annually are registered as having resulted directly from SAH (OPCS, 1989). Placed into context by a comparison of the annual fatality figures; SAH accounts for 128 deaths per million of population per year, malignant neoplasms of the brain for 102, cardiac dysrhythmias for 67, and pulmonary emboli for 49 (OPCS, 1985). The incidence of fatalities from SAH rises steadily with age and there is generally a 2:1 female to male predominance (OPCS, 1985), except in the very young where this ratio is reversed (Meyer et al., 1989).

Ruptured intracranial aneurysms account for 75 to 80%, and arteriovenous malformations for around 5%, of all cases of diagnosed haemorrhage into the subarachnoid spaces (McKissock and Paine, 1959; Pakarinen, 1967; Locksey, 1969; Phillips et al., 1980). The aetiology of SAH also varies with age, in the over 30 years age group, by far the commonest predisposing cause is a ruptured aneurysm, however in patients under 20 years of age it is more likely to be due to a congenital arteriovenous malformation. Other causes of SAH include head injury, bleeding disorders, primary or secondary tumours, while in approximately 15% of cases the causation remains unknown despite extensive investigation (Hayward, 1977; Shephard, 1984). The prevalence of
intracranial berry aneurysms detected at autopsy varies at between 0.2 - 9.6% in different series, with the average lying around 5% (Crawford and Sarner, 1965; Pakarinen, 1967; Phillips et al., 1980; Sekar and Heros, 1981). Between 20 - 80% of such autopsy diagnosed aneurysms also reveal evidence to suggest that they have undergone leak or rupture in the past (Wilson et al., 1954; Wiebers et al., 1981). These figures would suggest that intracranial aneurysms and subarachnoid haemorrhage may be a more frequent occurrence than an examination of hospital admission and mortality statistics alone might suggest. Intracranial aneurysms include the saccular or berry aneurysm, fusiform dilatations, mycotic, traumatic and dissecting aneurysms, as well as microaneurysms of Charcot and Bouchard (Ross Russell, 1963).

By far the most frequent cause underlying a SAH however, is the saccular or berry type of aneurysm. Some explanation of the structure of cerebral arteries and aneurysms is first necessary, in order to understand the mechanisms proposed for the formation of the latter. Cerebral arteries are composed of three layers - an outer collagenous adventitia, a circular smooth muscle media, and an inner intimal layer (Dahl, 1973). The intima is composed of a single layer of endothelial cells, resting upon a fenestrated internal elastic lamina. These fenestrations have a mean diameter of 2 um generally, but at vessel bifurcations there are a smaller number of larger (7 um) fenestrations that constitute a potential wall weakness (Campbell and Roach, 1981). The tunica media consists of several layers of
circularly arranged smooth muscle cells that diminish in number as the vessels decrease in size. The media at arterial branchings commonly shows defects or discontinuities in the muscle layer (Forbus, 1930). Although similar gaps occur in arteries elsewhere in the body (Hassler, 1963), no external elastic lamina as possessed by the extracranial arteries, is present in cerebral arteries (Crompton, 1976). Such defects, presumably must at least partly account for the prediliction of berry aneurysms to occur at branch points (Crompton, 1966). In the presence of an intact internal elastic lamina however, medial defects alone, do not appear to compromise the strength of the arterial wall (Glynn, 1940). Additional degeneration or weakness of the internal elastic lamina must therefore arise for aneurysm formation to occur. A saccular cerebral aneurysm arising at the apex of such an arterial bifurcation is generally thin walled, on histological examination consisting of an intima and adventitia alone and characterised by the virtual absence of any medial smooth muscle layer. The internal elastic lamina is always grossly diminished, usually being thinned and severely fragmented (Stehbens, 1975; Sekhar and Heros, 1981).

**Predisposing Factors**

**Arterial hypertension** increases haemodynamic stress and aggravates degenerative changes within the arterial wall. In clinical studies, conflicting findings were initially found regarding the effects of hypertension upon cerebral aneurysm formation and rupture (Black and Hicks, 1952; McCormick and Schmalstieg, 1977; Wiebers et al., 1981). Most of these
series are open to criticism, in that evidence of preictal hypertension was obtained by retrospective means, such as evidence of cardiac hypertrophy on ECG or at autopsy. In the Framingham study (Sacco et al., 1984) however, in which a defined population was studied prospectively, hypertension was found to be a definite risk factor, both for aneurysmal rupture and fatality of outcome.

Smoking The risk of aneurysmal SAH in smokers versus non-smokers, has been analysed both for men and women. In the case of the latter, this risk has also been examined in relation to use of oral contraceptives. Heavy cigarette smoking of >20/day appears to be significantly associated with SAH as a risk factor in women, although not in men (Bell and Symon, 1979; Sacco et al., 1984). Concurrent usage of cigarettes and oral contraceptives may synergistically raise the risk of SAH (Petitti, 1978; Sacco et al., 1984).

Other diseases Intracranial aneurysms have not infrequently been reported in association with polycystic renal disease, coarctation of the aorta, and connective tissue disorders such as Marfans, Ehlers-Danlos and pseudoxanthoma elasticum (Dixon, 1951; Bigelow, 1953; Stehbens, 1962; Rubenstein and Cohen, 1964; Stehbens et al., 1989). Such associations do not necessarily imply a causal predisposition, and no significant increase in incidence has actually been demonstrated in the above conditions. In some disorders, such as polycystic kidneys, any increased incidence might simply reflect the higher incidence of hypertension that occurs in this condition.
Connective tissue defects The fibrous structures of collagen, elastin and reticulin play an important role in the load-bearing capacity of the arteries. It has been reported that a deficiency may exist, both in levels of type III collagen and the amounts of reticular fibers, present within cerebral vessels from some patients suffering from ruptured intracranial saccular aneurysms (Neil-Dwyer et al., 1987; Ostergaard and Oxlund, 1987; Ostergaard et al., 1987; de Paepe et al., 1988). The cerebral vessels from such patients may also demonstrate altered mechanical properties with significant increases in arterial wall extensibility (Ostergaard and Oxlund, 1987).

Genetic factors A low familial incidence (0.7 - 3.2%) has been observed for intracranial saccular aneurysms (Carroll and Haddon, 1964; Hashimoto, 1977; Fukawa and Aihara, 1987). Within such families, the frequency of occurrence suggests an autosomal dominant mode of inheritance for some predisposing factor. Few investigations of associations between the major histocompatibility complex and saccular aneurysms in the general population have been performed. An increased occurrence of the HLA-DR2 antigen and C3-F gene has however been reported in patients with diagnosed ruptured intracranial aneurysms (Ostergaard et al., 1986a; Ostergaard et al., 1986b).

Experimental induction

Interesting insights into the pathogenesis of intracranial aneurysms, have been provided by a series of experimental studies in which aneurysm formation was successfully provoked, both in the monkey and in rats.
Cerebral aneurysms were produced by a combination of inducing systemic hypertension, ligating the common carotid artery on one side to produce a flow change, together with feeding on beta-aminoproprionitrile which causes defects in the linkages between the elastin and collagen of tissues.

Summary

Analysis of clinical observations and experimental studies, now strongly suggests that intracranial saccular aneurysms are acquired as the result of a complex interplay of predisposing factors and conditions. The most important of these would appear to be haemodynamic stress, arterial hypertension, and a derangement of the connective tissues - particularly affecting the internal elastic lamina. Because the media must also be absent or weakened for an aneurysm to develop, these tend naturally therefore to form at arterial branch points, where a discontinuity in the media and weakness in the elastica is already present.

Pathophysiology and Natural History:

Release of blood into the subarachnoid space from a ruptured vessel will result in an acute rise in intracranial pressure (ICP), and if bleeding continues, will reach a point within one to two minutes where the intracranial pressure equals diastolic blood pressure and effective cerebral perfusion is lost (Nornes, 1973). Unless a large haematoma is formed, ICP will then start to fall, and providing a fibrin-platelet plug has formed at the leak
point so bleeding does not recommence, perfusion can resume, averting immediate cerebral death. Severe primary brain damage may still have been occasioned however, by pressure effects upon the midbrain and brain stem or more directly from parenchymal disruption and haematoma formation within brain tissues adjacent to the point of blood release. Survival and morbidity thus far, will largely depend upon the severity and location of the initial bleed.

An accurate assessment of the natural history of subarachnoid haemorrhage is not easily attained, that which exists relates largely to the outcome of aneurysmal haemorrhage, and series based upon patients reaching neurosurgical centres will grossly underestimate the overall mortality and morbidity of the condition. It is necessary therefore, to look instead at defined population studies. One large community study in Finland (Pakarinen, 1967), found that 43% of patients died shortly after haemorrhage, and in those surviving their initial bleed another 35% died within one year. Combining the findings of a number of other published series, it appears that approximately 12% of patients that bleed from a saccular aneurysm will be dead before reaching hospital. Without treatment, the one month mortality in those surviving the initial ictus is approximately 50% with a significant morbidity of about 25%, while at the end of one year, mortality rises to around 60% (Crawford and Sarner, 1965; Jane et al., 1977; Phillips et al., 1980; Ljunggren et al., 1984; Rosenorn et al., 1987a).

In a large number of cases therefore, it appears to be the potentially preventable complications and
pathophysiological changes that can develop subsequent to a SAH, rather than the primary bleed itself, that will determine eventual patient outcome - both in terms of survival and of lasting disability. In the past, rebleeding was considered as practically the sole cause of delayed neurological deterioration in patients that had recently suffered an aneurysmal SAH. More recently, evidence whilst confirming that approximately two-thirds of sudden deteriorations are due to rebleeds, has also shown both that a number of acute deteriorations and the majority of cases in which neurological deterioration develops more gradually, are in fact due to the development of cerebral ischaemia (Maurice-Williams, 1982; Mickey et al., 1984; Mohsen et al., 1984; Vermeulen et al., 1984a).

Rebleeding

Approximately 20% of patients surviving an aneurysmal SAH will suffer a further bleed within two weeks, and up to 60% within the six months after the first haemorrhage, unless surgically treated (Locksey, 1966; Pakarinen, 1967; Jane et al., 1977; Kassel and Torner, 1983). Such repeated aneurysmal subarachnoid bleeding is associated with a greatly worsened prognosis, the overall mortality rate being nearly double that of those patients that do not rebleed (Rosenorn et al., 1987). The peak incidence for such rebleeds has been the subject of controversy. It is generally reported that the highest risk of such rebleeding occurs around the end of the first week following the initial event (Locksley, 1969; Nibblelink et al., 1975; Vermeulen et al., 1984; Rosenorn et al., 1987b). However,
two other studies have found that the highest rate of rebleeding occurs on the same day as the initial haemorrhage, with no later peak being identified (Kassel and Torner, 1983; Jane et al., 1985). Most authors however are in agreement, that the incidence of rerupture is highest in the first three weeks and, that after six months the risk of rebleeding for an untreated aneurysm levels off at around 3% per year (Winn et al., 1977; Kassell and Torner, 1983; Jane et al., 1985). After the first decade, this has been reported to fall further, to just under 1% per year, which is the same as for that of an unruptured incidental aneurysm (Nishioka et al., 1984). The risk of rebleeding from an arteriovenous malformation has been less extensively studied, but would appear to be approximately 6% in the first year then around 2 - 3% per year subsequently (Graf et al., 1983).

Certain diagnosis of rebleeding as the cause of an acute deterioration requires care. Although serial CT scans would appear to be of considerable value here in excluding other causes for deterioration, and confirming the correct diagnosis (Van Gijn and Van Dongen, 1980).

Cerebral Ischaemia, Blood Flow and Autoregulation

Clinically, cerebral ischaemia developing post-SAH is heralded by a gradual worsening in the patients level of consciousness, together with the onset of new focal neurological signs. Approximately 30% of all patients will develop delayed cerebral ischaemia post-SAH, the peak incidence being at around seven to ten days after the initial bleed (Pickard et al., 1989). The risk in the
individual patient appears to be directly related to the amount of blood released, and rises to about 60% for those patients with the most blood on CT scanning (Davis et al., 1980; Mizukami et al., 1980; Mohsen et al., 1984). Delayed cerebral ischaemia has been considered by some authors as actually causing more death and disability in patients admitted to neurosurgical units following SAH, than rebleeding itself (Illingworth, 1979; Maurice-Williams, 1982; Kassel et al., 1985). Repeat computed tomography in such cases usually demonstrates areas of low attenuation, and measurements of cerebral blood flow (CBF) reveal areas of well defined regional flow decrease (Ferguson et al., 1972; Symon et al., 1972; Grubb et al., 1977; Ishii, 1979; Mohsen et al., 1984; Mickey et al., 1984). Overall, CBF tends to decline in the first week after SAH and to remain low over the next two weeks. The reduction in flow is usually most marked in the older patients and in those that are drowsy or in poor clinical grades (Meyer et al., 1982; Meyer et al., 1983; Mickey et al., 1984). The exact mechanism whereby such a decreased CBF and delayed cerebral ischaemia arises is uncertain, although it is likely to be multifactorial in aetiology. One major factor would appear to be the development of angiographically detectable luminal arterial narrowing or "vasospasm" (vide infra). Delayed hypoperfusion is however also clearly associated with raised intraventricular pressure and ventricular dilatation (Ferguson et al., 1972; Ishii, 1979), falls in patient blood pressure, hypovolaemia, and surgical manipulation (Symon, 1978; Pickard et al., 1980; Solomon et al., 1984)
After subarachnoid haemorrhage a loss of the normal autoregulatory capacity of the cerebral circulation may arise. In health, cerebrovascular reflexes are capable of maintaining a constant blood flow to the cerebral tissues despite variations in perfusion pressure of between 60 - 150 mmHg (Lassen, 1959; Harper, 1966; Symon, 1978). Cerebral autoregulation to arterial hypotension appears though to be disturbed, even in mild cases of SAH. Generally the degree of vasoparalysis is related to the patients clinical grade, and becomes increasingly impaired in those patients in poorer neurological condition (Alvund et al., 1972; Voldby, 1988; Tenjin et al., 1988). As well as affecting autoregulation to pressure, subarachnoid haemorrhage may also impair cerebrovascular reactivity and blood flow changes produced by alterations in arterial pCO₂. Unlike autoregulation to pressure, this effect appears to be confined only to those patients in the worst clinical grades, with severe tissue acidosis. Here both modes of reactivity are affected - so called total vasoparalysis (Voldby et al., 1985a; Dernbach et al., 1988; Voldby, 1988).

Cerebral Vasospasm

A persistent constriction of cerebral arteries that followed SAH, was first described about 1950 (Robertson, 1949; Ecker and Riemenschneider, 1951). Considerable clinical and radiographic evidence now exists, to link the onset of such arterial narrowing with the development of delayed cerebral ischaemia (Allcock and Drake, 1965; Loach and De Azevedo-Filho, 1976; Fischer et al., 1977; Ishii,
1979; Voldby et al., 1985b). Although it was initially reported that vasospasm appeared relatively frequently, early in the period following SAH (Du Boulay, 1963; Allcock and Drake, 1965), more recent studies using greater patient numbers have shown it to be predominantly a delayed phenomena (Saito et al., 1977; Weir et al., 1978; Kwak et al., 1979). It is rarely seen within three days of a presenting bleed, has its onset about the fourth or fifth day after SAH, is maximal throughout the second week and largely gone by the end of the third week. A degree of delayed vasospasm is seen in approximately 50% of all cases of SAH from aneurysm rupture, however severe spasm, defined as a reduction in vessel luminal diameter of greater than 50% from normal, appears to affect only about 10% of patients (Boullin, 1985; Kassell, 1985).

The term vasospasm, readily came to be employed to describe such angiographically demonstrated delayed arterial luminal narrowing, as early investigators naturally assumed that it represented an active contraction of the medial smooth muscle of the vessels concerned (Simeone et al., 1968; Sundt et al., 1977). Such arterial narrowing has proven remarkably resistant to attempts to produce effective lasting reversal or prevention of such spasm, despite the use of a wide variety of pharmacological agents with known vasodilatory actions (Wilkins, 1980). In the light of these findings, and on the basis of a number of histopathological studies upon fresh and postmortem material, an alternative hypothesis proposes that delayed luminal narrowing represents a proliferative vasculopathy (Crompton, 1964;
Conway and McDonald, 1972; Mizukami et al., 1976; Peerless et al., 1980; Smith et al., 1985). These investigators and many others, having observed the occurrence of an early phase of medial necrosis followed by later subintimal thickening and fibrosis within affected vessels.

Vasospasm in experimental animal models commonly displays a biphasic pattern. An initial phase of marked constriction over hours, being followed by a phase of less intense but more prolonged narrowing (vide infra). An early phase of immediate vasospasm has been documented in man during angiography, when aneurysm rupture occurred under the procedure (Liliequist et al., 1976; Mohr and Kase, 1983). Odom (1975) however, saw no immediate vasospasm in four cases of aneurysm rupture during angiography, while Weir et al. (1978) found no spasm in 106 patients who had angiographic studies performed within 24 hours of their SAH. It is likely therefore, that an initial phase of acute vasospasm, unlike delayed vasospasm, is either uncommon or lasts for a few hours only in man.

Cerebral Metabolism

Under normal physiological conditions, CBF and cerebral metabolism are closely coupled, increases in cerebral oxygen utilisation (CMRO₂) leading to an increased blood flow. Conversely, a reduction in CBF will be compensated for by a rise in oxygen extraction and the arteriovenous difference of oxygen (AVDO₂) will increase (Lassen, 1959; Voldby et al., 1985). Subarachnoid haemorrhage produces a global decrease in CMRO₂ in all patients, but this is more marked in those with severe neurological deficits (Fein, 1975;
Grubb et al., 1977; Martin et al., 1984). Studies in which CBF, CMRO₂ and AVDO₂ have been simultaneously measured, reveal that this fall represents a true suppression of the oxidative metabolic rate, and that an uncoupling of CBF from cerebral metabolism occurs (Voldby et al., 1985b; Volby, 1988).

Cerebral Blood Volume

A striking finding post-SAH is the major increase in cerebral blood volume (CBV), up to nearly 60% above normal, that is seen in patients in poor clinical grades with severe neurological deficits and marked cerebral vasospasm on angiography. Such changes have been reported to be more consistently related to clinical progress, than measurements of cerebral blood flow (Eversden et al., 1977; Grubb et al., 1977). Using positron emission tomography, Martin et al. (1984) have demonstrated that patients without cerebral arterial narrowing show little change in CBV as compared with normal subjects, while patients with vasospasm consistently demonstrate marked increases in CBV. The large increases seen, suggests that while cerebral vasospasm may consist of a constriction of the large radiographically visible extraparenchymal vessels, that it is accompanied also by a massive dilation of intraparenchymal vessels (Grubb et al., 1977). Experimental studies have confirmed that a persistent elevation of CBV develops post-SAH, and have also revealed that a transient acute reduction in CBV occurs in the period immediately following the actual haemorrhage itself (Kuyama et al., 1984).
Hydrocephalus and raised Intracranial Pressure

Release of blood into the subarachnoid spaces, may lead to the development both of immediate acute hydrocephalus, and a more delayed chronic persistent hydrocephalus (Heros RC, 1984; Ojemann et al., 1987). Until quantitative methods for the assessment of ventricular dilation had been employed, the reported incidence of hydrocephalus post-SAH varied widely between 14 and 85 percent (Foltz and Ward, 1956; Vassilouthis and Richardson, 1979; Wenig et al., 1979; Kassel et al., 1984). However, more recent studies in which ventricular dilation has been carefully defined, using for example the bicaudate index on CT scanning, have shown the incidence of hydrocephalus on initial CT to be in the order of 20% (Van Gijn et al., 1985; Milhorat, 1987; Hasan et al., 1989). The pathophysiology of the disordered CSF dynamics that arise post-SAH are complex. In those patients in whom an intraventricular clot forms, an obvious mechanically obstructive element may occur immediately. Similarly, extensive clot in and around the basal cisterns can acutely obstruct both ventricular outflow and flow around the hemispheres. More distally, obstruction to CSF outflow and reabsorption at the level of the arachnoid villi, caused by an accumulation of erythrocytes within them, has been described (Ellington and Margolis, 1969). In some 10% of patients, a delayed hydrocephalus of a chronic communicating type will develop despite the clearance of all visible blood clot from the CSF containing spaces on CT scanning (Yasargil et al., 1973). In this latter instance it would appear that varying degrees of subarachnoid fibrosis develop, resulting
in persistent CSF outflow obstruction and impairment of reabsorption (Kibler et al., 1961; Ischii et al., 1979). In this context, it is relevant to note that treatment with the antifibrinolytic agent e-aminocaproic acid actually appears to increase the incidence of such hydrocephalus (Park, 1979; Kassell et al., 1984).

**Blood Brain Barrier**

The blood brain barrier (BBB) is an essential structure in the maintenance of a constant environment for the normal functioning of the tissues of the central nervous system. Blood-arterial wall barrier damage, and increases in capillary permeability can occur in a variety of conditions including; hypertension, ischaemia, trauma, seizures, and raised intracranial pressure (Rinder et al., 1968; Haggendal et al., 1970; Howse et al., 1974; O'Brien et al., 1974). Following subarachnoid haemorrhage, abnormal post-contrast enhancement can be seen in the region of the basal subarachnoid cisterns and occasionally far more extensively over the leptomeningeal surface (Moran et al., 1978 Davis et al., 1980; Sobel et al., 1981). Such changes have been considered to represent BBB breakdown with an increased permeability of the cerebral arterial wall. The actual degree of post-contrast enhancement seen being found by a number of authors, to be variously correlated with the development of neurological deficits, the occurrence of cerebral vasospasm, and poor patient outcome (Hirata et al., 1982; Tazawa et al., 1983; Doczi et al., 1984).
Hyponatraemia, Hypovolaemia and Central Neurendocrine Changes

Following subarachnoid haemorrhage, and in a variety of other intracranial disorders, patients are prone to the development of disturbances both in their sodium metabolism and circulating blood volume. The hyponatraemia seen in such cases was considered originally to represent a state of cerebral salt wasting. An impaired ability of the kidney to reabsorb sodium, despite a depressed concentration of sodium in the plasma, was thought to perhaps be due to the loss of some as yet unknown, cerebral salt conserving factor (Peters et al., 1950; Welt et al., 1952; Cort, 1954). The description however, by Schwartz et al. (1957) of a syndrome of inappropriate secretion of antidiuretic hormone (SIADH) that caused hyponatraemia and persistent natriuresis in patients with bronchogenic carcinoma, was followed by a general acceptance that SIADH was the probable cause also in many neurosurgical instances including SAH (Joynt et al., 1965; Hald et al., 1967; Maroon et al., 1970; Fox et al., 1971; Wise, 1978; Doczi et al., 1981). Indeed, in a number of such patients, actual measurement of the levels of the central neuroendocrine antidiuretic hormone - vasopressin, revealed an increased secretion of hypothalamic neuron derived vasopressin into the blood and CSF following SAH (Maroon and Nelson, 1979; Mather et al., 1981).

The important difference between these two aetiologies, is that in cerebral salt wasting there will be progressive sodium and extracellular volume contraction, whereas in SIADH there should be extracellular volume expansion. Some patients with SAH would appear to meet the diagnostic
criteria for SIADH; with hyponatraemia, hypo-osmolar plasma, hyperosmolar urine, and urinary sodium concentrations of >25 Meq/L in the presence of normovolaemia. However, latterly other studies upon patients with hyponatraemia following SAH, have clearly shown that a state of hypovolaemia with reductions in plasma volume (PV), red cell volume (RCV), and total circulating blood volume (TCBV), are in fact considerably more common (Maroon and Nelson, 1979; Nelson et al., 1981; Nelson et al., 1987). This would suggest therefore that in the majority of patients, SIADH is not the cause for the low sodium observed, and explains no doubt, why a regimen of fluid restriction to attempt to correct hyponatraemia post-SAH has been strongly associated with an increased incidence of cerebral infarction (Wijdicks et al., 1985).

The primary defect in the cerebral sodium wasting syndrome is as yet unknown. The central nervous system can affect sodium balance via neural, and by central neuroendocrine mechanisms. Both renin release and sodium reabsorption from the proximal tubule of the kidney can be altered by stimulation of the renal nerve (DiBona, 1985). The hypothalamus and associated nuclei control the thirst response, regulate ADH release and more recently have been shown to possess neurons containing atrial natriuretic polypeptide (Kawata et al., 1985; Standaert et al., 1986). Atrial natriuretic polypeptide (ANP) appears to play a role both in central and peripheral regulation of sodium homeostasis, and is largely produced and released from the atrial myocardium (Thibault et al., 1983). Recently, it has
been demonstrated that substantial elevations in plasma and CSF ANP levels, may arise in patients following SAH (Diringer et al., 1988; Doczi et al., 1988).

**Autonomic Dysfunction**

Indirect evidence of altered autonomic nervous system dysfunction following subarachnoid haemorrhage, is frequently seen. Arterial hypertension is extremely common, cardiac arrhythmias may occur and electrocardiographic alterations are well described. Direct evidence of sympathetic overactivity was first obtained, following the demonstration of an increased urinary excretion of free catecholamines and their metabolites in patients after SAH (Meyer et al., 1973; Neil-Dwyer et al., 1974). Elevated levels of circulating catecholamines and monoamine metabolites have been a uniform finding, some authors also documenting a correlation between their plasma concentrations, clinical grade and eventual outcome (Peerless and Griffiths, 1975; Benedict and Loach, 1978; Minegishi et al., 1987). Similarly, local amine levels in the cerebrospinal fluid have also been shown to be elevated following SAH (Cummins and Lothian, 1973), the increases being most marked in those patients with delayed cerebral ischaemia (Shigeno, 1982). Although considerable evidence now exists regarding the central sympathetic changes in animal studies, little direct information has to date been obtained in man. Tsukahara and his coworkers (1987) however, have shown that the noradrenaline content of human cerebral arteries is greatly reduced and also demonstrated concommitant alterations in cerebrovascular smooth muscle
Cardiovascular manifestations of subarachnoid haemorrhage have been recognised since 1947, when Byer et al. described large upright T waves and long Q-T intervals in a patient following SAH. Subsequent studies have confirmed that electrocardiographic (ECG) abnormalities are seen in at least 50% of patients with aneurysmal SAH, the commonest being broad or inverted T waves, Q-T prolongation, S-T segment elevation or depression and prominent U waves (Marion et al., 1986). Post-mortem studies and measurements of CPK isoenzymes indicate that SAH can actually result in subendocardial ischaemia and focal areas of necrosis (Koskelo et al., 1964; Doshi and Neil-Dwyer, 1977). Arrhythmias, both ventricular and supraventricular may develop (Wong and Cooper, 1969; Galloon et al., 1972) and can be life threatening (Parizel, 1973; Estanol and Marin, 1975). The mechanism by which these cardiac changes occur is less clearly understood, although the alterations in catecholamine levels are widely assumed to be implicated. It is certainly the case that an increased systemic catecholamine output post-SAH correlates well both with ECG changes and myocardial damage (Reichenbach and Benditt, 1970; Feibel et al., 1976).

As well as cardiac changes, acute pulmonary oedema has frequently been documented as developing in association with SAH (Ducker, 1968; Ciongoli et al., 1972). The actual incidence of pulmonary oedema following fatal SAH from aneurysm rupture is particularly high, at around 30% (Crompton, 1964; Weir, 1978). Acute pulmonary oedema is a
common event in those patients with acutely elevated ICP. The underlying cause is probably due to an inability, in some cases, of the heart to cope with the central neurogenically driven, sudden surge in sympathetic activity that leads to massive increases in systemic blood pressure, peripheral and pulmonary capillary resistance. While assisting the perfusion of the compromised brain, these extreme changes may lead to an acute failure in cardiac output, followed by left atrial distension and a transmitted rise in pulmonary venous pressure (Sarnoff and Sarnoff, 1952; Ducker and Simmons, 1968; Theodore and Robin, 1976).
1.2 PRESENT MANAGEMENT OF SUBARACHNOID HAEMORRHAGE

Diagnosis:

The management of patients surviving SAH, will naturally first include a confirmation of the correct clinical diagnosis having been made. Computed tomography (CT) is at present the initial investigation of choice, it will reveal blood within the subarachnoid spaces in approximately 90% of patients who have suffered a recent SAH (Scotti et al., 1977), as well as suggesting both the cause and the site of the bleed. In cases of uncertainty a lumbar puncture will show evenly bloodstained CSF, with a xanthochromic supernatant present in addition, if performed 12 to 24 hours after the bleed (Walton, 1956). Subarachnoid blood is rapidly removed however, and even a few days delay may make the diagnosis less certain since both the CT and CSF may now be clear. Persisting xanthochromia is a useful diagnostic aid here, even moderate degrees remain readily detectable by spectrophotometry, although it may also be detected by the naked eye alone for an average of 23 days (Walton, 1956; Van der Meulen, 1966). Subsequent angiography and visualisation of the vessels of the cerebral circulation, remains at present the essential investigation in order to identify the source of the bleed prior to definitive treatment.

Surgical Treatment:

Historical background

Treatment of SAH, due to aneurysm rupture and most other causes, is primarily a neurosurgical one. The procedures
employed aim to isolate the relevant pathology from the normal cerebral circulation, in order to prevent further bleeding. The first surgical procedure carried out on a patient suffering from SAH, due to a ruptured intracranial aneurysm, was performed in 1931 by Norman Dott at Edinburgh who successfully wrapped the aneurysm with muscle (Dott, 1933). Then in 1937, Walter Dandy foreshadowed the modern management of aneurysmal SAH, when, by using a McKenzie silver clip, he was the first person to selectively occlude the neck of an intracranial aneurysm (Dandy, 1938). Dandy and others (Dandy, 1944; Falconer, 1950; Norlen and Olivecrona, 1953) went on to promote a more widespread policy of neurosurgical intervention, with the use of both clipping and wrapping of intracranial aneurysms, for the prevention of further bleeding after SAH. Despite the enthusiasm of these early pioneers, there was little evidence at this time to indicate that the outcome following intracranial surgery, was in any way superior to that of a period of bed-rest alone (McKissock et al., 1958). It was only in 1960, that a controlled trial, comparing the results of conservative against surgical treatment, in SAH due to ruptured internal carotid artery aneurysms, was published (McKissock et al., 1960). This study demonstrated for the first time, that a lowering of morbidity and mortality, could be achieved in such cases, through surgical intervention to prevent rebleeding. Subsequently, the introduction of the operating microscope with application of microsurgical techniques to such operations (Pool and Colton, 1966; Krayenbuhl et al., 1972; Yasargil and Fox,
1975), together with the development of self-closing crossed action arterial clips (McFadden - review, 1989), has led to further improvements in the results of surgical treatments over conservative therapy. The mortality ascribed to operative intervention for aneurysmal SAH having been reduced as a consequence of such advances, to below 3% since the end of the seventies (Yasargil, 1984).

Timing of surgical intervention

The timing of surgical intervention for aneurysmal SAH has been, and continues to be, the subject of great interest and debate (Drake, 1978; Sano and Saito, 1979; Kassel and Drake, 1982; Flamm, 1986; Ljunggren and Brandt, 1986; Maurice-Williams, 1987). Early surgery obviates the risk of rebleeding, allows the removal of the subarachnoid clot, and permits the use of aggressive volume expansion and hypertensive therapy. Against this must be weighed the greater technical risks posed by brain swelling, and an increased risk of intraoperative aneurysm rupture. Later surgery carries a lower perioperative mortality, but a greater patient wastage may occur from rebleeding and other complications during the period of delay. In assessing the results of surgery in SAH performed at different surgical timings, it is important to apply the concept of total management morbidity and mortality, as opposed to surgical morbidity and mortality alone (Lougheed, 1969). The International Cooperative Study on the Timing of Aneurysm Surgery, was set up in an attempt to resolve the issue, but failed to demonstrate a clear advantage for either modality (Adams, 1986). There have been few non-randomised
retrospective, and only one prospective randomised trial of different timing policies, to date. Chyatte et al. (1988) in a retrospective analysis of Mayo Clinic records, found there was little difference in either morbidity or mortality. Ljunggren et al. (1988) comparing the results of an early surgical treatment policy carried out in Sweden against later surgical intervention in Denmark, found a similar morbidity but lower mortality with earlier treatment. In the one prospective study published (Ohman and Heiskanen, 1989), 216 patients were randomised to either acute (0-3 days), intermediate (4-7 days) or late surgery (>8 days). This analysis revealed no significant differences in the mortality rates between the operation groups, although morbidity was probably higher in the intermediate group. The surgeons dilemma therefore has apparently not changed since Maurice-Williams (1987) wrote, that "Until there is a large scale randomised trial of different timing policies applied to a single patient population the matter is likely to remain unsettled".

Evacuation of intracerebral and subarachnoid clot

A special group of patients constitute the 4-17% of all cases of aneurysmal SAH in which a clinically significant intracerebral haematoma (ICH) occurs (Bohm and Hugosson, 1978; Sano, 1979; Tapaninaho et al., 1988). The surgical treatment of such cases has been contentious, some authors reporting the results of acute surgical intervention under such circumstances as differing little from those of conservative management (Pia, 1979). Others have recommended urgent evacuation of the haematoma as the sole procedure
(Lougheed and Marshall, 1973; Symon, 1988) while a further body of opinion favours a policy of acute evacuation of ICH and clipping of aneurysm at the same time (Ljunggren et al., 1981; Wheelock et al., 1983; Tapaninaho et al., 1988). Mortality in the group of patients treated either conservatively or by evacuation of ICH alone, has been reported as between 69-100%. In those patients treated by evacuation of ICH with simultaneous clipping, the mortality rate has varied widely between 0-100%. One randomised prospective study (Heiskanen et al., 1988) that compared the results of conservative therapy against emergency ICH evacuation and definitive clipping, found a mortality rate of 80% in the conservative group and 27% in the surgical group, the difference being statistically significant.

Clearly however, it is not only mortality figures alone that should be considered when assessing patient outcome from such therapies, but also the quality of life in those that survive. This has been seldom analysed or commented upon in papers on this subject. In the series of Wheelock et al. (1983), eleven moribund patients underwent combined clipping of their aneurysm and evacuation of an associated ICH. Although there were six survivors, only one patient recovered sufficiently to be discharged home. Differences in survival outcome appear to exist, depending upon the vessel of origin of the aneurysm from which ICH arises. Auer (1985) reported four patients in each of whom a symptomatic ICH arose from the anterior cerebral circulation. Despite emergency evacuation of clot and successful clipping of the aneurysm all cases died. More encouraging results would seem
achievable however, in the salvage of patients moribund from temporal lobe ICH due to middle cerebral artery aneurysm rupture. Brandt et al. (1987) reported a series of four patients, with abnormal respirations and decerebrate posturing, in whom CT scan revealed SAH with a large ICH characteristic for a middle cerebral artery aneurysm. Rather than delay surgery by performing angiography, these cases were taken immediately to theatre for evacuation of haematoma and clipping of the expected underlying aneurysm. In this series three of the four patients did recover sufficiently to return home.

The volume and location of actual subarachnoid blood clot as revealed by CT scanning, has been shown to be directly related to the risk of subsequently developing delayed cerebral arterial narrowing and cerebral ischaemia (Davis et al., 1980; Mizukami et al., 1980; Mohsen et al., 1984). A number of surgeons therefore, have advocated early surgical evacuation of subarachnoid haematoma as a potential means of preventing such complications (Johnson et al., 1958; Dolenc et al., 1982; Mizukami et al., 1982). The feasibility of achieving a significant, early removal of subarachnoid blood has been demonstrated both by cisternal irrigation with artificial CSF (Kennady, 1966) and by using a blunt-tip suction tube (Wakabayashi and Fujita, 1984). Experimental evidence, obtained in a primate model of SAH, has suggested that a thorough removal of subarachnoid clot within 48 hours after SAH may help to prevent vasospasm (Nosko et al., 1982; Handa et al., 1987). At present however, there is no clinical evidence to suggest that such procedures
significantly lessen the risk of vasospasm or ischaemia in patients, and an extensive application of the procedure may indeed actually worsen acute cerebral swelling (Drake, 1981; Ohta et al., 1982).

Hydrocephalus

Some degree of asymptomatic ventricular enlargement is commonly seen following SAH, it is important however to recognise two groups of patients in whom either acute or delayed ventricular enlargement occurs, and which if left untreated by shunting, results in further neurological deterioration. It is worth noting here, that the intracranial pressure/volume relationship may be greatly altered acutely following SAH, and some patients do not tolerate well even small rises in ICP (Brock and Jane, 1980). An immediately beneficial effect upon the conscious level, of early CSF drainage in cases of acute hydrocephalus post-SAH, has now been well documented by a number of authors (Kusske et al., 1973; Suzuki et al., 1974; Hartmann et al., 1977; Hasan et al., 1989). The overall outlook in these patients, that present immediately following SAH with acute hydrocephalus, remains however, extremely unfavourable (Van Gijn et al., 1985; Doczi et al., 1983). A policy of reducing ICP by ventricular drainage before control of the source of the bleeding, for example prior to aneurysm clipping, appears to be associated with an increased risk of early rebleeding of around 40%, and an incidence of ventriculitis of approximately 50% after 3 days drainage (Brock and Jane, 1980; Hasan et al., 1989). In 10% of SAH patients, chronic hydrocephalus of the communicating type
may persist or develop after a delay. Unsurprisingly, in these patients, the results of shunting are usually good, with significant improvements both in their mental state and CBF following ventricular shunting procedures (Vasargil et al., 1973; Vassilouthis and Richardson, 1979; Mickey et al., 1984).

Medical treatment:

General measures

Patients when first seen are rapidly assessed as to their airway, and adequacy of ventilation and oxygenation ensured. If the patient is unable to maintain these parameters adequately, or if there are signs of increasing intracranial pressure with tentorial herniation, then intubation and hyperventilation is indicated. The blood pressure should initially be monitored continuously, and subsequently at frequent intervals. Arterial hypotension should be corrected vigorously as it may result in cerebral ischaemia in the presence of impaired autoregulation and raised intracranial pressure. The temptation to reduce the commonly found transient arterial hypertension should probably be resisted also for the same reasons, although some authors have advocated treatment to lower blood pressure if excessively high, to reduce the risk of rebleeding. Patients are placed upon strict bed-rest, stimuli minimised and the numbers of visitors restricted. In those patients in the better clinical grades, severe headache and neck pain will require adequate analgesia with codeine phosphate being given as necessary. Stool softeners should also be given regularly
so as to avoid the development of constipation, with resultant excessive straining. Poorer grade and incontinent patients will require the insertion of a closed-system continuous catheter drainage system, while a careful fluid-balance record should be kept in all cases. Prophylaxis of venous thromboembolism is performed by active and passive leg movement, together with the wearing of graded compression stockings.

Volume expansion, haemodilution, induced hypertension

Hypervolaemic haemodilution, combined in some instances with induced hypertension, has been widely recognised as a valuable adjunct in the prevention and treatment of delayed cerebral ischaemia arising following SAH. There is now good evidence available to show that SAH results in many patients in a contraction of the extracellular and vascular spaces, perhaps via a cerebral salt-wasting syndrome (Maroon and Nelson, 1979; Kudo et al., 1981; Pritz, 1984). Volume expansion is primarily aimed at correcting this contraction, thereby leading to an improvement in cardiac output and cerebral perfusion (Pritz et al., 1978; Finn et al., 1986). In addition, volume expansion using crystalloid or colloid agents can be employed to achieve a haemodilution, aiming at a haematocrit of 30-35%. Such haemodilution results in a lowering of blood viscosity (Stone et al., 1968; Wood and Kee, 1985), and leads to improved blood flow in the cerebral microcirculation (Sundt and Waltz, 1967; Grotta et al., 1982). The optimum haematocrit to maximize oxygen delivery to the cerebral tissue has been estimated at around 33% (Thomas, 1985; Wood and Kee, 1985). Kosnick and Hunt (1976)
were the first to report, that ischaemic neurological deficits that arose after intracranial aneurysm surgery could be successfully reversed by induced hypertension, raising the mean arterial pressure by some 20-40 mmHg above pretreatment values. Subsequent authors have confirmed the effectiveness of such therapies, and that approximately 60% of ischaemic deficits following SAH can be reversed using hypervolaemic haemodilution therapy combined with induced hypertension; aiming for a haematocrit of 33-38%, a central venous pressure of 10-12 mmHg (or a pulmonary wedge pressure of 15-18 mmHg) and a systolic arterial pressure of 160-200 mmHg (Symon, 1978; Kassell et al., 1982; Awad et al., 1986). An obvious risk of such therapy is rebleeding from an uncontrolled aneurysm, and most authors would reserve or modify this therapy in the non-operated patient. More recently, it has been reported that early treatment with fludrocortisone acetate may result in a reduction in the incidence of those patients that develop volume depletion after aneurysmal SAH (Wijdick et al., 1988).

Antifibrinolytic agents

Surgical intervention in patients that have recently suffered aneurysmal SAH, is primarily aimed at the prevention of rebleeding. It was natural therefore, that medical therapy with antifibrinolytic agents should have been introduced, in an attempt to achieve a similar aim. After initial favourable reports (Gibbs and O'Gorman, 1967; Mullan and Dawley, 1968), the widespread adoption of such agents by neurosurgeons worldwide soon followed. Mostly, two chemically related agents came into common usage - epsilon
aminocaproic acid (EACA) and tranexamic acid (TEA). Both act by similar mechanisms. Firstly by inhibiting the conversion of plasminogen to plasmin, and also by partially blocking the action of plasmin upon fibrin clot (Kassell et al., 1986). Early studies using these agents, appeared to confirm their clinical efficacy in the setting of aneurysmal SAH, with a reduction in rebleeding rates of up to fifty percent being reported (Nibbelink et al., 1975; Adams et al., 1981). Subsequent studies however, in which the overall mortality outcome was examined, have since shown that any reduction obtained in the mortality from rebleeding, is equally offset by a concommitant increase in cerebral ischaemia and infarction (Kassell et al., 1984b; Vermeulen et al., 1984). At present therefore, antifibrinolytic agents would appear to be of no overall benefit, and to have no place in the current management of patients following rupture of an intracranial aneurysm (Lindsay, 1987).

Autonomic blockade

A considerable body of evidence has demonstrated that following SAH a state of increased sympathetic outflow exists, with elevated levels of central and circulating catecholamines (Meyer et al., 1973; Neil-Dwyer et al., 1974; Benedict and Loach, 1978; Minegishi et al., 1987). Cardiac manifestations that can arise following SAH, include ventricular and supraventricular arrhythmias, as well as actual injury in the form of subendocardial necrosis (Marion et al., 1986). Attempts have been made to modify this response pharmacologically. A beneficial effect of
adrenergic blockade, especially beta-blockade with propanolol, being reported by some authors in the overall management of patients after aneurysmal SAH (Weider, 1974; Walter et al., 1982; Neil-Dwyer, 1986). Cardiac dysrhythmias as a cause of sudden death in such circumstances however, are probably most common immediately post-haemorrhage (Secher-Hansen, 1964; Estanol and Marin, 1975). Prophylaxis against these early events is clearly difficult, while the importance of later occurring events as a significant contribution to morbidity and mortality in patients attending hospital has not in fact been established (Marion et al., 1986).

**Prevention and treatment of cerebral vasospasm and ischaemia**

It remains unclear whether the delayed cerebral arterial narrowing with a partly resultant cerebral ischaemia, following SAH, represents an active contraction together with a failure of relaxation of cerebrovascular smooth muscle, or whether it instead results from a proliferative inflammatory vasculopathy of the vessel wall itself. Many attempts, using varied pharmacological agents, have been aimed at preventing or reversing such changes. The results of such therapies have been extensively reviewed previously (Wilkins, 1980; Wilkins, 1986). Wilkins concluding that "the numerous attempts made, ingenious though they have been, have largely been unsuccessful". Recently however, some encouraging results have been obtained in reducing the incidence of cerebral ischaemia after SAH, by early treatment with calcium channel blockers; particularly those of the dihydropyridine family. Calcium channel blockers
inhibit the influx of calcium through receptor- or voltage-operated channels in cell membranes. They have been widely used, both for prophylaxis against, and in the treatment of cardiac arrhythmias and angina (Triggle and Swamy, 1980; Braunwald, 1982). Dihydropyridines such as nimodipine readily bind to human brain membranes and selectively affect cerebrovascular smooth muscle to a greater extent than they do systemic arterial smooth muscle (Kazda and Towart, 1982; Peroutka and Allen, 1983). Their experimental administration can result in moderate cerebral vasodilation, which is most pronounced in the penetrating arterioles (Harper et al., 1981; Takayasu et al., 1988). However, they do not appear to reduce the frequency or severity of cerebral vasospasm following SAH (Espinosa et al., 1984a; Nosko et al., 1985).

The first multicentre controlled trial of nimodipine therapy in aneurysmal SAH was conducted by Allen et al (1983), and involved 121 patients. Cerebral arterial vasospasm was not reduced, nor was eventual recovery from ischaemic neurological deficits affected. Temporary neurological deficits did however occur with greater frequency in the placebo group. Two subsequent uncontrolled studies, and a further somewhat smaller controlled study, suggested that nimodipine therapy reduced both the incidence of delayed ischaemic dysfunction and permanent neurological deficits (Ljunggren et al., 1984a; Auer et al., 1986; Philippon et al., 1986). It was only in 1989 however, that the results of a substantial (540 patients), multicentre, double-blind, placebo-controlled trial on the efficacy of oral nimodipine therapy in this condition was published
This study showed a significant effect of treatment for all clinical grades of patients, both before and after operation, when started within 96 hours after the ictus. Oral nimodipine given to patients as 60mg four hourly, was shown to significantly reduce the incidence of cerebral infarction by 34%, and to reduce the incidence of poor outcome by 40%, as compared with placebo therapy.

The exact mechanism of action of these compounds in reducing cerebral ischaemia after SAH remains unclear. As previously indicated, dihydropyridines do not appear to actually prevent cerebral vasopasm and do not in themselves lead to consistent increases in cerebral blood flow (Neil-Dwyer, 1987; Schmidt et al., 1988; Young and Chien, 1989). In animal models of cerebral ischaemia however, nimodipine has been shown to reduce both infarct size and mortality, and to improve eventual neurological outcome (Harris et al., 1982; Kazda, 1983; Bederson et al., 1985; Steen et al., 1985; Gotoh et al., 1986b). Ischaemia leads to a rapid influx of calcium across neural membranes, and to an impaired mitochondrial ATP production that directly results in water accumulation within cells due to loss of ATP-dependent Na⁺/K⁺ membrane pump function (Siesjo, 1981; Hossman et al., 1983; Mohamed et al., 1985). Calcium channel antagonists such as nimodipine, by inhibiting neuronal calcium entry, may prevent or reduce secondary metabolic changes, thereby preserving neuronal ATP during the course of an ischaemic insult (Mabe et al., 1985; Kucharczyk et al., 1989)
1.3 CEREBRAL ARTERIES AND CONTROL OF THE CEREBRAL CIRCULATION

Anatomy:

The blood supply to the brain in man is predominantly derived from the two internal carotid and two vertebral arteries which lie within the subarachnoid space (Warwick and Williams, 1973). This simple arrangement is not uniform throughout the mammalian species, a point often overlooked in animal studies of experimental SAH. Primates and laboratory rodents, share with man, a virtual isolation of the intracranial and extracranial blood supply, with negligible anastomoses between them. In the dog and cat however, it is the external carotid artery that supplies the larger share of blood to the brain, via the rete mirabile a network of anastomoses that exists between extracerebral and intracerebral arterial branches (Kety, 1972).

The carotid and vertebrobasilar arteries communicate with each other at the base of the brain, through the cerebral arterial circle named after Thomas Willis (1621-1675), who published in 1664 a work based upon brain dissections he had performed, illustrated by Christopher Wren. The circle of Willis is regarded as providing a potential anastomotic circle, allowing collateral circulation in the event of interruption of one of the main input vessels (Fields et al., 1965).

The larger cerebral vessels branch to form successively smaller cerebral arteries, some also being given off directly. In contrast to some other regional circulations, the large arteries, including internal carotid and basilar
arteries, circle of Willis, and their direct branches, are important resistance vessels, accounting for almost half of total cerebral vascular resistance (Heistad et al., 1978; Abboud, 1981). The smaller arteries reach the neural surface in the pia mater, dividing to form muscular arterioles that together with the smallest arteries penetrate perpendicularly into the neural tissue (Warwick and Williams, 1973). The intracerebral arterioles within the neuropil are surrounded by a perivascular space, bounded by a basement membrane. Nerve fibers derived from the extracerebral perivascular plexus may penetrate as far as this space. As arteriolar calibre is reduced, the perivascular space narrows and is eventually obliterated by fusion of the vascular and parenchymal basement membranes. Smooth muscle cells are replaced by pericytes at this point, and the vessel becomes a true capillary (Jones, 1970; Rennels and Nelson, 1975).

Structure:

Cerebral arteries, arterioles and capillaries, conform in general terms to the same plan as that for vessels in the remainder of the body. Intracranial arteries are composed of the usual three layers - named from within outwards; the tunica intima, tunica media, and tunica adventitia (Dahl, 1973). The intima consists of a luminal lining composed of a single layer of endothelial cells, resting upon a fenestrated internal elastic lamina. These fenestrations have a mean diameter of 2 um, but at vessel bifurcations there are a smaller number of larger (7 um) fenestrations (Campbell and Roach, 1981). The tunica media is composed
of several layers of circularly arranged smooth muscle cells, that become fewer in number as the vessels decrease in size. In all vessels, membrane contacts between smooth muscle cells are found, these appear to allow to a limited extent the direct spread of electrical activity (Keatinge, 1978; Dahl, 1986). The media at cerebral arterial branchings commonly shows defects or discontinuities in the muscle layer (Forbus, 1930). Unlike the extracranial arteries, apart also from the ovarian artery, the cerebral arteries do not possess an external elastic lamina (Crompton, 1976). The adventitial layer is largely composed of collections of collagen bundles, between which fibroblasts are found. Within this adventitial layer, run bundles of nerve fibers forming a neural plexus that extends inwards to supply the surface of the smooth muscle media layer, in a manner typical of that observed for the autonomic innervation of vascular beds (Dahl and Nelson, 1964; Nelson et al., 1972; Burnstock, 1977). Vasa vasorum are largely absent (Scaravilli, 1985). Cerebral arterioles essentially resemble their parent arteries, but the media is now composed of only one to three layers of smooth muscle cells, and the adventitial layer is relatively thin. Cerebral capillaries are of the continuous variety, being composed of a non-fenestrated, complete tube of endothelial cells. Pericytes may partially surround this endothelium, which otherwise is separated only by its basement membrane from the surrounding neuropil (Warwick and Williams, 1973; Dahl, 1986). Pericytes have long been regarded by some workers, on the basis of their cytological features, to be a primitive
or undifferentiated form of vascular smooth muscle, and to subserve a role in capillary contractility (Krogh, 1929; Farquhar and Hartmann, 1956; Maynard et al., 1957). Similarly, on the basis both of cytological and biochemical evidence, it has been suggested that vascular endothelial cells possess a contractile capability (Rostgaard et al., 1972; Yohro and Burnstock, 1973). It is interesting in this context therefore, to note the observations made by a number of workers, of axon terminals directly apposed to and apparently innervating both pericytes and endothelial cells of CNS capillaries directly (Edvinsson et al., 1973; Swanson, 1977; Itakura et al, 1977; Itakura et al, 1985).

Cerebral vessels, especially the small arterioles and capillaries, present both physical and biochemical barriers to the passage of substances from blood to the brain. The physical barrier is formed by the presence of a continuous endothelium with its tight junctions, while the biochemical barrier is due to the presence of degradative enzymes that effectively prevent many substances from reaching the brain in any appreciable concentration (Rapoport, 1967).

Control of the cerebral circulation:

The normal CBF to grey matter is in the region of 60-70 mls/100g/min in the adult, and to white matter is approximately 25 mls/100g/min. This gives a mean hemisphere flow of around 50-60 mls/100g/min (Thomas and Crockard, 1985). Symon (1978), has detailed the effects upon the primate brain of a gradually reduced blood flow. At about 20 mls/100g/min, electrical activity ceases. At a CBF of around 12-20 mls/100g/min, membrane pump function fails, with
potassium losses from the cell together with sodium and water entry leading to cytotoxic oedema. Finally at a CBF of below 12 mls/100g/min, irreversible cellular damage is sustained.

In normal health however, the cerebral circulation displays an exquisite responsiveness, maintaining normal levels of flow despite fluctuations in systemic arterial pressure, regulating normal levels of oxygen and carbon dioxide within the brain tissues, and modulating flow locally to the demands of increased neuronal activity. Although a large number of mechanisms may influence the cerebral circulation, they must all ultimately act by producing change in one of two ways, either in the cerebral perfusion pressure - the difference between cerebral arterial and venous pressures, or the resistance to flow of blood through the cerebral vessels themselves.

Factors concerned with the regulation of the cerebral circulation can be considered conveniently under the following headings:

- Regulation to changes in systemic arterial pressure
- Regulation to changes in blood gases
- Regulation to changes in cerebral metabolism

In addition, it is useful to consider separately,

- Regulation by neural and vascular endothelial influences

since more is known regarding the actions of the various neurotransmitters upon the cerebral circulation, than is presently understood regarding their exact role in the above situations.
Regulation to changes in systemic arterial pressure

The cerebral circulation maintains the CBF at a remarkably constant level, over a wide range of systemic arterial pressures. This autoregulatory response is capable of keeping CBF close to normal, between mean arterial pressures of approximately 60-150 mmHg (Harper, 1966). Autoregulation also acts to maintain a constant CBF, when cerebral perfusion pressure (CPP) is similarly reduced by increases in CSF or cerebral venous pressure (Miller et al., 1972). These autoregulatory responses occur fairly rapidly, within 10-90 seconds (Symon, 1973). Autoregulation appears to be mediated by a vasodilation or vasoconstriction of the cerebral vessels, in response to increases or decreases in CPP respectively (Fog, 1938; Mchedlishvili, 1976). The exact mechanism underlying the responses of cerebral vessels in autoregulation remains unclear, proposed mechanisms have included myogenic, metabolic, neurogenic, and endothelial factors.

In initial studies, it was felt that regulation to pressure change must be a function of the vascular smooth muscle itself, hence the term autoregulation; for a full description of the myogenic hypothesis see Folkow (1964). There is no doubt, that in certain vascular beds that myogenic factors play a significant role. For example in the rat portal vein, stretch or relaxation appears to elicit sustained coordinated electrical and mechanical activity directly (Johansson and Mellander, 1975). In the isolated cat middle cerebral artery, a pressure-dependent reduction in internal diameter occurs, which is mediated by muscle
cell membrane depolarisation with action potential
generation, and which continues in the presence of neuronal
blockade (Harder, 1984; Harder and Lombard, 1985). The
subject of intrinsic myogenic responses in cerebral vessels
has been extensively reviewed previously (Orlov, 1979;
Mamisashivili, 1979), the evidence indicates that although a
short acting direct vessel response may occur, that it does
not satisfactorily wholly explain the mechanism of
autoregulation.

The metabolic theory, supposes that changes in perfusion
pressure may alter the local concentrations of substances
affecting vascular tone. Agents commonly proposed include
adenosine, K\(^+\), and H\(^+\). Adenosine is a well known potent
dilator of cerebral vessels (Berne et al., 1974). Its
concentration in the brain rises as a result of increasing
hypotension within the autoregulatory range (Winn et al.,
1980a; Bockman et al., 1981). Theophylline, a
methylxanthine that blocks the cerebrovascular effects of
adenosine, reduces cerebral vascular resistance at normal
perfusion pressure, but actually increases it at decreased
perfusion pressure (Oberdorster et al., 1975). Morii et al.
(1989) have shown that caffeine, another methylxanthine that
blocks adenosine receptors, attenuates the vasodilatory
response of cerebral vessels to induced hypotension, further
supporting a role for adenosine in CBF autoregulation.

Extracellular K\(^+\) and H\(^+\) concentrations have profound
effects upon cerebral arteriolar diameters, and as a result
blood flow (vide infra). It might be expected, that their
local concentrations would alter, depending upon
perfusion pressure. However, measurements of perivascular $K^+$ and $H^+$ concentration near pial vessels, when arterial pressure has been varied by 60-200 mmHg, have failed to demonstrate any change despite these vessels constricting or dilating in response to the induced pressure changes (Siesjo and Zwentnow, 1970; Wahl and Kuschinsky, 1979).

The existence of a neural role in autoregulation in man, is suggested by the observation that autoregulation of CBF is altered in patients with severe impairment of their autonomic system, such as occurs in Shy-Drager syndrome (Gotoh et al., 1972). Of the neurogenic influences, the most studied in this context have been that of the sympathetic nerves (Bill and Linder, 1976; Edvinsson et al., 1976a; Strandgaard et al., 1976; Busija et al., 1980; Heistad et al., 1982). From these studies, it now appears that although the sympathetic nerves are not of necessity required for the autoregulatory response of cerebral vessels, that they can however, exert a strong protective effect against increases in CBF during acute hypertension. In effect, shifting the upper limit of CBF autoregulation to higher levels. A converse effect has been noted following acute sympathectomy, when the lower limit of autoregulation is reduced by approximately 30 mmHg (Fitch et al., 1975).

More recently, a significant vascular endothelial contribution towards autoregulation has also been suggested, as it has been shown that contraction of isolated cerebral vessels to stretch or transmural pressure rise is effectively abolished by removal of the endothelial layer (Katusic et al., 1986; Harder 1987).
Regulation to changes in blood gases

CBF is profoundly affected by alterations in arterial pCO₂. Hypercapnia leading to a reduced cerebrovascular resistance, as a result of arterial dilation, with a consequently increased CBF. Conversely, hypocapnia has precisely the opposite effects (Raper et al., 1971). The cerebrovascular tone is also responsive to changes in arterial pO₂, but less so, than to similar changes in pCO₂. When PaO₂ falls below 40-50mm Hg, arterial dilation occurs and CBF rises (Borgstrom et al., 1975; Jones et al., 1981). Raising the PaO₂ at normal atmospheric pressure, has little effect upon CBF when PaCO₂ remains constant, although the use of hyperbaric oxygen will produce a fall in CBF (Lambertsen et al., 1953; Jones et al., 1981).

Cerebrovascular reactivity to arterial pCO₂ is not due simply to blood pH change, as alterations in arterial pH, provided the pCO₂ is kept constant, have little effect upon CBF (Harper and Bell, 1963). However, CO₂ is freely diffusible across the blood brain barrier, and after its passage will inevitably give rise to a change in brain extracellular pH. These alterations in extracellular H⁺ concentration have a profound effect upon cerebral arteries; decreased pH leading to dilation and augmenting CBF, while increased pH leads to constriction with a reduction in CBF (Betz and Heuser, 1967; Lassen, 1968; Kuschinsky et al., 1972). It is now generally considered that changes in CBF due to alterations in PaCO₂, are largely mediated via changes in brain extracellular pH (Skinhoj, 1966; Kontos et al., 1972; Kontos et al., 1977). The concept however, of a
brainstem center that additionally modulates the cerebrovascular response to PaCO$_2$ change, is suggested by studies utilising acute brain sectioning. These have demonstrated the importance of the brainstem at the level of the Pons in moderating this PaCO$_2$ response (Shalit et al., 1967; Capon, 1975). More recently, it has shown that stereotactic lesioning of the locus coeruleus, a nucleus within this region that diffusely projects central noradrenergic fibers to the cerebrum, results in a significant reduction in the slope of the CBF - PaCO$_2$ response curve (Reddy et al., 1986).

Hypoxia appears to be capable of directly affecting the responses of cerebrovascular smooth muscle. Isolated cerebral vessels exposed to hypoxia show inhibition both of contractility to transmural pressure change, and vasoactivity to applied agents such as serotonin (Lombard et al., 1986; Vinall and Simeone, 1986). This impairment of reactivity, appears to reflect a reduced calcium uptake into the smooth muscle cell under hypoxic conditions (Ebeigbe et al., 1980; Vinall and Simeone, 1986). It has also been proposed that the rise in CBF induced by hypoxaemia is due to an accumulation of lactic acid and resultant acidosis. However, in hypoxia studies the CBF increase is clearly initiated before lactate rises, and while pH is unchanged (Nordberg and Siesjo, 1975; Astrup et al., 1979; Javaheri, 1986). As hypoxia continues however, tissue acidosis develops due to lactic acid accumulation, and this may then significantly contribute to the maintenance of vasodilation (Astrup et al., 1979). A contribution by K$^+$ to the
initiation of hypoxic hyperaemia, is suggested by the finding of a moderate increase in extracellular brain $K^+$ shortly after the onset of hypoxia (Krishner et al., 1975; Astrup et al., 1976). It has also been proposed that adenosine may play a role in the hypoxic response. During hypoxia, adenosine is known to accumulate as a result of the dephosphorylation of adenosine monophosphate (AMP) by 5-nucleotidase (Rubio et al., 1975). Brain tissue and interstitial adenosine concentrations rise up to sixfold, within seconds of hypoxia (Winn et al., 1981; Zetterstrom and Fredholm, 1982). During shorter periods of hypoxia, of up to 30 seconds, these temporal changes in brain tissue adenosine are inversely correlated with the changes in cerebrovascular resistance (Winn et al., 1984). Further support for the role of adenosine, is given by the finding that administration of the adenosine inhibitor theophylline attenuates both the vasodilation and hyperaemic response produced by hypoxia (Emerson and Raymond, 1981; Morii et al., 1987).

Regulation to changes in metabolisms

The concept that cerebral blood flow can be adjusted either to or by neuronal activity, was elegantly hypothesised towards the end of the last century by Roy and Sherrington (1890) who wrote that "The chemical products of cerebral metabolism contained in the lymph that bathes the walls of the arterioles of the brain can cause variations of the calibre of the cerebral vessels. In this reaction the brain possesses an intrinsic mechanism by which its vascular supply can be varied locally in correspondence with local
variations in functional activity." The expectations of Roy and Sherrington that increased neuronal activity was accompanied by increased metabolism and that an increased CBF would follow, has been amply confirmed in numerous experimental studies (Sokoloff, 1961; Olesen, 1971; Greenberg et al., 1979). However, the precise mechanism by which neuronal function and cerebral perfusion are so closely and intimately coupled, remains less certain (Sokoloff, 1981).

Suggested factors for this coupling have included changes in extracellular pH, PO$_2$, CO$_2$, K$^+$, and adenosine. Using the activation of neurons by electrical stimulation, it has been demonstrated that local blood flow increases within 1-2 seconds of enhanced activity, and that the zone of increased CBF does not extend beyond approximately 250 um from the site of increased activity (Moskalenko, 1975; Silver, 1978). Following local neuronal stimulation, hyperaemia is initiated in spite of an initial fall in extracellular H$^+$ concentration and a rise in PO$_2$ (Lubbers and Leniger-Follert, 1978; Silver, 1978). Both these latter authors demonstrated that an immediate rise in local extracellular K$^+$ concentration, of the order of 0.5-5 Meq/L, occurs with such enhanced neuronal activity. The extracellular K$^+$ has similarly been shown, to rise at the onset of bicuculline-induced seizures (Astrup, 1978). Extracellular K$^+$ acts as a potent vasodilator and promotor of increased CBF (Kuschinsky et al., 1972; Cameron and Caronna, 1976). Increases in extracellular K$^+$ in the range demonstrated to occur with increased neuronal activity, will
cause small cerebral arterioles to dilate, by an increase in conductance of K⁺ channels and resultant hyperpolarisation of smooth muscle cells (Silverberg, 1989).

In neuronal coupling, as in hypoxia and hypotension, again it has been proposed that adenosine may be partly involved in mediating vasodilation. Both electrical stimulation and bicuculline-induced seizures have been shown to produce an increase in brain adenosine concentration (Pull and McIlwain, 1972; Wahl et al., 1978; Winn et al., 1980b). While extracellular K⁺ concentrations increase with the activation of neuronal metabolism, extracellular Ca²⁺ concentrations show a decrease. These changes in Ca²⁺ have been observed both with somatosensory cortex activation by peripheral field stimuli (Heinemann et al., 1977) and bicuculline-induced seizures (Heuser, 1978). The reductions in Ca²⁺ reported from these experiments were however, less than 1mM, which suggests they form only a minor contribution towards the vasodilatory response (Betz et al., 1973).

Recently, attention has become increasingly focused on the role of the neural plexus that surrounds cerebral vessels and microvessels, both in terms of local coupling of neuronal function to blood flow, and in causing a more widespread activation of cerebral metabolism and increased blood flow (vide infra).

Regulation by neural and vascular endothelial influences

Neural effects

The cerebrovascular bed is known to be innervated by a quite extensive system of perivascular nerve fibers, that utilises a wide variety of vasoconstrictor and vasodilator
transmitters. The cerebral vasculature would appear to be unique, in that it receives nerve fibers from both the peripheral autonomic system and central neuronal sources directly. The larger cerebral vessels of the circle of Willis, small arteries and penetrating arterioles, are innervated by nerve fibers that arise from the cervical and cranial ganglia. As these vessels successively branch and decrease in diameter, so there is a concommitant reduction in the number of perivascular nerve fibers that accompany them. There are also, significant variations in the density of innervation received by vessels, in different areas of the brain. Approximately 75% of the arterioles of the thalamus, hypothalamus and parietal cortex are innervated, while in the medulla, cerebellum and occipital cortex, only some 25% of vessels receive any identifiable innervation (Edvinsson and Owman, 1977). The intracerebral capillaries in contrast, would appear to receive a completely separate more limited innervation by nerve fibers that arise from the ascending central aminergic pathways of the brainstem. In addition, both capillary endothelial cells and pericytes possess contractile proteins (Owman et al., 1977), and may therefore be capable of reacting directly to the release of neurotransmitters from nearby intracerebral neurons.

NORADRENALINE

Cerebral vessels both in humans and in laboratory animals receive a rich innervation of noradrenergic nerve fibers. The arteries of the anterior circulation are largely supplied from the ipsilateral superior cervical ganglia (Nielsen and Owman, 1967; Edvinsson et al., 1972; Edvinsson et al.,
The arteries of the posterior circulation, receive in addition a contribution from the lower cervical ganglia, and tend to have a greater overlap of innervation between the two sides (Edvinsson, 1975). Detailed immunoelectron microscopic examination both of the hypothalamus and cerebral cortex, has demonstrated that noradrenergic nerve fibers are also found in close association with intracerebral capillaries, in some cases with no glial process between them (Swanson et al., 1977; Papadopoulos et al., 1989). These latter nerve fibers originate centrally, the majority arising from neurons in the locus coeruleus (Fuxe et al., 1968).

Cerebral arteries are however, relatively insensitive to noradrenaline as compared with other vessels (Duckles and Bevan, 1976). Studies though, have indicated the presence on human cerebral vessels of both \( \alpha_1 \)- and \( \alpha_2 \)-adrenoceptors (Toda, 1983; Tsukahara et al., 1985). Alpha-adrenoceptor function seeming to predominate in the large cerebral arteries, and causing constriction. However, \( \beta_2 \)-adrenoceptors appear to predominate in smaller vessels where their activation produces a vasodilation (Edvinsson et al., 1979; Harik et al., 1980; Nathanson, 1980). The noradrenergic sympathetics appear predominantly to play a role in acute hypertension and autoregulation, whilst the central noradrenergic projection of the locus coeruleus may be involved in modulating the hypercapnic response (Vida supra). The role of noradrenergic nerves in the regulation of resting CBF has been more disputed (Heistad and Marcus, 1978; Purves 1978). Overall findings, and more recent
investigations, do lend support to the concept that peripheral noradrenergic sympathetic stimulation can reduce cerebral perfusion, larger pial vessels constricting whilst smaller intraparenchymal vessels dilate (Edvinsson and MacKenzie, 1977; Gotoh et al., 1986a). The effects upon the resting CBF, of lesions made within the locus coeruleus have been largely inconclusive (Bates et al., 1977; Dahlgren et al., 1981a), stimulation however of this nucleus has been reported to result in a global reduction in CBF (Raichle et al., 1975).

NEUROPEPTIDE Y

Neuropeptide Y (NPY) coexists with noradrenaline in the great majority of the sympathetic nerves supplying cerebral blood vessels (Edvinsson et al., 1983; Matsuyama et al., 1985). In addition more recently, it has also been shown to coexist in a smaller proportion of the cerebrovascular parasympathetic cholinergic/VIPergic nerves (Cavanagh et al., 1989; Suzuki et al., 1989a). NPY causes potent concentration-dependent contractions of cerebral vessels in man and animals, and does not appear to require an intact endothelium for its action (Edvinsson et al., 1983a; Allen et al., 1984; Mejia et al., 1988). The primary action of NPY upon cerebrovascular smooth muscle would appear to be mediated by a direct stimulation of specific receptors. This is quite unlike the situation in peripheral arteries, where NPY does not cause a direct contraction per-se, but instead potentiates the contractions induced by electrical or aminergic nerve stimulation (Edvinsson, 1985; Edvinsson et al., 1987).
The cerebral circulation is endowed with a cholinergic, and vasoactive intestinal peptide (VIP)ergic innervation that is more diffuse in its origin than that of the noradrenergic innervation. Cholinergic / VIPergic nerves arise both from cranial microganglia, and cells in the otic and sphenopalatine ganglia that project via the greater petrosal branch of VII to the carotid plexus (Chorobiski and Penfield, 1932; Edvinsson et al., 1980; Gibbins et al., 1984; Hara et al., 1985; Saito et al., 1985). A proportion of such nerves and their cell bodies will be either purely cholinergic or VIPergic alone, whilst in others the two neurotransmitters coexist (Suzuki et al., 1989). The density of the innervation by cholinergic and VIPergic nerves appears to be considerably greater in the anterior, as compared with the posterior cerebral circulation (Gibbins et al., 1984; Saito et al., 1985). Stimulation of the greater petrosal / VII pathway or sphenopalatine ganglia directly, produces increases in CBF that occur quite independently of changes in cerebral metabolic activity (D'Alecy and Rose, 1977; Goadsby, 1989; Seylaz et al., 1989). Petrosal denervation studies however, have failed to show any involvement in either hypercapnic or hypoxic responses (Hoff et al., 1977; Busija and Heistad, 1981). Hence the physiological role of these nerves remains unclear at present.

In vitro studies show that cerebral vessels possess cholinergic receptors of the muscarinic type. Application of acetylcholine in low concentrations to isolated vessels
produces vasodilation, while higher concentrations result in vasoconstriction (Edvinsson et al., 1977a; Lee et al., 1980). The dilatory response produced by cholomimetic stimulation, is attenuated following removal of the vascular endothelium. Experiments indicate that in low concentrations, acetylcholine causes vasodilation by inducing the release of an endothelium derived relaxing factor – EDRF (Furchgott and Zawadzski, 1980; Lee, 1982). There is also now evidence for intrinsic cholinergic neuronal activation of the cortical microvasculature. Acetylcholine released by cortical cholinergic neurons as a result of stimulation of the fastigial nucleus of the cerebellum, being capable of effecting a regional increase in CBF without concommitant changes in glucose utilization (Nakai et al., 1983; Iadecola et al., 1986; Arneric et al., 1987).

In vitro, VIP dilates both small and large cerebral arteries, even at quite low concentrations. This response is blocked by indomethacin, a cyclo-oxygenase inhibitor, indicating that it may in part be mediated via local prostaglandin synthesis (Wei et al., 1980a; Edvinsson et al., 1981). It is of some interest therefore, that indomethacin administered in a number of species including humans can produce cerebral vasoconstriction and reduce CBF without affecting cerebral metabolism (Vide infra). Many VIP-containing neurons have now been demonstrated to exist diffusely in the cerebral cortex (Fuxe et al., 1977). The possibility of a direct action of VIP, released from such cells, upon the cerebral microcirculation is lent weight by
the electron microscopic observation of close associations existing between some intracortical blood vessels with nearby VIPergic terminals and neurons (Eckenstein and Baughman, 1984).

TACHYKININS/CGRP

The cerebrovasculature receives a nociceptive/vasomotor sensory innervation, that largely originates from cell bodies within the trigeminal ganglion (Mayberg et al., 1981; Ruskell and Simons, 1987; Suzuki et al., 1989b). Tachykinins are a family of small peptides that produce cerebral vasodilation. The tachykinins; substance P (SP), neurokinin A (NKA) and neurokinin B (NKB), coexist together with another peptide - calcitonin gene related peptide (CGRP), within these sensory nerve fibers and their parent cell bodies. (Liu-Chen et al., 1983; Uddman et al., 1985; Saito et al., 1987). Few studies involving either trigeminal stimulation or denervation have been performed. One study however, has shown that trigeminal ganglionectomy, but not rhizotomy, attenuates the increase in CBF induced by severe hypertension by 25-30% (Sakas et al., 1981). CGRP and the tachykinins, produce relaxation of precontracted isolated cerebral arteries in the following order of potency: CGRP > SP > NKA. The relaxation induced by the tachykinins are all endothelial-dependent, while that of CGRP is not (Hanko, 1985; Edvinsson and Jansen, 1987; Mejia et al., 1988).

SEROTONIN

The widespread presence of serotonin (5-HT) within the brain was first demonstrated, by biochemical means, over
thirty years ago (Twarog and Page, 1953). With the development of immunohistochemical staining methods employing specific antibodies raised against 5-HT, it has been made possible to visualise the origin and distribution pattern of this neurotransmitter directly (Steinbusch, 1978). The cerebral structures in most species studied, receive a rich monosynaptic innervation that arises from cell bodies located within the dorsal and medial raphe nuclei of the midbrain (Steinbusch, 1981; Lindvall and Bjorklund, 1984; Morrison and Foote, 1986). Detailed immunohistochemical studies upon the raphe nuclei, have demonstrated the occurrence of intimate contacts between serotonergic neurons and brainstem microvessels, for which a vasosensorimotor role was postulated (DiCarlo, 1977; DiCarlo, 1984; Kapadia and DeLanerolle, 1984). Biochemical and immunoelectron microscopic studies have also furnished evidence for a more widespread innervation of the cerebral microcirculation by axonal projections from these same nuclei (Reinhard et al., 1979; Itakura et al., 1985; Scatton et al., 1985). More recently, evidence for the existence of a serotonergic innervation of the major cerebral vessels of various species, apparently arising peripherally from the superior cervical ganglia, has been reported (Griffith et al., 1982; Griffith and Burnstock, 1983; Alafaci et al., 1986; Cowen et al., 1987). Although this finding has not been supported by others (Itakura et al., 1985). Stimulation of the raphe nuclei, in intact conscious animals, causes an increase in thalamic and cortical neuronal glucose utilisation, and produces heterogeneous increases in blood
flow (Cudennec et al., 1987; Cudennec et al., 1989). As with noradrenaline, serotonin has both vasoconstrictor and vasodilator effects upon the cerebral vessels. The direct effects of 5-HT has been studied both upon isolated vessels in vitro, and by its microapplication to pial vessels in vivo. Vessels with a resting diameter of 200 um or more constrict, whilst smaller arterioles of less than 70 um dilate (Edvinsson et al., 1977b; MacKenzie et al., 1978). The 5-HT induced vasoconstriction is largely mediated by the 5-HT_{2} receptor, and can be inhibited by the specific antagonist ketanserin (Edvinsson et al., 1978; Chang and Owman, 1986). This vasoconstriction is highly dependent upon extracellular calcium, and is more sensitive to calcium antagonists than is the case for peripheral vessels (Allen and Banghart, 1979; Shimizu et al., 1980; Towart, 1981). The vasodilator action of 5-HT upon small arterioles is also dependent upon the mean arterial pressure. The higher the vascular tone, the greater the dilator response to applied 5-HT (Harper and MacKenzie, 1977a). This dilator response is entirely unaffected by the classic 5-HT antagonists, but is however inhibited by propanolol, which suggests that the response is in some way mediated by an interaction with beta-adrenoceptors (Edvinsson et al., 1977b).

ADENOSINE NUCLEOSIDE AND NUCLEOTIDES

These include; adenosine, adenosine 5'-monophosphate (AMP), adenosine 5'-diphosphate (ADP), and adenosine 5'- triphosphate (ATP). The potent, and widespread effects of the adenine compounds upon the vasculature have long been recognised (Drury and Szent-Gyorgi, 1929). Their actions
upon vascular smooth muscle are mediated by specific purinoceptors, that can be subclassified, partly on the relative potency of the adenine compounds (Burnstock, 1978; Van Calker et al., 1979; Londos, 1980; Burnstock, 1987). Purinoceptors can be divided into $P_1$- and $P_2$- purinoceptors. $P_1$-purinoceptors can also be subclassified into $A_1$ (or $R_1$) and $A_2$ (or $R_2$) receptors, whilst $P_2$-purinoceptors may be further subclassified as $P_{2x}$- and $P_{2y}$- purinoceptors. The vasodilatory action of adenosine and AMP upon cerebral vessels is largely mediated by the $A_2$ subtype of the $P_1$-purinoceptor. Its activation leads to adenylate cyclase stimulation, and can be antagonised by the methylxanthines (Edvinsson and Fredholm, 1983; McBean et al., 1988). This adenosine induced relaxation is endothelium independent (Hardebo et al., 1985). ATP and ADP act predominantly via the $P_2$-purinoceptor, the occupation of which, in some cases, leads to increased prostaglandin synthesis (Burnstock, 1987). In cerebral vessels, ATP exerts effects both by activation of the $P_{2x}$-purinoceptor causing vasoconstriction and via the $P_{2y}$-purinoceptor to produce vasodilation, the latter effect probably being mediated via inhibitory endothelial receptors (Burnstock and Kennedy, 1986; Alborch et al., 1989). Adenosine and its nucleotides share a ubiquitous distribution in all tissues, and are essential for a variety of biochemical and metabolic processes (Bowman and Rand, 1980). Local neuronal release may play an important role in the cerebrovascular responses to changes in perfusion pressure, hypoxia and cerebral metabolism (vide supra). In addition, there is now considerable biochemical
evidence that ATP is stored as a cotransmitter with noradrenaline in sympathetic neurons and noradrenergic nerves (Stjarne and Lishajko, 1966; Geffen and Livett, 1971; Medrum and Burnstock, 1983; Sneddon and Westfall, 1984) including the sympathetic nerves supplying cerebral vessels (Muramatsu et al., 1981).

Endothelial effects

The intimal endothelial layer, lying directly interposed between circulating blood and vascular smooth muscle, is strategically placed to influence vessel responses. Endothelial cells take up and metabolise vasoactive substances such as noradrenaline, serotonin and substance P. In response to neurohumoral mediators and other stimuli, they are capable of releasing both endothelial-derived relaxant and constrictor substances. In addition, endothelial cells can produce and secrete vasoactive prostaglandins, most notably prostacyclin.

ENDOTHELIUM-DEPENDENT RELAXING FACTORS

The necessity for an intact endothelium in the cholinergic vasodilator response, and the existence of a potent relaxing agent released by vascular endothelial cells in response to acetylcholine, and later termed endothelium-derived relaxing factor (EDRF), was first reported by Furchgott and Zawadski (1980). Subsequently, the presence of an intact endothelium has also been shown to be required for the vasodilation elicited in cerebral vessels, not only by acetylcholine but in addition by ATP, tachykinins and bradykinin (Verrecchia et al., 1986;
Edvinsson and Jansen, 1987; Alborch et al., 1989). EDRF, probably only one of a number of relaxant substances released by the vascular endothelium, is a very labile diffusible substance, readily degraded by superoxide anions. It has recently been shown to be identical with nitric oxide (NO) (Palmer et al., 1987). EDRF appears to act by stimulating the soluble guanylate cyclase of vascular smooth muscle cytosol. This leads to increases in cGMP levels, and relaxation of the vascular smooth muscle (Holzmann, 1982; Furchgott, 1983).

ENDOTHELIUM-DEPENDENT CONTRACTING FACTORS

The development in isolated cerebral arteries of contractions to either anoxia or changes in transmural pressure, are dependent upon the presence of an intact endothelial layer, and can therefore be explained on the basis of the release of some endothelium-derived contracting factor(s) (Katusic and Vanhoutte, 1986; Harder, 1987). It is quite likely that more than one factor is involved in this endothelium-dependent constriction. One such factor is endothelin, a recently identified, 21 amino acid vasoconstrictor peptide produced by vascular endothelial cells (Yanagisawa et al., 1988). Endothelin appears to be one of the most vasoconstrictive of substances yet known, and endothelin-induced constriction is both long lasting and extremely resistant to reversal.

Endothelin-like immunoreactivity is present in the endothelium of cerebral vessels. Studies show that the application of endothelin to cerebral vessels both in vivo and in vitro produces a strong sustained vasoconstriction,
that is at least partly calcium-dependent (Edvinsson et al., 1989; Mima et al., 1989; Robinson et al., 1989; Saito et al., 1989).

ENDOTHELIAL-DERIVED EICOSANOIDS

The endothelial cells of the cerebrovasculature, are known to be capable of synthesising and releasing a variety of arachidonic acid derivatives. These include; prostaglandin E\textsubscript{2}, prostaglandin F\textsubscript{2}, and the vasodilator prostacyclin (Boullin et al., 1981; Pickard and Walker, 1984). Information on the physiological effects of these endogenously synthesised eicosanoids, has largely been inferred from studies that attempt to impair their production using the cyclo-oxygenase inhibitor indomethacin. Works by numerous investigators have demonstrated that the inhibition of cyclo-oxygenase activity by indomethacin, induces cerebral vasoconstriction, reducing CBF in man and a variety of animals by 30-50\% during normocapnia, without similarly affecting metabolism (Bill, 1979; Pickard, 1981; Dahlgren et al., 1981b; Crockard et al., 1982; McCulloch et al., 1982; Quintana et al., 1983; Pickles et al., 1984).

More controversial, has been a reported role of prostaglandins in increasing CBF during hypercapnia. Of the previously mentioned workers, some have also found that indomethacin reduces the rise in CBF elicited by hypercapnia. In contrast, others (Wei et al., 1980b; Busija and Heistad, 1983; Jackson et al., 1983) have been unable to demonstrate any effect of indomethacin upon the hypercapnic response.
1.4 AIMS OF THE STUDY

These were to examine the effects of subarachnoid haemorrhage upon various neuropharmacological, physiological, and morphological aspects of the major cerebral arteries. It was hoped that a productive investigations of changes in the above parameters, of possible relevance to man, could be obtained using a small animal model of experimental SAH developed in the laboratory rat. In particular, it was considered of potential benefit towards achieving a greater understanding of the pathophysiology of the condition to explore the following avenues of research interest.

(1) To elucidate the normal relationship of cerebrovascular 5-HT containing nerve fibers with other nerve fibers, particularly catecholaminergic sympathetics, within the major cerebral arteries.

(2) To characterise and compare any changes that might arise following SAH in the levels of vasoconstrictor neurotransmitters such as 5-HT and NPY contained within these perivascular nerve fibers.

(3) To determine the precise time-course of any alterations in CBF following experimental SAH in the rat, and to attempt to correlate this against any alterations in CPP, ICP, or other changes that might develop over the same time period.

(4) To undertake scanning and transmission electron microscopic studies upon the major cerebral arteries at different time points following experimental SAH, to
establish if changes in ultrastructural morphology similar to those that have been associated with vasospasm in man and larger animals, also develop after SAH in the rat.

(5) To ascertain the mechanism, by which subarachnoid blood clot is so effectively and rapidly cleared from the perivascular and other CSF spaces, following experimental SAH in this animal model.
2. GENERAL METHODS

Details of the experimental methods and materials that were utilised to obtain the data presented and analysed in this thesis, are provided in each of the individual experimental chapters that follow. Certain general principles, and points of specific note with regards to some of the techniques employed are first briefly considered.

2.1 IMMUNOFLUORESCENCE AND IMMUNOELECTRON MICROSCOPY

The two-layer, indirect immunofluorescence technique, as described by Coons et al. (1955), was used for the demonstration of immunoreactivity to the neurotransmitters 5-HT and neuropeptide Y at the light microscopic level. By this method, a primary antibody raised against the substance being investigated, is first reacted with the tissue under study. A second-layer antibody, directed against the gamma-globulin of the species in which the first antibody was raised, is then applied. This second-layer antibody, to which a fluorescent label such as fluorescein isothiocyanate (FITC) has firstly been preconjugated, binds to the first antibody, allowing visualisation of the primary antibody-antigen complex location when viewed under an ultraviolet microscope.

Localisation of immunoreactivity for 5-HT at the ultrastructural level, by transmission electron microscopy, was performed by a three-layer peroxidase anti-peroxidase (PAP) technique (Sternberger, 1979). Visualisation of the resulting antibody-antigen complex was achieved using Graham and Karnovsky's (1966) diaminobenzidine (DAB) reaction for
peroxidase. The final reaction product obtained, being made electron-dense following treatment with osmium tetroxide.

2.2 HIGH PERFORMANCE LIQUID CHROMATOGRAPHY

The concentrations of 5-HT and noradrenaline in pooled samples of cerebral vessels, were determined by HPLC with electrochemical detection, using the method of Reinhard et al. (1980). Tissue samples were homogenised in 0.1 M perchloric acid using a motor-driven glass-glass homogeniser, cell debris being removed by low speed centrifugation with filtration of the resultant supernatant. Aliquots were then injected through a reverse phase column (C-18 microbondapak, Radial Pak, Waters, UK); compounds being separated out on the basis of their overall polarity. Subsequent detection and quantitations were performed with a glassy electrode set at a potential of 0.72V (Roth Metrohm detector, Roth Scientific). Supernatant aliquots used for the determination of noradrenaline concentration were firstly treated by alumina extraction (Keller et al., 1976) followed by the addition of sodium heptane sulphonate, then reverse phase ion-pair separation, by passage through the column. Noradrenaline levels were corrected for alumina recovery losses using dihydroxybenzylamine (DHBA) as an internal standard.

2.3 HYDROGEN-CLEARANCE MEASUREMENT OF LOCAL CEREBRAL BLOOD FLOW

The measurement of local blood flow, using implanted platinium electrodes to record tissue hydrogen concentrations following the inhalation of hydrogen gas, was
first described by Aukland et al. (1964). The necessary calculations involved being derived from Kety and Schmidt's (1948) interpretation of the Fick principle (Fick, 1870). Kety and Schmidt recognised that "the amount of an inert gas taken up by tissue per unit of time is equal to the quantity brought to the tissue by the arterial blood minus the quantity carried away in the venous blood".

\[ WdCi = (Ca - Cvi) Fdt \]

Where \( Ci \) is the concentration in the tissue with volume \( W \), \( Ca \) is arterial concentration, \( Cvi \) the concentration in venous blood from that particular tissue, and \( F \) is the blood flow.

The theoretical principles and validation of the hydrogen clearance method have been fully described (Auckland et al., 1964), and its applicability to the measurement of local cerebral blood flow extensively reviewed (Farrar, 1987). Further technical consideration and an assessment of the suitability of the technique for the obtaining of cerebral blood flow measurements chronically in small animals, forms the subject of experimental chapter IV of this thesis. The basic principle utilised, is that a platinum-wire electrode is inserted into the cerebral tissues under study, then a positive potential created between that electrode and a remote reference electrode. Hydrogen gas administered by inhalation, results in the formation of an electrode current generated by the oxidation of molecular hydrogen to hydrogen ions upon the platinum surface. Electrode current will then be a linear function of tissue hydrogen concentration (\( Ci \)), provided other parameters remain stable. Standard clearance
Curves for hydrogen can therefore be readily obtained, by recording the resultant electrode current over time elapsed since hydrogen inhalation was discontinued. The slope of the resultant tissue desaturation clearance curve when plotted on a semilogarithmic scale against time, permits the obtaining of flows in cc/min/cc of tissue.

Since: \[ F = \frac{0.693}{T/2} \] (2)

Where \( T/2 \) is the time in min for tissue hydrogen concentration and the resultant current to be reduced to half of its numerical value.

2.4 Acid-Etch Scanning Electron Microscopy

The general principles, and practical application of both transmission and scanning electron microscopy have been so extensively described as to need no further introduction here. However, to allow the three-dimensional organisation of the perivascular schwann-cell / neural plexus within the adventitia of major cerebral vessels to be visualised, before and after SAH, it was necessary to use an acid-etch microdissection technique. This method, used as a preliminary step in the preparation of specimens prior to scanning electron microscopy, was originally described by Evan et al. (1976) for the study of cell surfaces. It has subsequently been adapted by Uehara and his coworkers for the study of vascular nerve plexuses (Uehara et al., 1981). In brief, specimens after initial perfusion-fixation then post-fixation in aqueous osmium tetroxide and washings in distilled water, are then further treated by exposure to 8N HCl for 10-30 min at 60°C prior to further washing,
dehydrations and processing through to critical point
drying. The addition of the 8N HCl treatment step results
in a partial hydrolysis of the connective tissues. This
permits vessels to be mechanically fractured at their
adventitial medial border, thereby fully exposing the
perivascular neural plexus, prior to final sputter coating
with gold for viewing by scanning electron microscopy.
3. EXPERIMENTAL CHAPTER I:

ULTRASTRUCTURE OF SEROTONIN-CONTAINING NERVE FIBRES IN THE MIDDLE CEREBRAL ARTERY OF THE RAT AND EVIDENCE FOR ITS LOCALISATION WITHIN CATECHOLAMINE-CONTAINING NERVE FIBRES BY IMMUNOELECTRON MICROSCOPY
Summary: This study has examined the ultrastructural characteristics of serotonin (5-HT)-like immunoreactive (5-HT-I) nerves in the perivascular plexus of a major cerebral blood vessel and their relationship with catecholaminergic (CA) nerves. This was achieved by immunohistochemistry for 5-HT used alone and in combination with simultaneous 5-hydroxydopamine (5-OHDA) false transmitter preloading to demonstrate catecholaminergic nerves examined by electron microscopy. In immersion-fixed vessels about 15% of perivascular nerves showed some 5-HT-like immunoreactivity. These nerves were characterised by a predominance of small pleomorphic vesicles with a diameter of 40-50 nm and a few large granular vesicles about 100-150 nm diameter. 5-HT-I nerves were found at all levels of the perivascular plexus, from the outer adventitia where they were formed together with non-reactive nerves into bundles by Schwann cells and their processes, to more singly at the adventitia-media border, sometimes in close apposition to smooth muscle cells, separated by some 125 nm. Examination of animals treated by 5-OHDA preloading revealed that these 5-HT-I nerves were almost invariably identified as catecholaminergic. However, CA nerves did not always show 5-HT-like immunoreactivity, some 25% only, being doubly labelled in this study. Treatment of animals with the catecholaminergic nerve uptake blocker Desmethyli mipramine (DMI) resulted in abolition of 5-HT-like immunoreactivity. These results provide direct ultrastructural evidence for the localisation of 5-HT within the perivascular catecholaminergic nerves of a major cerebral vessel and
suggest that the contained 5-HT is mainly derived by uptake rather than synthesis within them.
3.1 INTRODUCTION

The presence of serotonin as a putative neurotransmitter in cerebral perivascular nerves was first suggested by autofluorescence studies in a lower vertebrate, the lamprey (Baumgarten, 1972). Subsequent autoradiographic and autofluorescence studies in the rat and monkey supported its presence in the plexus about major cerebral vessels (Chan-Palay, 1976) and the microvasculature of the brainstem raphe region (Di Carlo, 1977). Work followed that suggested a serotonergic innervation of the small intraparenchymal vessels of the rat forebrain (Reinhard et al., 1978; Reinhard et al., 1979) and studies showed depletion of 5-HT from small pial and intraparenchymal vessels after lesions of the raphe nuclei (Reinhard et al., 1979; Mackenzie and Edvinsson, 1980; Edvinsson et al., 1983b). The first immunohistochemical evidence for its existence in a major cerebral vessel was presented in a light microscopic study of the rabbit vertebral artery (Griffith et al., 1982) and in addition, some of its pharmacological properties characterised. More recently, 5-HT-like immunoreactive nerves have been demonstrated in the major cerebral arteries of the gerbil and have been shown to share a common origin, with noradrenergic nerves, from the superior cervical ganglion (Alafaci et al., 1986; Cowen et al., 1986), although it was not possible to say if there were two distinct populations of nerves or the two substances co-existed within the same nerves and their parent neurones.

A complex pharmacological interrelationship exists
between 5-HT and noradrenaline in the vasoconstrictor response elicited by each of these substances released upon stimulation of the perivascular nerves (Marin et al., 1981; Morotoki and Su, 1981; Kawasaki and Takasaki, 1984). In addition it has been demonstrated that 5-HT may be taken up and accumulated in catecholaminergic nerves of peripheral and cerebral arteries and released by subsequent nerve stimulation (Verbewen et al., 1983; Kawasaki and Takasaki, 1984).

There have been no published immunohistochemical studies upon 5-HT-like immunoreactive (5-HT-I) nerves of a major cerebral vessel at the electron microscopic level. Previous studies have looked at the innervation of the brainstem microvasculature and small intraparenchymal cerebral vessels (Kapadia and De Lanerolle, 1984; Itakura et al., 1985). It was decided therefore to examine 5-HT-I nerves in the perivascular plexus of a major cerebral blood vessel by immunohistochemistry at the electron microscopic level and to investigate a possible interrelationship with catecholaminergic nerve fibres.
3.2 MATERIALS AND METHODS

A total of 18 Wistar rats (150-200 g) were used. Ten animals were subjected to immunohistochemistry utilising anti-5-HT antibody either alone or preabsorbed with a variety of control substances to test the specificity of the antiserum. Four animals were given 5-OHDA (100 mg/kg i.p.) 24 and 12 h before sacrifice to demonstrate catecholaminergic nerves (Dail and Evans, 1978), in addition to being processed for 5-HT immunoreactivity subsequently. Four animals were given the catecholaminergic nerve uptake blocker Desmethylimipramine (DMI) (28.5 umole/kg i.p.) at 48 h, 24 h and 1 h prior to sacrifice and subsequently processed for 5-HT immunoreactivity. All the animals were deeply anaesthetised with ether, and either perfused via the ascending aorta with approximately 200 ml of fixative composed of 4% paraformaldehyde, 0.1% glutaraldehyde, in 0.1 M phosphate buffer pH 7.3, and followed by 500 ml of the same fixative minus glutaraldehyde or the brains were rapidly removed after decapitation and immersed in glutaraldehyde-free fixative for 2h then placed in 0.1 M phosphate buffer overnight at 4 °C . The Circle of Willis was then removed from the brain and the middle cerebral arteries dissected out from their origin to first major divisions and rinsed in 0.1 M phosphate buffer for 1 h. The brains of two of the animals that had been treated with 5-OHDA and the four animals that had been treated with DMI were cut by vibratome into 50 um thick coronal sections and rinsed in 0.1 M phosphate buffer for 3 h.
**Immunohistochemistry**

Immunohistochemistry was performed on vessel whole-mount and vibratome brain section preparations. All steps took place at room temperature unless otherwise stated. Following rinsing in 0.1 M phosphate buffer, the vessels and sections were incubated in 10% normal goat serum in 0.1 M phosphate buffer for 1 h, then transferred into the primary antibody (anti-5-HT antibody, Immunonuclear Corp.) diluted 1:500 in 0.1 M phosphate buffer for 1 h at room temperature, then for 40-44 h at 4 °C followed by 1 h rewarming to room temperature. Vessels and sections were then washed in 0.1 M phosphate buffer and incubated for 1.5 h in goat-rabbit immunoglobulin G (Seralab) diluted 1:50 in 0.1 M phosphate buffer. They were then washed in 0.1 M phosphate buffer for 3 x 10 min and placed in rabbit peroxidase-antiperoxidase complex (DACO) diluted 1:50 in 0.1 M phosphate buffer for 3 h. The vessels and sections were then rinsed in 0.1 M phosphate buffer followed by 0.1 M Tris HCl buffer pH 7.4. They were then preincubated for 10 min in 0.1% 3,3'-diaminobenzidene tetrahydrochloride (DAB) (Sigma) dissolved in 0.1 M Tris HCl buffer, then reacted for 10 min in 0.1% DAB containing 0.01% H₂O₂. After rinsing in 0.1 M Tris HCl buffer followed by 0.1 M phosphate buffer, the vessels and sections were postfixed for 30 min in 1% OsO₄ in 0.1 M phosphate buffer. They were then rinsed in 0.1 M phosphate buffer followed by sodium acetate buffer and stained in 1% aqueous uranyl acetate for 30 min. Vessels and sections were next dehydrated through a series of alcohols into propylene oxide and flat embedded in Araldite. Serial
ultrathin sections were cut from the midpoint of the vessels and mounted on 200-mesh grids. The ultrathin sections were stained with aqueous uranyl acetate followed by lead citrate and examined with a Philips-300 electron microscope. The flat embedded vibratome sections were examined by light microscopy and the raphe-nuclei and nucleus of the solitary tract identified. Appropriate regions were marked, cut and remounted on araldite stubs. After examination of semithin sections, these regions were trimmed to allow serial ultrathin sections of portions of the nuclei to be cut, stained and examined as described above.

**Immunohistochemical controls**

Specificity of the anti-5-HT antibody was tested by preabsorption of the primary antiserum with 5-HT (10^{-3} M) for 24h at 4 °C prior to use, and omission of the primary antiserum, both of which resulted in abolition of immunostaining of nerves. Cross-reactivity of the anti-5-HT antibody for histamine, dopamine, 5-OHDA and noradrenaline was excluded by preabsorption of the primary antiserum with each of these substances (10^{-3} M) for 24 h at 4 °C when immunostaining of nerves was not abolished.
3.3 RESULTS

In this study, about 15% of the perivascular axons and vesicle-containing profiles visualised showed some degree of 5-HT-like immunoreactivity. 5-HT-I nerves were identified from the outer adventitia, where they were usually grouped together with non-reactive nerves in bundles containing some 12-25 nerves ensheathed by the cytoplasm of Schwann cells, down to the adventitia-media border where smaller groups of nerves more loosely surrounded by Schwann cell processes lay (Fig. 1). In axons the immunoprecipitate was found diffusely in the cytoplasm and also associated with microtubules. Vesicle-containing profiles immunoreactive for 5-HT were also identified (Fig. 1) and again were present through all levels of the adventitia and occasionally in close apposition to smooth muscle cells. In 5-HT-I vesicle-containing profiles, the immunoprecipitate was associated with the membrane of both large and small vesicles as well as diffusely in the cytoplasm (Fig. 1).

Animals treated with the false transmitter 5-OHDA develop intensely electron-dense granules in both small and large vesicle populations of catecholaminergic nerves (Fig. 2). This reaction is specific for catecholaminergic nerves (Gershon et al., 1976; Rothman et al., 1976), and we found no evidence of 5-OHDA uptake into the truly serotonergic neurons, dendrites and axons of the brainstem raphe-nuclei or terminals in the nucleus of the solitary tract. In this study about 50% of the total perivascular nerves were found to be catecholaminergic, which agrees with previous descriptions (Iwayama et al., 1970). The vesicle content of
the 5-HT-I profiles and CA profiles appeared similar and consisted predominantly of a pleomorphic population of small 40–50 nm diameter agranular vesicles and a few large dense-cored vesicles 100–150 nm in diameter. Vesicle-filled profiles which were immunoreactive for 5-HT and simultaneously labelled with 5-OHDA were easily identified (Fig. 2). In these animals which were treated with 5-OHDA, 5-HT-I nerves were invariably found to be catecholaminergic also. However, of the total of CA nerves visualised, only some 25% demonstrated immunoreactivity for 5-HT.

In those animals that had been treated with the catecholaminergic nerve uptake blocker Desmethylimipramine, it was found that 5-HT-I was now absent in the perivascular nerves of the middle cerebral artery. However, 5-HT-I in the truly serotonergic neurons and axons of the brainstem raphe-nuclei was unaffected by this treatment.
3.5 DISCUSSION

5-HT is a potent and well documented constrictor of a variety of vessels, including the major cerebral vessels. Its mechanism of action is complex, interacting with that of adrenergic nerves. Exogenous application in low doses to isolated, peripheral and cerebral arteries in vitro causes constriction by a direct activation of postjunctional 5-HT$^2$ receptors (Toda and Fujita, 1973; Allen et al., 1974a; Edvinsson et al., 1978; Medgett et al., 1984). In subcontractile doses, application potentiates the vasoconstrictor response elicited by stimulation of perivascular sympathetic nerves by an identical mechanism (Morotoki and Su, 1981; Van Nueten et al., 1981; Medgett et al., 1984). Application of higher doses causes constriction by a prejunctional mechanism, 5-HT causing displacement release of noradrenaline to act upon alpha - adrenoceptors (Fozard and Mwaluko, 1976; Marin and Sanchez, 1980; Itakura et al., 1985). Griffith et al. (1982) were able to demonstrate that the transmural nerve stimulation evoked neurogenic response in the rabbit basilar artery that persists after adrenoceptor blockade (Lee et al., 1980) could be blocked by the 5-HT2 receptor antagonist ketanserin. In addition to this pharmacological interaction, it is now becoming increasingly apparent that perivascular catecholaminergic nerves in peripheral and cerebral arteries can take up 5-HT and release it subsequently upon nerve stimulation (Verbewen et al., 1983; Kawasaki and Takasaki, 1984).

In this study of rat middle cerebral arteries prepared by
immersion-fixation, 5-HT-I was found to be present only in nerves that were shown also to be catecholaminergic in nature. Approximately 25% of the perivascular catecholaminergic nerves displaying some degree of 5-HT-like immunoreactivity. These findings providing direct ultrastructural evidence to support recent physiological and pharmacological studies which have suggested co-localisation of these transmitter substances in the perivascular nerves of major cerebral vessels.

It is unclear, however, whether the 5-HT-I demonstrated in these perivascular catecholaminergic nerves represents synthesis or simply uptake and storage of 5-HT. A recent study (Sah and Matsumoto, 1987) has shown that some rat superior cervical ganglion sympathetic neurons grown in culture can release 5-HT in a Ca$^{2+}$ dependent manner. The study also showed that the released 5-HT could be derived both by uptake and by synthesis. When the neurons were grown in 5-HT-containing culture medium, their 5-HT stores were derived from uptake. However, when they grown in a 5-HT-free medium that had been conditioned by cardiac myocytes, some neurones were able to synthesise 5-HT from tryptophan.

To ascertain whether the 5-HT-I observed in the perivascular nerves of major cerebral vessels is derived from uptake or synthesis, rats were examined that had been treated with the catecholaminergic nerve uptake blocker DMI in a dose that has been previously shown to reduce uptake activity by 83% (Wong et al., 1974). Following treatment with DMI 5-HT-I could not be demonstrated in any of the
catecholaminergic nerves of the rat middle cerebral artery, whilst the 5-HT-I of the raphe-nucleus neurons and terminals in the solitary tract nucleus were unaffected. This would support the premise that the 5-HT-I seen in a proportion of perivascular catecholaminergic nerves is due to uptake rather than intrinsic synthesis.

The possibility still remains, however, that if the level of synthesis of 5-HT was low, then by preventing reuptake of released 5-HT with DMI the nerves could be depleted of 5-HT to a level insufficient to be visualised by immunohistochemical methods.
Fig. 1. Immunoelectron photomicrographs showing 5-HT-I axons and vesicle-filled profiles in the middle cerebral artery of the rat without 5-OHDA treatment. A: an immunoreactive axon is seen in a periadventitial bundle consisting of some 25 nerves surrounded by the cytoplasm of an adjacent Schwann cell. Bar = 1 um. B: an immunoreactive axon is seen together with 4 non-immunoreactive axons at the adventitia-media junction close to a smooth muscle cell (sm). Bar = 0.3 um. C: the same immunoreactive axon as is seen in Fig. 1A, in a subsequent serial section at a higher magnification. As in Fig. 1B, the immunoprecipitate is found diffusely in the cytoplasm and associated with microtubules. Bar = 0.2 um. D: an immunoreactive vesicle-containing profile within an adventitial nerve bundle. Bar = 0.3 um. E: a solitary immunoreactive vesicle-containing profile filled with a rich collection of small pleomorphic vesicles and a few large vesicles. Bar = 0.4 um. Inset: higher magnification of another vesicle-containing profile showing a large granular vesicle (arrowhead) with core material partially lost during preparation and many small pleomorphic vesicles. Bar = 0.2 um.
Fig. 2. Immunoelectron photomicrographs showing axons and vesicle-filled profiles in the middle cerebral artery of the rat with 5-OHDA treatment to demonstrate catecholaminergic nerves. A: an adventitial bundle from a vessel not processed for 5-HT-I staining. Note the distinct electron-dense granules that develop within both large and small vesicles. Sm, smooth muscle cell. Bar = 1 um. Inset: at higher magnification. Bar = 0.2 um. B: an adventitial nerve bundle from a vessel processed for 5-HT-I staining. Vesicle-filled profiles that simultaneously demonstrate 5-OHDA-induced electron-dense granules and 5-HT-I staining are easily identified (straight arrows) as well as non-immunoreactive profiles that only contain 5-OHDA-induced electron-dense granules (curved arrows). This suggests that only a proportion of CA profiles contain 5-HT. Bar = 1 um. Inset: at higher magnification. Bar = 0.2 um.
4. EXPERIMENTAL CHAPTER II:

5-HYDROXYTRYPTAMINE DEMONSTRATED IMMUNOHISTOCHEMICALLY IN RAT CEREBROVASCULAR NERVES LARGELY REPRESENTS
5-HYDROXYTRYPTAMINE UPTAKE INTO SYMPATHETIC NERVE FIBRES
Summary: This chapter re-examines by means of fluorescence immunohistochemical techniques, a proposed serotonergic innervation of major cerebral vessels in the rat. Earlier studies had demonstrated a dense perivascular plexus of 5-hydroxytryptamine immunoreactive nerve fibres upon major cerebral vessels in this and many other species. In this investigation however, it was found that 5-hydroxytryptamine immunoreactive nerve fibres are rarely observed in cerebral vessels prepared by perfusion-fixation in situ, and only form a well-developed plexus in vessels prepared, as in previous authors studies, by immersion-fixation. Prior treatment with a predominantly noradrenergic uptake inhibitor desmethylimipramine but not the serotonergic uptake inhibitor fluoxetine produced a major diminution in the 5-hydroxytryptamine immunoreactive plexus visualised in these immersion-fixed vessels. In addition, 5-hydroxytryptamine immunoreactive nerves were only occasionally observed in immersion-fixed vessels from animals that had been pretreated with 6-hydroxydopamine to produce adrenergic denervation. The removal firstly, of vessel-contained blood, by left ventricular perfusion with Krebs, prior to vessel dissection and immersion-fixation, resulted in an absence of 5-hydroxytryptamine immunoreactivity in perivascular nerves. Immunoreactivity could then be restored by briefly incubating vessels in krebs containing either blood or 5-hydroxytryptamine, before fixation. It would appear therefore that 5-hydroxytryptamine is rarely present under normal circumstances in the perivascular nerves of major cerebral
vessels and that previous descriptions of a dense serotonergic nerve plexus, represent 5-hydroxytryptamine in blood released during vessel dissection, being taken up via the noradrenaline-uptake system into perivascular sympathetic nerves. The possibility is thus raised that 5-hydroxytryptamine uptake and interaction within perivascular adrenergic nerves could occur in those cerebrovascular disorders where blood is released.
4.1 INTRODUCTION

Serotonin is well known for its marked effects upon the cerebral circulation (Harper and Mackenzie, 1977b; Mackenzie and Edvinsson, 1980). Intracarotid administration reduces cerebral blood flow profoundly by constriction of large calibre vessels and arteriovenous anastomoses, whilst small arterioles dilate (Verdouw et al., 1984; Saxena and Verdouw, 1985). The possibility that perivascular nerves upon the major cerebral vessels are involved in this response was raised, when studies in a variety of species (Baumgarten, 1972; Chan-Palay, 1976; Griffith et al., 1982; Edvinsson et al., 1983b; Chang et al., 1987; Cowen et al., 1987) including man (Griffith and Burnstock, 1983) suggested the presence of a specific serotonergic innervation.

The immunohistochemical studies employed, however, have all utilised immersion-fixation of vessels, and in some cases vessel incubation in vitro with 5-HT or the monoamine oxidase inhibitor pargyline to enhance 5-HT-like immunoreactive (5-HT-I) nerve staining. The virtual disappearance of such nerves following superior cervical ganglionectomy (Marco et al., 1985; Cowen et al., 1986; Cowen et al., 1987), which also results in the loss of noradrenergic sympathetic nerves (Nielsen and Owman, 1967; Iwayama et al., 1970), together with recent evidence demonstrating an ability of sympathetic nerves to accumulate exogenous 5-HT in vitro, (Verbeuren et al., 1983; Levitt and Duckles, 1986; Saito and Lee, 1987) has led to two questions being raised. Firstly, are all, or at least a majority of the serotonin-containing nerves demonstrated
previously in major cerebral vessels in fact sympathetic nerves, and secondly what is the origin of the serotonin that has been demonstrated within them.

The rat is an animal in which a serotonergic innervation of major cerebral vessels has been well described, (Edvinsson et al., 1983b; Chang and Owman, 1986; Cowen et al., 1987) it was decided therefore to undertake further studies upon it in order to elucidate these questions. This study was designed to assess whether the 5-HT-I plexus demonstrated in previous studies, was as equally developed in perfusion-fixed material, and if not, whether it could represent 5-HT uptake into sympathetic nerves during tissue dissection and preparation procedures.
4.2 MATERIALS AND METHODS

A total of 33 male Sprague-Dawley rats weighing 350-450g were used in this study. All animals were anaesthetised with 70% nitrous oxide, 30% oxygen and 1.5% halothane and were allowed to breathe spontaneously.

In the first group of experiments, seven rats were sacrificed by perfusion with 150 mls of fixative composed of 4% paraformaldehyde in 0.1 M phosphate buffered saline (PBS) through the left ventricle. The brains were then removed and all branches of the circle of Willis dissected out to yield a single preparation that included the anterior cerebral, middle cerebral, internal carotid, posterior cerebral, superior cerebellar, basilar and vertebral arteries. These vessels were then further fixed in the same fixative for 2 to 24 h.

In the second group, fourteen rats were utilised; all these animals were killed by decapitation under anaesthesia. The brains were rapidly removed, and the circle of Willis dissected out in a Krebs solution composition (in M): NaCl 0.12, KCl 0.1, KH₂PO₄ 0.05, MgSO₄ 0.03, NaHCO₃ 0.12, glucose 0.07, CaCl₂ 2.5. The vessels were then fixed by immersion in 4% paraformaldehyde in 0.1 M PBS for 2 to 24 h. Within this group of animals, 5 rats received no pretreatment and 9 rats, three in each category, were pretreated with one of the following drugs: the adrenergic neurotoxin 6-hydroxydopamine (6-OHDA) (100-150mg/kg) i.v. 72 and 24h before sacrifice, the noradrenergic uptake antagonist desmethylimipramine (28.5 umole/kg) i.p. 1h before sacrifice, or the serotonergic uptake antagonist fluoxetine
(28.5 umole/kg) i.p. 1h before sacrifice.

In the third group, 12 rats were utilised. These animals were killed by perfusion with Krebs through the left ventricle, the brains then rapidly removed and the circle of Willis and cerebral vessels dissected out. The circle of Willis from 3 rats were then immediately fixed by immersion in 4% paraformaldehyde in 0.1M PBS. Those from the remaining 9 animals were incubated at 37°C for 30 minutes in either Krebs alone (3 rats), Krebs containing 5-HT (10^-6 M), or Krebs containing 10% by volume of blood previously collected from another rat. Following 30 mins incubation, the circle of Willis vessels were fixed by immersion in 4% paraformaldehyde in PBS for 2-24 h. The Krebs solution used in incubation procedures was constantly aerated with a gas mixture of 95% O_2 / 5% CO_2.

**Immunohistochemistry**

Immunohistochemistry was performed on vessel whole-mount preparations. All steps took place at room temperature. After fixation all vessels were rinsed in PBS. To facilitate antibody penetration the vessels were then rinsed in 80% alcohol for 1h, and given short rinses in PBS with 0.1% Triton X-100. Vessels from one half of each circle of Willis were then incubated for 12-18 h in anti-5-HT antibody (Immunonuclear Corp.) diluted 1:400 with PBS containing 0.1% Triton X-100. After 3 PBS washes, vessels were exposed for 2 h to fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG (Nordic) at a dilution of 1:50. Vessels were briefly rinsed in PBS then placed in PBS with 0.05% pontamine sky blue and 1% dimethyl sulphoxide for 5-10 mins
in order to reduce background autofluorescence (Cowen et al., 1985). Vessels were next air-dried onto slides and coverslipped in an anti-fade mountant (Citifluor, City University, London, U.K.) for viewing. Vessels from the other half of each circle of Willis, utilised for the demonstration of neuropeptide-Y-like immunoreactive (NPY-I) nerves, were treated similarly but using an anti-NPY antibody (Immunonuclear Corp.) diluted 1:400.

**Immunohistochemical controls**

Specificity of the primary antibody was tested by preabsorption with excess antigen \(10^{-3}\)M for 24 h at 4°C prior to use and omission of the primary antiserum, both of which resulted in abolition of immunoreactivity. Cross-reactivity of the anti-5-HT antibody for histamine, dopamine and noradrenaline was excluded by preincubation with each of these substances at \(10^{-3}\)M when immunoreactivity was maintained.

**6-hydroxydopamine induced adrenergic denervation**

6-Hydroxydopamine was used to induce a selective degeneration of adrenergic nerve terminals (Thoenen and Tranzer, 1968; Costa et al., 1982), in order to investigate the effect of the resultant sympathetic denervation, upon the ability to demonstrate 5-HT containing nerves upon circle of Willis vessels. Confirmation of the effectiveness of the adrenergic denervation produced was ascertained as cerebral vessels from 6-OHDA treated and untreated animals were examined for the presence or absence of NPY-I nerves. NPY-I has been demonstrated previously, as localised within
the adrenergic nerves of cerebral vessels (Matsuyama et al., 1985; Edvinsson et al., 1987), and NPY-I in adrenergic nerves is markedly reduced, together with noradrenaline, following 6-OHDA treatment (Morris et al., 1986a; Edvinsson et al., 1987). In those animals in this study that received 6-OHDA, there was always a major reduction of NPY-I nerves throughout the circle of Willis whilst untreated animals in all other groups always demonstrated a dense plexus as previously described (Edvinsson et al., 1987).
4.3 RESULTS

**Group I Perfusion-fixed vessels:** In cerebral vessels taken from animals killed by perfusion-fixation, 5-HT-I nerves were only inconstantly and most infrequently observed (Fig. 3). In five of the seven animals employed, there were no 5-HT-IR nerves seen throughout any of the vessels comprising the circle of Willis. Whilst in the remaining two animals there were a few faint 5-HT-I nerves seen restricted to the vertebral and basilar arteries, the remaining superior cerebellar, posterior cerebral, internal carotid, middle cerebral and anterior cerebral arteries were otherwise entirely devoid of 5-HT-I nerve fibres.

**Group II Immersion-fixed vessels:** Within this group of animals killed by decapitation, followed by vessel dissection and subsequent immersion-fixation, the results were affected by the prior treatment received. In all five animals that received no pretreatment, a well defined plexus of 5-HT-I nerves was consistently demonstrated throughout all the vessels of the circle of Willis (Fig. 4). Pretreatment with the serotonergic uptake inhibitor fluoxetine (3 animals), produced only a minor diminution in the extent and intensity of the 5-HT-I nerve plexus (Fig. 5A and 5B). However, pretreatment with an equimolar dose of the noradrenergic uptake inhibitor desmethylimipramine (3 animals) resulted in the virtual abolition of the 5-HT-I nerve plexus (Fig. 5C and 5D). Similarly, in three animals pretreated with 6-OHDA to induce degeneration of adrenergic varicosities, it was found that 5-HT-I nerves were only seldom observed (Fig. 6).
Group III Krebs perfused vessels: In this group of animals perfused with Krebs to remove blood from the vessels prior to dissection, with or without an incubation, followed by immersion-fixation, the results obtained depended upon the incubation procedure. In the three animals whose vessels were immersion-fixed without any incubation, 5-HT-I nerves were never observed. Similarly in those vessels (3 animals) incubated in Krebs alone no 5-HT-I nerves were seen (Fig. 7A and 7B). However, in vessels incubated in either Krebs plus 5-HT (Fig. 7C and 7D) or Krebs plus blood (Fig. 7E and 7F) prior to immersion-fixation, the results differed. In these latter vessels, a strongly staining plexus of 5-HT-I nerves was once again seen throughout the circle of Willis.
4.5 DISCUSSION

Serotonin is predominantly synthesised by the chromaffin cells of the gastrointestinal tract (Erspamer and Testini, 1959). It is then released into the portal circulation, where that fraction that escapes inactivation by hepatocytes and subsequently the pulmonary bed, is taken up and stored within the granules of platelets. Serotonin synthesising neurons have also been demonstrated in the enteric plexus of the gut and the raphe nuclei of the brainstem (Dahlstrom and Fuxe, 1964; Dreyfus et al., 1977; Steinbusch et al., 1978; Costa et al., 1982). On the basis of these and other findings a variety of roles have been suggested for serotonin, in health and disease, in a number of sites. Serotonin released during platelet aggregation has been proposed to act upon the smooth muscle of injured vessels to produce local vasoconstriction and aid haemostasis (Van Zwieten, 1987). In the enteric plexus, serotonergic neurons and their processes would seem to be involved in the regulation of intestinal motility (Gershon and Sherman, 1987). With regard to the cerebral circulation, the serotonergic neurons of the raphe nuclei appear to make somal and dendritic contacts locally with parenchymal blood vessels of the brainstem, and also contact more distal intracerebral parenchymal blood vessels through long projection axons (Scheibel et al., 1975; Reinhard et al., 1979; Di Carlo, 1984; Kapaidia and de Lanerolle, 1984; Scatton et al., 1985). This has led to the suggestion of a possible role for serotonin in cerebral vasomotor control, regulating overall parenchymal cerebrovascular resistance,
in response to changes in brainstem blood flow, as sensed by the raphe neurons.

A completely separate, serotonergic innervation of the circle of Willis and large extraparenchymal cerebral vessels then appeared to exist, following the demonstration by fluorescence immunohistochemistry of a well developed 5-HT-I plexus about such vessels in a number of species, including man. Unlike the innervation of the smaller intraparenchymal cerebral vessels, this plexus was largely lost following removal of the superior cervical ganglia, which are known also to be the major source of adrenergic innervation to these major cerebral vessels. However, from an examination of perfusion-fixed material and by using uptake studies, this experimental chapter now clearly demonstrates that the major cerebral vessels of the rat, are normally only seldom and inconstantly innervated by occasional faint 5-HT-I nerves limited strictly to the vertebrobasilar arteries. When the same vessels are examined from animals that have been killed by decapitation, followed by vessel dissection and immersion-fixation, then a rich network of 5-HT-I nerves is readily demonstrated throughout the circle of Willis.

The hypothesis, that this difference is due to 5-HT from blood, released during dissection, being taken up via the noradrenaline uptake mechanism into sympathetic nerves was tested. Firstly, development of a well-defined 5-HT-I plexus in immersion-fixed material could largely be prevented by pretreatment with a predominantly noradrenergic uptake inhibitor but not with a more selective serotonergic
uptake inhibitor. Similarly, the development of a 5-HT-I plexus could largely be prevented, by prior treatment of animals with 6-OHDA which produces a selective degeneration of adrenergic nerve terminals.

In a separate series of experiments, the careful removal of vessel-contained blood by perfusion with Krebs solution prior to brain removal and vessel dissection, also resulted in the absence of any visible 5-HT-I plexus. Such a plexus was readily restored if vessels were then briefly incubated in Krebs containing either 5-HT (10^-6 M) or freshly collected blood (10% Vol).

Taken together, these results indicate that the well developed 5-HT-I nerve plexus previously described upon major cerebral vessels largely represents an in vitro uptake phenomenon, serotonin from blood released during vessel dissection being readily taken up into sympathetic nerves where it can subsequently be demonstrated by immunohistochemistry. The very small and inconstant number of 5-HT-I fibres that were occasionally seen in perfusion-fixed animals, largely restricted to the basilar artery, might represent centrally derived serotonergic fibres that have passed out along the perforating arteries to the raphe complex. Another possibility might be that 5-HT in locally elevated levels, derived from nearby raphe neurons, is taken up into adjacent sympathetic nerves.

The reasons why perivascular sympathetic nerves supplying cerebral vessels are capable of avid uptake of 5-HT needs to be examined. For example, is it a protective mechanism to remove potent concentrations of 5-HT arising in platelet
aggregation and other conditions or is it a biological device for reinforcing sympathetic vasoconstriction arising during certain physiological and pathological states. These and other questions need to be resolved in future studies.

The ready demonstration of such an ability in vitro is of interest, in that a similar mechanism of blood-derived 5-HT uptake and interaction in perivascular sympathetics may occur in vivo in cerebrovascular disorders such as following stroke, migraine attacks and in vasospasm following subarachnoid haemorrhage, where serotonin has frequently been implicated in the aetiology (Sicuteri et al., 1961; Anthony et al., 1969; Allen et al., 1974a; Voldby et al., 1982; Tagari et al., 1983; Richardson et al., 1987).
Fig. 3. Whole-mount preparation of perfusion-fixed vessels from rat circle of Willis. 5-HT immunofluorescence. (A) Basilar artery (B) Vertebral artery (C) Internal carotid artery (D) Anterior cerebral artery. In five out of seven perfusion-fixed animals no 5-HT-IR nerve fibres were seen in any of the major cerebral vessels. In the remaining two, one animal had a few faintly staining fibres upon the vertebral and basilar arteries while the other had occasional fibres upon one vertebral artery. Bar = 75 um. All micrographs have the same magnification.
Fig. 4. Whole-mount preparation of immersion-fixed vessels from rat circle of Willis. 5-HT immunofluorescence. (A) Basilar artery (B) Vertebral artery (C) Internal carotid artery (D) Anterior cerebral artery. In all five animals a well defined plexus of 5-HT-I nerve fibres was consistently demonstrated upon all major cerebral vessels. Bar = 75 um. All micrographs have the same magnification.
Fig. 5. Immunofluorescence photomicrographs of immersion-fixed cerebral vessel whole-mounts from rats pretreated with fluoxetine (A and B) or desmethylimipramine (C and D). A and C are corresponding portions of vertebral arteries whilst B and D are similar regions of anterior cerebral arteries. Compared as pairs: A against C, and B against D, it can be seen that numerous 5-HT-I fibres persist following fluoxetine administration whilst after desmethylimipramine, only a few fibres remain faintly immunoreactive. Bar = 75 um. All micrographs have the same magnification.
Fig. 6. Immunofluorescence photomicrographs of immersion-fixed rat Internal carotid artery whole-mounts. After treatment with 6-hydroxydopamine to produce a chemical sympathectom}y (B and D). Control vessels (A and C). Following 6-OHDA treatment, 5-HT-I fibres virtually disappeared (B) as compared with control vessels (A). The effectiveness of adrenergic denervation was evidenced by a concomitant loss of NPY-I fibres in 6-OHDA treated animal vessels (D) over controls (C). Bar = 50 um. All micrographs have the same magnification.
Fig. 7. Whole-mount preparation of circle vessels, perfused with Krebs to remove contained blood, and incubated in either Krebs alone (A and B), Krebs plus 5-HT (C and D) or Krebs plus blood (E and F) prior to immersion-fixation. In (A and B) 5-HT-I fibres were completely absent. Following incubation with 5-HT or blood a well-developed plexus of 5-HT-I nerves fibres was restored. Bar = 100 um. All micrographs have the same magnification.
5. EXPERIMENTAL CHAPTER III:

ALTERATIONS IN SEROTONIN AND NEUROPEPTIDE Y CONTENT OF CEREBROVASCULAR SYMPATHETIC NERVES FOLLOWING EXPERIMENTAL SUBARACHNOID HAEMORRHAGE
Summary: The effect of an experimental subarachnoid haemorrhage (SAH) upon neurotransmitter content in sympathetic nerves supplying the major cerebral arteries of the rat has been examined both by an immunohistochemical analysis, and by high performance liquid chromatography with electrochemical detection (HPLC-ECD). In particular, changes that occur in sympathetic nerve content of the vasoconstrictor agents serotonin (5-HT) and neuropeptide-Y (NPY), which are colocalized with noradrenaline were assessed. Subarachnoid haemorrhage was induced by a single injection of autologous arterial blood into the cerebrospinal fluid (CSF) space of the cisterna magna. The density of 5-HT-containing and NPY-containing perivascular nerves fibres per unit area of vessels was measured at defined intervals from 15 min to 5 days post-SAH. In addition, an HPLC study was performed to quantify the actual amounts of 5-HT and noradrenaline present in circle of Willis vessels at 3 h post-SAH. Comparison was made with sham-operated animals and animals that received a cisternal injection of buffered saline in place of blood. The results reveal a major increase in cerebrovascular sympathetic nerve content of serotonin, arising by uptake, presumably from subarachnoid blood clot, within the first 3 h post-SAH. Neuropeptide Y content, however, decreased from 3 h up to 48 h posthaemorrhage. By 3 days post-SAH, when the majority of subarachnoid clot had resorbed, the sympathetic nerve content of both NPY and 5-HT was restored to normal. This pattern of change was not observed in either sham-operated or saline-injected controls.
5.1 INTRODUCTION

Subarachnoid haemorrhage (SAH) is known, both clinically in humans (Grubb et al., 1977; Symon, 1978; Kwak et al., 1979), and from experimental studies in animals (Kuwayama et al., 1972; Petruk et al., 1972; Umansky et al., 1983; Svendgard et al., 1983) to be associated with the development of angiographically defined cerebral arterial spasm, reduced cerebral blood flow, and impaired autoregulation. The neural plexus that surrounds and innervates the major cerebral vessels is one of the factors known to control the calibre of these vessels (Edvinsson and MacKenzie, 1977), with sympathetic noradrenergic nerves making up a major component of this plexus (Iwayama et al., 1970). Studies, both in human SAH and from animal models, have revealed the development posthaemorrhage of sympathetic nerve dysfunction in cerebral vessels, with an impairment of noradrenaline uptake and synthesising ability (Lobato et al., 1980; Edvinsson et al., 1982; Tsukahara et al., 1988).

It has become clear from recent discoveries that the cerebrovascular sympathetic nerves contain other substances, in addition to noradrenaline, that can act as neurotransmitters and neuromodulators. One of these, the peptide neuropeptide Y (NPY), is known to cause strong concentration-dependent contractions of cerebral vessels (Edvinsson et al., 1987). Cerebrovascular sympathetic nerves in vitro have additionally been shown to readily take up serotonin (5-HT) into nerve terminals (Verbeuren et al., 1983; Levitt and Duckles, 1986; Jackowski et al., 1989a), from whence it can act as a vasoconstrictor false
transmitter (Saito and Lee, 1986). Following rupture of an intracranial aneurysm, 5-HT-containing platelets are released about vessels locally and throughout the subarachnoid space, with elevated 5-HT levels being found in the cerebrospinal fluid (CSF) post-SAH (Voldby et al., 1982; Tagari et al., 1983). In this study, the effects of an induced subarachnoid haemorrhage upon the content of 5-HT and NPY in the sympathetic nerves of major cerebral vessels in the rat, as revealed by immunohistochemistry, was examined for up to 5 days postictus. In addition, the levels of 5-HT and noradrenaline within the circle of Willis vessels in the period immediately following SAH, was measured by high performance liquid chromatography (HPLC) with electrochemical detection (ECD).
5.2 MATERIALS AND METHODS

Animal Preparations

One hundred and ninety male Sprague-Dawley rats weighing 350-450 grams were used in the experiments. The animals were anaesthetised with 70% nitrous oxide, 30% oxygen and 1.5% halothane during operative procedures and perfusions. While cisternal injections were made, the halothane was reduced to 0.75%. All animals were allowed to breathe spontaneously. In each animal the left femoral artery was cannulated and connected to a Statham P50 transducer. The arterial pressure was continuously recorded with a Lectromed recorder and blood gases were intermittently monitored.

In addition to a control group of 30 normal animals, the animals were divided into three groups, consisting of one group that underwent experimental subarachnoid haemorrhage (SAH), and two further groups of controls that received either a subarachnoid injection of buffered saline or sham operation only.

Induction of subarachnoid haemorrhage

In 78 animals, SAH was induced by a single injection of 0.3 mls of autologous arterial blood, freshly withdrawn from the femoral cannula and replaced with an equal volume of normal saline. The blood was injected at a rate of 0.1 mls/minute under operating microscope vision, via a 30 gauge needle passed through the previously exposed atlanto-occipital membrane, into the cisterna magna of the prone head-down animal.

In 50 animals, 0.3 mls of buffered normal saline was injected in place of autologous blood. Buffered normal
Saline was prepared by the addition of 25 meq/1 of NaHCO\textsubscript{3} and equilibrated at 37°C with a gas mixture of 5% CO\textsubscript{2}/95% O\textsubscript{2} to pH 7.4.

In a further 32 animals, a sham operation only was performed, the rats undergoing the same procedures to those receiving an SAH, except that no injections were given into the cisterna magna.

**Immunohistochemistry**

Immunohistochemistry was performed on vessel whole-mount preparations. All steps took place at room temperature. Animals for the immunohistochemical determination of the density of nerves upon major cerebral vessels were killed post-SAHI at times 15 min, 1 h, 3 h, 6 h, 1 day, 2 days, 3 days, and 5 days. Groups examined included normal, sham-operated, saline-injected, and SAH rats. Under anaesthesia, the left ventricle was perfused with 150 mls of fixative composed of 4% paraformaldehyde in 0.1M phosphate-buffered saline (PBS). The brain was immediately removed and all branches of the circle of Willis dissected out to yield a single preparation that included the anterior cerebral, middle cerebral, internal carotid, posterior cerebral and basilar arteries. Vessels were further fixed for 2 h, then rinsed in 3 changes of PBS for 1 h. To facilitate antibody penetration the vessels were then rinsed in 80% alcohol for 1 h and given short rinses in PBS with 0.1% Triton X-100. Vessels were then incubated for 16 h in anti-5-HT antibody (Immunonuclear Corp.) diluted 1:400 with PBS containing 0.1% Triton X-100. After three PBS washes, vessels were exposed for 2 h to fluorescein isothiocyanate (FITC)-conjugated
goat anti-rabbit IgG (Nordic) at a dilution of 1:50. Vessels were briefly rinsed in PBS, then placed in PBS with 0.05% pontamine sky blue and 1% dimethyl sulphoxide for 5-10 min in order to reduce background autofluorescence (Cowen et al., 1985). Vessels were next air-dried onto slides and coverslipped in an anti-fade mountant (Citifluor, City University, London, U.K.). Vessels used for the immunohistochemical demonstration of NPY content within perivascular nerves were treated similarly but using an anti-NPY antibody (Immunonuclear Corp.) diluted 1:400. Slides were viewed, and selected areas photographed at X 100 using a Zeiss microscope equipped for viewing FITC fluorescence.

Specificity of the anti-5-HT and anti-NPY antibody was tested by preincubation with $10^{-3}$ M concentration of 5-HT or NPY, respectively for 24 h at $4^\circ C$ prior to use, or omission of the primary antibody, both of which resulted in elimination of immunostaining of nerves.

The densities of nerves containing 5-HT or NPY were measured from photomicrographs taken at uniform exposure, film speed and development time. In every animal five standardised areas were sampled from the circle of Willis: one each from the anterior cerebral, middle cerebral, internal carotid, posterior cerebral, and basilar arteries. The number of nerve fibers seen crossing the diagonal of a rectangular area measuring 0.3 x 0.2 mm$^2$ in each of these five unit areas was counted and averaged per animal. Results are expressed as the overall mean ± SEM and data compared using the Student's $t$ test. Significance was accepted for
6-OHDA and Clomipramine experiments

To ensure, as seemed probable from previous studies, that the 5-HT- and NPY-immunoreactive nerve fibers counted following SAH truly represented sympathetic nerves, six animals that had been pretreated with the adrenergic neurotoxin 6-hydroxydopamine (6-OHDA) were also examined. Administration of 6-OHDA has been shown to induce degeneration of adrenergic nerve terminals (Thoenen and Tranzer, 1968) including those upon major cerebral vessels (Matsuyama et al., 1985) and was administered as two doses (100 and 150 mg/Kg i.v.) 72 and 24 h before SAH and/or killing.

To investigate further a supposition that the marked increase post-SA H of 5-HT-containing nerves was a consequence of 5-HT, released from subarachnoid blood clot, being taken up into sympathetic nerves, the effect of the noradrenergic nerve uptake antagonist clomipramine was studied. A single administration of clomipramine in a dose (57 umole/kg i.p.) previously shown to reduce adrenergic uptake activity by 83% (Wong et al., 1974) was administered 1 h prior to SAH in four animals.

High performance liquid chromatography

Animals for the HPLC determination of the total amounts of 5-HT and noradrenaline present in circle of Willis vessels were killed at 3h post-SA H. Three groups of animals were examined: normal, saline-injected, and SAH rats. Under anaesthesia the left ventricle was perfused with Krebs
solution to remove contaminating blood, contained within the cerebral vessels. Next, the brains were rapidly removed and transferred into ice-cold Krebs solution and the circle of Willis arteries resected and dissected clean under the operating microscope. Vessel preparations were briefly rinsed in three changes of ice-cold Krebs solution and the excess liquid blotted off before being placed in cryotubes for rapid freezing in liquid nitrogen, with storage at -20°C until later analysis. Vessels from six animals were pooled for each determination of 5-HT and noradrenaline content by HPLC. The total sample wet weight was approximately 9 mg. Samples were homogenised in 500 ul of ice-cold 0.1M perchloric acid containing 0.4 mM sodium bisulphite and 25 ng of dihydroxybenzylamine (DHBA) using a motor-driven glass-glass homogeniser, and cell debris was removed by low speed centrifugation. Aliquots (100 ul) of the supernatant were injected without further treatment for the determination of 5-HT by HPLC with electrochemical detection (HPLC-ECD) using the method of Reinhard et al. (1980). The remainder of the supernatant was subjected to alumina extraction (Keller et al., 1976). Noradrenaline and DHBA were separated on a 10 u Bondapak C-18 reverse phase column (Radial Pak, Waters, Harrow, U.K.) using a mobile phase of 0.1 M sodium dihydrogen phosphate (pH 5.0) containing 5mM heptane sulphonate, 0.1mM ethylenediaminetetraacetic acid, and 13% (vol/vol) methanol (Moyer and Jiang, 1978). Quantitation was performed with a glassy carbon electrode set at a potential of +0.72 V. Noradrenaline levels were corrected for recovery using the
DHBA internal standard. Results are expressed as the mean ± SEM and the data compared using the Student's t test. Significance was accepted for p<0.05.
5.3 RESULTS

Macroscopic appearance

In all animals examined shortly following SAH, there was extensive blood clot within the basal cisterns that extended throughout the subarachnoid space and about the circle of Willis (Fig. 8A). In the rat, there appears to be a rapid removal of blood from the CSF-containing spaces and, by the third day after haemorrhage, nearly all clot had disappeared (Fig. 8B). Preparation of clean vessels for examination was facilitated, since separation of overlying meninges resulted in removal of the majority of any adherent clot.

Changes in nerve density of 5-HT- or NPY-containing nerves (Figs. 9A-9C)

5-HT

In normal rats, there was as previously described, an extremely low and variable density of 5-HT-I nerve fibers found upon the circle of Willis vessels (Fig. 10A), the mean density being 0.08 ± 0.08. Following SAH however, the number of nerves containing 5-HT rose rapidly: to 10.2 ± 0.51 at 15 min, reaching a peak of 12.0 ± 0.34 at 1 h post-SAH (Fig. 10B). The numbers declined rapidly over the next 24 h to 1.8 ± 0.18, and once more were an infrequent finding by 3 days post-SAH. In the saline-injected and sham control groups, no significant changes in the density of 5-HT-containing nerves occurred.

NPY

In normal rats (Fig. 11A), the mean density of NPY-containing nerve fibers found in vessels from the
circle of Willis was 14.6 ± 0.35. Following SAH, the number of NPY-containing nerves declined gradually to reach a nadir at 24 h post-SAH (Fig. 11B) of 6.1 ± 0.23, a figure representing 42% of normal values. Recovery was seen by day 2, and a return to pre-SAH levels had occurred by day 3 post-SAH. In the sham control group, no significant changes in the density of NPY-containing nerves took place. However, in the saline-injected animals, a similar but far less marked diminution of nerve density occurred, with a maximum decrease to 76% of normal values, occurring on day 1 postinjection.

Effects of 6-OHDA or clomipramine treatment

The prior administration of the sympathetic neurotoxin 6-OHDA effectively prevented the marked increase in 5-HT-containing nerve fibres following SAH that had previously been observed (Fig. 10C). Similarly, in those animals that received the adrenergic nerve uptake inhibitor clomipramine 1h prior to SAH, any increase in 5-HT-containing nerves was also largely prevented (Fig. 10D). Following the administration of 6-OHDA, NPY-containing nerve fibres were only infrequently observed, both in control animals and in animals post-SAH (Fig. 11C and 11D).

Results of HPLC study (Table 1)

The amount of 5-HT present in circle of Willis vessels from normal animals was low (0.049 ± 0.015 ug/g). Experimental SAH caused nearly a threefold increase in the amount of 5-HT present in these vessels at 3 h post-SAH.
However, in saline-injected animals, at 3 h post-injection the level was not found to differ significantly from that of normal animals. The amount of noradrenaline present in circle of Willis vessels from normal animals was $1.80 \pm 0.03$ ug/g. Neither experimental SAH nor saline-injection caused a statistically significant change in the amount of noradrenaline present 3 h postinjection, although there was always a trend towards a slight reduction following SAH.
5.5 DISCUSSION

Cerebrovascular dysfunction remains a major cause of mortality and lasting disability in patients following subarachnoid haemorrhage. The results of a number of studies have indicated that vasoactive agents are released as a result of the blood clot that forms about major vessels and within CSF cisterns after SAH (Boullin, 1985). The CSF from patients following SAH has been found to possess significant vasoconstrictor activity upon cerebral vessels (Allen et al., 1976), in addition, the degree of contractile activity exhibited has been shown to be greatest in those patient with evidence of severe angiographic spasm (Kaye et al., 1984).

Increased levels of a number of putative mediators of cerebral vasospasm have been found in CSF following SAH, including 5-HT, noradrenaline, prostaglandins, free radicals, and oxyhaemoglobin (Shigeno et al., 1982; Tagari et al., 1983; Sakaki et al., 1988). While it has not been clearly defined what role the neural plexus that innervates cerebral vessels plays in the regulation of cerebral blood flow under normal conditions, there is abundant evidence to suggest that the sympathetic nerves are involved in autoregulation (James et al., 1969; Hernandez et al., 1971) and may exert a protective influence under certain pathological conditions (Heistad et al., 1982). Following SAH, it has been demonstrated that there arises a dysfunction of the noradrenergic component of perivascular sympathetic nerves. However, very little is known of the effects of SAH upon the content of other vasoactive
transmitters and neuromodulators such as NPY and ATP, which are contained also within sympathetic nerves. In addition, no studies to date have examined the consequences of SAH upon the 5-HT content of cerebrovascular sympathetic nerves, which are capable of an efficient uptake of 5-HT through their amine uptake mechanism (Levitt and Duckles, 1986). It is perhaps relevant that a recent study in the dog coronary artery following experimental coronary occlusion, with platelet aggregation, has demonstrated an accumulation of 5-HT into adrenergic nerves with subsequent adrenergic dysfunction (Cohen et al., 1987).

The results of this study clearly demonstrate that, following SAH, the 5-HT content of cerebral vessels increases, and that 5-HT is rapidly taken up via their amine uptake mechanism into sympathetic nerve terminals, from which it can be readily detected by specific immunofluorescence.

Other workers have previously demonstrated that SAH may transiently decrease the noradrenaline content of cerebrovascular sympathetic nerves (Edvinsson et al., 1982; Delgado et al., 1985). A finding from this study is that the NPY content of sympathetic nerves is similarly decreased, by nearly 60%, in a transient manner following SAH. A smaller, 24% decrease in content following saline-injection was also noted, and may represent a stress-release response, as has been demonstrated to occur peripherally by other workers (Morris et al., 1986b).

It has been suggested that the observed decrease in catecholamine content in sympathetic terminals could be due
to an increased release of contained neurotransmitter and is in some way triggered by SAH. It is well known that 5-HT directly causes vasoconstriction of cerebral vessels by specific receptors (Chang and Owman, 1987). However, in addition, in higher concentration, 5-HT taken up into nerves in vitro can trigger, by a tyramine-like effect, the displacement release of noradrenaline from sympathetic nerves (Marin et al., 1981) and, presumably other, contained vasoactive agents such as NPY and ATP that are also displaced by tyramine (Cheng and Shen, 1987; Kirpatrick, personal communication) and that would then be free to act directly upon the vessel wall.

This study has demonstrated following experimental SAH in vivo in the rat, that 5-HT is indeed taken up into the perivascular sympathetic nerve endings of major cerebral vessels and that coincident with this there is a temporary decrease in NPY content. Whether these two phenomena represent a causal or casual relationship remains to be determined, but is yet further evidence of a transient marked dysfunction of cerebrovascular sympathetic nerves that arises post-SAH.
Fig. 8. Ventral view of brain from animals in which subarachnoid haemorrhage has been induced by the cisterna magna injection of 0.3 ml of autologous arterial blood. A: SAH induced 1h prior to killing showing extensive blood throughout the subarachnoid space and surrounding major cerebral vessels. B: SAH induced 3 days prior to killing showing the rapid resolution of subarachnoid blood clot in the rat.
Fig. 9. Graphs showing changes in the mean densities of 5-hydroxytryptamine and neuropeptide Y-containing perivascular nerves of major cerebral vessels up to 5 days following SAH or control procedures. A: Following subarachnoid haemorrhage. B: Following injection of buffered saline in place of blood. C: Following a sham operation only.

The vertical bars represent 2 SEM (* p <0.001, ** p <0.01).
nerve density

A

SAH GROUP

• NPY
○ 5-HT

B

SALINE GROUP

• NPY
○ 5-HT

C

SHAM GROUP

• NPY
○ 5-HT
Fig. 10. Photomicrographs of rat internal carotid arteries prepared to show 5-HT immunofluorescence. A: From a normal control animal, no 5-HT-containing nerve fibres are visible. B: From an animal at 1h following induction of subarachnoid haemorrhage, a well developed plexus of 5-HT-containing nerve fibres is now seen in the wall of the vessel. C: At 1h following subarachnoid haemorrhage from an animal previously treated with 6-OHDA to produce adrenergic denervation, virtually no 5-HT-containing nerve fibres appear. D: At 1h following subarachnoid haemorrhage from an animal treated 1h prior to SAH with the noradrenergic uptake antagonist clomipramine, the development of 5-HT-containing nerve fibres is markedly diminished. Bar = 100 um. All micrographs have the same magnification.
Fig. 11. Photomicrographs of rat internal carotid arteries prepared to show NPY immunofluorescence. A: From a normal control animal, a dense plexus of NPY-containing nerve fibre is always visible. B: From an animal at 1 day following induction of subarachnoid haemorrhage, a plexus of NPY-containing nerves is still apparent but this is markedly reduced both in its density and intensity. C and D: Following pretreatment of animals with 6-OHDA to produce adrenergic denervation, NPY-containing nerve fibres have virtually disappeared both in normal controls (C) and in animals at 1 day post-SAH (D). Bar = 100 um. All micrographs have the same magnification.
Table 1. Total amounts of serotonin (5-HT) and noradrenaline (NA) in circle of Willis vessels 3 h post-SAH, compared with control vessels.

<table>
<thead>
<tr>
<th>Animal group</th>
<th>Weight (mg)</th>
<th>5-HT (ug/g tissue)</th>
<th>NA (ug/g tissue)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAH rats (n = 30)</td>
<td>8.69 ± 0.30</td>
<td>0.121 ± 0.012</td>
<td>1.76 ± 0.04</td>
</tr>
<tr>
<td>Normal rats (n = 18)</td>
<td>9.18 ± 0.45</td>
<td>0.049 ± 0.015</td>
<td>1.80 ± 0.03</td>
</tr>
<tr>
<td>Saline injected rats (n = 18)</td>
<td>8.87 ± 0.59</td>
<td>0.035 ± 0.008</td>
<td>1.98 ± 0.12</td>
</tr>
</tbody>
</table>

Values are means ± SEM. A significant increase in 5-HT occurred in SAH vs. control animals (*p < 0.02). There was no significant difference between saline-injected and normal animals and no significant difference in weight between groups. n = number of animals in each group.
6. EXPERIMENTAL CHAPTER IV:

SERIAL DETERMINATIONS OF REGIONAL CEREBRAL BLOOD FLOW IN THE RAT USING SIMPLE CHRONICALLY IMPLANTED PLATINUM WIRE MICROELECTRODES
Summary: The ability to reliably obtain serial determinations of regional cerebral blood flow (rCBF) over time in small animals using simple and inexpensively constructed platinum-wire microelectrodes has been investigated. Repeated measurements of local cortical flow were obtained on a daily basis using the hydrogen clearance technique in a group of 18 animals, in each of which, six electrodes were chronically implanted. These studies have shown the simple microelectrodes utilised to give a low variability of results, as well as being well tolerated and provoking minimal tissue reaction even over prolonged periods of time. Under identical steady state conditions, serial determinations of rCBF were obtained for up to one week, with a maximum variation in mean values of only 12%, a result that compares favourably with the known serial determination error for the technique in the shorter term.
6.1 INTRODUCTION

Methodologies currently employed for the measurement of cerebral blood flow (CBF) in small animals include the radioactive microspheres technique (Heymann et al., 1977), $^{14}$C-Iodoantipyrine autoradiography (Sakurada et al., 1978), and hydrogen clearance (Auckland et al., 1964). Of these, the autoradiographic technique is unsuitable for repeated studies in the same animal since it requires tissue sections while the microspheres technique is limited by the total number of microspheres that can safely be injected into small animals (Solomon et al., 1985). The hydrogen clearance technique using cortically implanted platinum electrodes, has regularly been used for acute studies of cerebral ischaemia in small animals. It was desired however, to extend the technique to allow repeated determinations of CBF over longer periods of time to be obtained in the same animal.

Platinum microelectrodes have only infrequently been employed before for determining regional CBF (rCBF) in chronic experiments, usually in larger animals such as the cat (Kummer 1984), and no systematic investigation of the reliability of the hydrogen clearance technique in chronic studies has been reported. Although platinum is known to be well tolerated and to provoke minimal tissue reaction when implanted in the brain (Summerlee et al., 1982), it was not known whether tissue trauma over time would create a changing or excessive zone of damaged cells and tissue fluid around the electrode site. Such a zone could potentially create unstable electrical baselines and act as a diffusion
barrier, affecting the sensitivity and shape of the hydrogen clearance curves obtained, rendering the technique impractical for measurements over any prolonged period of time. This study was designed to answer these questions and to assess also the duration of functionally useful electrode survival times.
6.2 MATERIALS AND METHODS

Construction of microelectrode assembly

The microelectrode used was constructed from a commercially available electronics printed-circuit board socket (30S/093/V Oxley Developments, Ulverston, Cumbria, UK) to the connector pin of which, a short length of 100 μm diameter teflon-coated platinum-iridium wire (Clarke Electromedical Instruments, Pangbourne, Reading, UK) was soldered. The pin and soldered join was then electrically insulated with quick setting epoxy-resin (RS Components, Corby, Northants, UK) applied to form a tear-drop shape (Fig. 12). When the resin had hardened the platinum-wire was shortened to a final length of 1.5mm, and the distal 0.5mm of the electrode bared of teflon insulation.

Physiological preparations and measurement of CBF

A total of 24 male Sprague-Dawley rats weighing between 350-450 g were used in the experiments. Implantation and fixation of the electrode-assembly was performed in animals anaesthetised on a mixture of 70% nitrous oxide: 30% oxygen, with 1.5% halothane added and who were allowed to breathe spontaneously. In each animal, six electrodes were implanted as left and right pairs into parietal, occipital and cerebellar cortical regions, respectively. Following cleansing of the scalp with 70% alcohol, the superior aspect of the skull was exposed via a single midline sagittal incision and the skin edges drawn back. The pericranium was then excised and burr holes made over the specified cortical areas, with saline irrigation to avoid local heating. The electrode assembly, together with a small drop of
cyanoacrylate glue (RS Components) applied to the base of the epoxy-resin tear drop, was then inserted into the burr hole. By making the burr hole and the tear drop of similar diameters, immediate fixation followed contact of the epoxy-resin upon the burr hole shoulder. Each electrode was then firmly secured by running further epoxy-resin onto the tear drop and surrounding bare cortical bone. The scalp edges were approximated with sutures and a small scalpel blade nick made in the skin directly over each electrode to allow the connector sockets to protrude snugly through the scalp, enabling lead connections to be made via a matching male connector plug (30P/093 Oxley Developments). The reference electrode employed was constructed from 320 µm diameter Ag/AgCl wire passed during each set of recordings via a 21 gauge needle subcutaneously into the tissues on the back. CBF was measured at each electrode site by the hydrogen clearance technique as previously detailed (Auckland et al., 1964; Pasztor et al., 1973; Young 1980). Hydrogen gas was administered in the inhaled anaesthetic gases to a final concentration of approximately 7% and was given via a parallel circuit to ensure that inspired concentrations fell immediately to zero at onset of desaturation. Hydrogen inhalation times were kept at a constant 3 min for all flow measurements which, for the rates of flow obtained in our animals, should result in a more than 90% saturation and subsequent desaturation of tissues within this time period (Farrar, 1987). CBF was then calculated from the resultant clearance curves using the initial slope method (Olesen et al., 1971; Doyle et al.,
For flow recordings, animals were induced and initially anaesthetised on a mixture of 70% nitrous oxide : 30% oxygen together with 1.5% halothane. After being allowed to settle on this mixture for 30 min, the halothane was then reduced to 0.75% for a further 30 min before flow recordings were undertaken. The electrical apparatus together with the polarising voltage (+400 mv) was turned on for 30 min before recordings, to allow electrode stabilisation to steady baseline values to take place. Three hydrogen clearance recordings were then made separated by 10 min intervals, the CBF being calculated from the final curve obtained. Satisfactory electrode function was defined by: the attainment of a stable electrical baseline at 30 min following the application of the polarising voltage, electrode sensitivity remaining constant over the period of three clearance recordings, and finally, the formation of clearance curves of typical shape and appearance. Only flow recordings from electrodes that fully satisfied these criteria were utilised for the serial determination of cortical flows. Recordings were made for up to 10 days following implantation; the CBF being calculated then expressed as a percentage of each animals initial CBF value, obtained 2h following electrode implantation on day 0.

Histological analyses

In six rats, electrodes were implanted as previously described and the animals sacrificed subsequently for the microscopic localisation and assessment of electrode tracks at 2h, 1day, 3days, 5days, 7days and 10days.
postimplantation. Under deep inhalational anaesthesia, animals were perfused through the left ventricle with approximately 150 ml of fixative composed of 4% paraformaldehyde in 0.1M phosphate buffered saline. The overlying skull with microelectrodes still attached was carefully removed and the brains immersed in the same fixative overnight at 4 °C. The brains were then rinsed in 0.1M phosphate buffered saline and processed for embedding in paraffin wax. Sections 7um thick were cut and stained with Haematoxylin and Eosin for viewing by light microscopy. Using a calibrated eyepiece, the diameter of the electrode track, distance and numbers of neurons, reactive glia and macrophages from the electrode track was assessed.
6.3 RESULTS

Histological findings

There was only a minimal immediate tissue reaction provoked by the insertion and presence of the platinum-wire microelectrodes; this did not appear to change appreciably with the subsequent passage of time. The resultant track cavity was largely similar in diameter to that of the platinum wire used (100 um) (Fig. 13). At 2h following implantation, the track boundaries consisted of the brain parenchyma itself together with a small number of extravasated erythrocytes. By the third day post-implantation, fibroblasts appeared in the superficial portion of the track and by seven days, a smooth fibroblast lining in continuity with the overlying meninges had formed. Immediately surrounding the microelectrode track was an annulus of approximately 125 um in which a mild microglial reaction occurred. Neurons of normal appearance were usually present at a distance of some 50 um from the track boundary.

Electrode tolerance

Animals appeared to tolerate the electrodes well, feeding and behaving normally. No cases of infection were seen at any electrode site and there were no instances of detectable loosening of the microelectrode assembly up to seven days postimplantation. After this time however, some instances did occur, and after ten days, 24% of electrodes showed evidence of loosening in their attachment to the skull.
Functional electrode survival times

108 electrodes were implanted in 18 animals for the serial determination of regional cortical flow by the hydrogen clearance technique. Out of the total implanted, some 103 microelectrodes (95%) fulfilled our criteria for acceptable electrode function, producing satisfactory hydrogen clearance curves on day 0 at 2h postimplantation. The subsequent time course for continued satisfactory electrode function is detailed in Fig. 14. These results indicate that for periods of study of up to five days, when over 60% of implanted microelectrodes functioned satisfactorily, that a serial assessment of CBF during this time at least is practicable. After five days the numbers of satisfactory electrodes declined more rapidly, with less than 20% of microelectrodes continuing to function satisfactorily into the second week.

CBF variability

The mean ± SEM values of regional blood flow obtained 2 h postimplantation were 148 ± 4 ml/100g/min (parietal cortex), 158 ± 5 ml/100g/min (occipital cortex), and 93 ± 2 ml/100g/min (cerebellar cortex), figures in general agreement with values obtained previously in the rat (Cremer and Seville, 1983; Kangstrom et al. 1983). Subsequent serial determinations of flow under identical steady-state conditions demonstrated that repeated blood flow measurements could be obtained over periods of several days with, in some individual cases, quite remarkably low variability (Fig. 15). The overall results for regional flows measured at increasing times up to ten days from
microelectrode implantation, is detailed in Fig. 16. The results demonstrate, that for periods up to seven days following microelectrode implantation, that it is possible under the same steady-state conditions to obtain mean values of regional cortical flow that vary by a maximum of only 12% from those values obtained shortly after implantation.
6.5 DISCUSSION

rCBF changes may develop immediately, or evolve following a delay, after a widely varied number of pathophysiological events. Study of such changes for periods greater than a few hours in small animals such as the rat, has generally necessitated the use of a number of matched groups of animals at multiple time points to acquire the same data as would have been obtained by serial flow determinations over time in a single group of animals. The usage of simple chronically implanted platinum-wire microelectrodes as we describe, allows just such a serial determination of CBF to be readily performed. As well as allowing the use of smaller animal numbers to achieve the same data, such a technique should also reduce variability due to normal interanimal flow differences, as the flow data in each individual animal may now be expressed as a percentage of its own control value.

Such theoretical advantages can only be realised, however, if the serial determination of CBF by this method can be shown to be both practical, and capable of achieving a similar degree of accuracy over periods time, as has been demonstrated already in the shorter term. The hydrogen clearance technique has been shown over short term observations to have a serial determination error of up to 10%, other conditions being equal (Pasztor et al., 1973). In this study it has been possible under identical steady-state conditions to achieve serial determinations of regional cortical flow for up to one week with no more than a 12% variation in the mean flow values obtained.
It is felt therefore, that the serial assessment of CBF in small animals using simple platinum-wire microelectrodes is both practical and for periods of study up to one week, possesses a margin of experimental error not dissimilar from that for the technique in the shorter term.
Fig. 12. Diagramatic representation of the construction and approximate dimensions of a microelectrode assembly ready for implantation. a: printed-circuit board socket. b: epoxy-resin tear-drop insulation. c: platinum-wire soldered to connector pin. d: teflon insulation coating. e: 0.5 mm exposed electrode tip.
Fig. 13. Photomicrographs of cortical implantation sites showing microelectrode tracks. Animals sacrificed at 2h postimplantation on day 0 (upper figure) and day 10 postimplantation (lower figure). Scale bar = 100 um.
Fig. 14. Graph showing the time course of continued satisfactory electrode function that was obtained in 108 chronically implanted microelectrodes, studied for up to ten days following regional cortical insertion.
Percent Electrodes functioning

Days post implantation

Percent. Electrodes functioning
Fig. 15. Typical examples of the hydrogen clearance curves and semilogarithmic plots with calculated flows, obtained in an animal 2h post electrode implantation (day 0), and one week subsequently (day 7). At day 0 all six electrodes function satisfactorily. Although by day 7 only four electrodes now remain satisfactory, these produce clearance curves and reveal flows very similar to those obtained initially.
Fig. 16. Regional cortical blood flows measured serially under identical steady-state conditions, plotted for up to 10 days after microelectrode implantation. Flows are expressed as a percentage of 2h postimplantation control blood flow value. (—●—, parietal cortex; —●—, occipital cortex; —○—, cerebellar cortex). Values are means ± SEM.
7. EXPERIMENTAL CHAPTER V:
SUBARACHNOID HAEMORRHAGE IN THE RAT MODEL: THE TIME COURSE
OF ALTERED CEREBRAL BLOOD FLOW, CEREBRAL PERFUSION AND
INTRACRANIAL PRESSURE
Summary: The rat subarachnoid haemorrhage (SAH) model was studied in order to establish the precise time course of the globally reduced cerebral blood flow (CBF) that follows, and to ascertain whether related concomitant temporal changes in cerebral perfusion pressure (CPP) and intracranial pressure (ICP) take place. Studies were carried out upon spontaneously breathing animals under inhalational anaesthesia. SAH was induced in animals by a single intracisternal injection of 0.3 ml of blood; controls received either 0.3 ml of buffered saline in place of blood or underwent sham procedures only. Serial measurements of regional cortical cerebral blood flow were obtained by the hydrogen clearance technique, both acutely and for up to 5 days subsequently. Experimental SAH in the rat resulted in an immediate 50% global reduction in cortical flows that persisted for up to 3 h post-SAH. At 24 h, flows were still significantly reduced at 85% of their control values (p<0.05), but by 48 h, flows had regained their normal values and these were maintained up to 5 days post-SAH. ICP rose acutely after haemorrhage to nearly 50 mmHg with C-type pressure-waves being present. ICP then fell slowly, only fully returning to control levels at 72 h. Acute hydrocephalus was observed in autopsy studies of SAH animals but not in controls. Reductions in CPP occurred post-SAH, but only in the order of 15%, which could not alone account for the fall in CBF that took place. These findings indicate that this increasingly utilised small animal model of SAH develops only acute global flow changes, and it is likely that early vasospasm, secondary to released blood products
rather than pressure changes per-se, is responsible for the initial cerebral ischaemia that develops.
Subarachnoid haemorrhage (SAH) resulting from the rupture of an intracranial aneurysm in man, may lead to decreased cerebral blood flow (CBF), raised intracranial pressure (ICP) with hydrocephalus and the development of prolonged cerebral arterial spasm. Considerable evidence points to cerebral ischaemia resulting from reduced cerebral perfusion as the major cause of delayed neurological deterioration following SAH (Heilbrun et al., 1972; Ischii, 1979; Kutsuzawa et al., 1983; Voldby et al., 1985b), the peak incidence in man being around seven to ten days after a bleed (Pickard et al., 1989). Cerebral arterial spasm is one factor that has been significantly associated with a reduced CBF, and the presence of vasospasm is also known to be related both to the patient's clinical grade and eventual outcome (Loach and De Azevedo-Filho, 1976; Weir et al., 1978). It is likely however that the cause of the cerebral ischaemia that occurs post-SAH is multifactorial in origin, delayed hypoperfusion having been found to be associated not only with angiographically demonstrated luminal narrowing of vessels but also with the development of episodes of hypovolaemia, raised intraventricular pressure and ventricular dilatation (Ferguson et al., 1972; Ischii, 1979; Wijdicks et al., 1985).

Increasingly, in place of larger animal models such as the dog, cat, and monkey (Brawley et al., 1968; Kuwayama et al., 1972; Petruck et al., 1972; Umansky et al., 1984), the laboratory rat has been employed for experimental studies of SAH (Barry et al., 1979; Lacy and Earle, 1982; Solomon et
al., 1985; Svendgaard et al., 1985). Studies in the rat have demonstrated acute reductions in CBF, a biphasic pattern of vasospasm, and changes in cerebral metabolic activity and perivascular neurotransmitter content, consequent upon the release of homologous arterial blood into the subarachnoid spaces. Surprisingly however, little information exists regarding the subsequent time course of the globally reduced CBF that develops acutely post-SAH in the rat. Similarly, few details are known regarding the temporal development of changes in intracranial and cerebral perfusion pressure (CPP), and their interrelationship with reduced CBF following experimental SAH in the rat model. This study was designed therefore, to precisely determine the exact time course of such changes in the CBF occurring both within minutes and up to several days following the event, together with the obtaining of serial measurements of ICP and CPP over the same period of time.
7.2 MATERIALS AND METHODS

Animal Preparations

Experiments were performed on 90 male Sprague-Dawley rats weighing between 350 and 450 g. Animals were allowed to breathe spontaneously on an anaesthetic mixture consisting of 70% nitrous oxide : 30% oxygen, with 1.5% halothane added during operative procedures; reduced to 0.75% halothane 30 min before cisternal injections or measurements of CBF were performed. The left femoral artery was cannulated; arterial blood pressure was monitored with a Statham P50 pressure transducer (Statham Instruments Inc., Oxnard, California) and was recorded on a Lectromed MX6 chart recorder (Ormed Ltd., Welwyn Garden City, U.K.). Arterial blood samples were obtained anaerobically in microhaematocrit tubes (0.1 ml), and systemic $P_{a}CO_2$, $P_{a}O_2$ and pH determined using an ABL 30 blood-gas analyser (Radiometer, Copenhagen, Denmark). The haematocrit was measured. Body temperature was maintained at 37 °C by external warming.

Induction of Experimental SAH

SAH was induced in 30 rats by a posterior craniocervical approach using a modification of a technique described previously (Solomon et al., 1985). Briefly, utilising the operating microscope the posterior cervical musculature was separated in the midline, exposing the arch of the atlas, occipital bone and atlanto-occipital membrane. For production of haemorrhage, 0.3 ml of arterial blood was withdrawn by syringe from the femoral arterial line and replaced with an equal volume of normal saline. The freshly withdrawn blood was then injected into the subarachnoid
space via a 30 gauge needle passed into the cisterna magna. Injections were given as six equal aliquots over a period of 3 to 4 minutes. Control groups comprised: one group of 30 rats that received an intracisternal injection of 0.3 ml buffered saline (normal saline with 25 meq/1 NaHCO₃ equilibrated at 37 °C with 5% CO₂ : 95% air to pH 7.4), and a second sham-operated group of 30 rats that underwent identical operative procedures including cisternal needle puncture to the other two groups but in which no injections were given.

**CBF Measurements**

Regional cortical grey matter CBF (rCBF) was measured by the hydrogen clearance technique (Auckland et al., 1964; Young, 1980) using a chronically implanted platinum-wire microelectrode array (Jackowski et al., 1989c). In six animals from each treatment group, three pairs of microelectrodes were inserted into the parietal, occipital and cerebellar cortical regions of both left and right-hand hemispheres. In each animal the rCBF was determined pre-injection, after 3, 15, 30, 60, 120, 180 min and 1 to 5 days following SAH, saline or sham operation. The values of rCBF were calculated from the hydrogen clearance curves obtained, using the initial slope method (Oleson et al., 1971; Doyle et al., 1975) between 30 and 120 sec. The regional flows derived were then expressed as a percentage of each animal's control rCBF value obtained prior to injection on day 0.
ICP CPP and Other Measurements

ICP was measured via a saline-filled 30 gauge cannula placed into the cisterna magna and connected to a Statham P50 pressure transducer. Data values for ICP, CPP, MABP, haematocrit and blood gases for the times immediately post-SAH up to 3 h post event were acquired in the same animals as were employed also for the serial assessment of rCBF. Studies of these same parameters at the time points 1 to 5 days post-SAH, were performed in separate groups of animals, each group consisting of six animals, studied at those specified time points alone.

Statistical Analysis

All values are given as the mean ± SEM unless otherwise stated. Statistical comparison between experimental groups at different time points was performed using a two-tailed t test. p < 0.05 was taken as the level of significance.
7.3 RESULTS

rCBF Changes

Regional cortical flow changes were analysed separately for parietal, occipital, and cerebellar areas. The alterations in flow observed however were essentially global, affecting all cortical areas to the same degree (Fig. 17). The regional flow data from each animal have therefore been combined to give a mean cortical flow value that was used for further comparisons and in the statistical analysis between treatment groups. The mean cortical flow changes following experimental SAH, saline-injection, or sham-operation are summarised in Table 2. Following experimental SAH there was an immediate and marked reduction in flow, the overall CBF falling by 15 minutes to $50 \pm 2\%$ of pre-injection values ($p < 0.001$). This reduction persisted up to 3 h post-haemorrhage. At 24 h, flow was just significantly reduced still at $85 \pm 7\%$ of control values ($p < 0.05$). Pre-haemorrhage values were regained on the second day and remained normal up to 5 days subsequently. The mean cortical flow change in animals receiving cisternal saline-injection was similar to that obtained in sham-operated animals. Neither underwent rapid or marked changes in flow; in both groups after 30 minutes of the acute study there was a gradual reduction in the mean cortical flow reaching 80-85\% of pre-haemorrhage values at 3 h. In both groups of control animals, the mean cortical flow had fully recovered to pre-haemorrhage values by 24 h.
ICP Changes

The changes in ICP differed markedly in animals that underwent experimental SAH, from those that received cisternal saline-injection or underwent sham procedures. Differences in response occurred both in the period during and acutely following injection as well as in the time course of subsequent recovery. Administration into the subarachnoid space of 0.3 ml of blood as six aliquots was accompanied by a step-like rise in ICP that reached a maximum value of approximately 50 mmHg and only slowly declined upon completion of the injection (Fig. 18). During the period of acutely raised ICP, pressure-waves were commonly observed, these occurred at a frequency of about 8 per minute and were usually closely related to similar variations in the arterial pressure. Also during this period, an increased cerebrospinal fluid pulse pressure was often evident. In those animals in which 0.3 ml of buffered saline was administered in place of blood, the ICP attained roughly the same immediate peak values, but fell rapidly towards control values both between individual aliquots and upon final completion of injection. The subsequent time course of ICP recovery differed also between the groups (Fig. 19). Following experimental SAH the ICP remained elevated for up to 3 h at a pressure of 16 mmHg compared with a normal value of only 4 mmHg in the rat. At 24 h the ICP was still mildly elevated at 8 mmHg, and it only fully returned to control levels after 48 h. In the saline-injected animals the ICP normalised far more rapidly, falling to 8 mmHg by 15 minutes and returning fully to
control values by 2 h. There were no significant changes in the ICP of sham-operated animals. No formal analysis of ventricular volumes was performed in this study, however on sectioning the brains of animals used in delayed ICP studies it was observed that hydrocephalus occurred in animals 24 h post-SAH, but not following control procedures.

**MABP and CPP Changes (Fig. 19)**

In both the SAH and saline-injected animals there was a rapid rise in MABP following the injection of an 0.3 ml volume load into the subarachnoid space via the cisterna magna. The pressure reached a mean peak value of around 125 mmHg then fell slowly after 3 minutes, recovering to normal values by 3 h in both groups of animals.

CPP fell immediately by approximately 20 % following either SAH or saline-injection as the rise in MABP produced failed to match the peak ICP that resulted. In the SAH animals, CPP then recovered to normal within 3 minutes only to fall again by 15 % at 1 to 3 h post-SAH as ICP remained elevated in the presence of a now normal MABP. In the saline-injected animals, as ICP returned rapidly to normal values, the CPP after its initial fall became elevated by 20 % only to then fall in parallel with the recovery in MABP. In both SAH and saline-injected animals the MABP and CPP were maintained subsequently at normal levels from 24 h onwards. There were no significant changes in the MABP or CPP of sham-operated animals.
Other Physiological Parameters

The blood gases and haematocrit for the experimental groups at different times are given in Table 3. These parameters remained largely constant and approximate values were as follows: the haematocrit 45 %, the $P_a CO_2$ 37 mmHg, the $P_a O_2$ 139 mmHg and the pH 7.4. There were no significant differences in these values between the experimental groups.
7.5 DISCUSSION

A significant global reduction in CBF during the period acutely following experimental SAH has been demonstrated in animal studies performed in the monkey, dog, cat (Petruck et al., 1972; Brawley et al., 1968; Umansky et al., 1983) and more recently in the rat (Solomon et al., 1985). However details of the subsequent time course to recovery of such alterations in CBF, and of any subsequent development of delayed ischaemic changes have been described far less commonly. In the monkey, persistent reductions in CBF have been demonstrated up to 3 days post-SAH (Jakubowski et al., 1982; Sahlin et al., 1987), while in the rat a rather uniform 20% reduction in CBF has been reported to occur at 2 days post-haemorrhage (Delgado et al., 1986). The latter authors, when previously reporting their angiographic findings following experimental SAH in the rat, described a biphasic pattern of vasospasm with a maximum acute spasm at 10 minutes and a maximum late spasm at 48 h. By contrast, another earlier study (Barry et al., 1979) documented only an immediate onset of spasm which recovered by day 3 and was followed by vasodilatation from day 5 onwards. The rat has also been used to study delayed changes in cerebral metabolic activity following experimental SAH, one group (Solomon et al., 1987) reporting a significant decrease in metabolic activity of < 34% at 48 h while conversely another found a 30% increase (Delgado et al., 1986).

The measurements of CBF obtained in our study clearly demonstrate the occurrence of an acute decrease in mean cortical flow of approximately 50% for up to 3 h following
haemorrhage. A smaller, but still statistically 
significant, reduction in CBF persisted up to 24 h post-SAH. However, at 48 h during the period when a second lesser 
phase of vasospasm has been described, the mean cortical CBF 
was now fully recovered to normal values. Such a lack of 
absolute correlation between the time course of a reduced 
CBF and the previously described angiographic changes in the 
rat is not altogether surprising. Studies upon SAH in humans 
have shown the precise relationship of vasospasm to impaired 
CBF to be uncertain. Many investigators have found the 
presence of cerebral vasospasm in patients, to be associated 
with a reduced CBF both as a global and focal phenomena 
(Heilbrun et al., 1972; Ischii, 1979; Voldby et al., 1985b). Other authors however, have shown a less well-defined 
relationship (Bergvall et al., 1973; Grubb et al., 1977; 
Mickey et al., 1984).

Several patient studies have also shown a relationship 
post-SAH between hydrocephalus, raised ICP and reduced 
rCBF (Ferguson and Harper, 1971; Hartmann et al., 1977; 
Ischii, 1979). Similarly, hydrocephalus and vasospasm 
following SAH have been found to be significantly associated 
(Black, 1986), and the therapeutic benefits from ventricular 
drainage in cases of raised ICP, upon CBF and clinical grade 
have been well described (Kusske et al., 1973; Suzuki et 
al., 1974; Hartmann, 1980). A finding from this experimental 
chapter has been to emphasise the differences in ICP 
response following the administration of either blood or 
saline into the rat cisterna magna. Injection of 0.3 ml 
saline caused only a short-lived elevation of ICP to a peak
of approximately 50 mmHg. In contrast, an equal volume of released blood produced the same peak ICP, but the pressure then fell only slowly; remaining elevated four-fold at 3 h, approximately twice normal at 24 h and recovering to normal at 72 h. Concomitant with these changes in ICP, pressure-wave phenomena were observed following SAH, but not after control procedures. These waves most closely resembled Traube-Hering-Mayer related "C" waves, were seen only when the pulse-pressure amplitude was increased in the presence of an elevated ICP and are almost certainly evidence of a greatly reduced intracranial compliance post-SAH.

The changes in CBF that were observed following experimental SAH in the rat cannot simply be explained on the basis of concurrent changes in ICP and CPP alone. In the saline-injected control animals, despite 20% fluctuations in CPP, the CBF remained relatively constant, while in the SAH animals the maximum sustained reduction in CPP was 15%, which, even in the presence of impaired autoregulation, would not alone account for the 50% decline in CBF that took place.

The time course of reduced CBF in the rat does correlate well however with previously made observations (Barry et al., 1979; Jackowski et al., 1989b) that subarachnoid-released blood diminishes and is largely resorbed within 48 h in the rat. This suggests that the reduction in cerebral flow that occurs is closely related to the actual presence of blood and blood breakdown-products within the perivascular CSF spaces themselves. Acting either
directly upon the cerebral vessel wall or perhaps more indirectly via perivascular nerves and central brainstem afferent connections (Svendgaard et al., 1985; Solomon et al., 1987; Jackowski et al., 1989b) to produce an early phase of vasospasm. Although from this study it now appears that significant delayed cerebral ischaemia fails to develop in the rat after haemorrhage, this model still remains a cheap and readily reproducible one, demonstrating many of the pathophysiological changes that occur shortly following SAH in man. Use of the rat for further investigative studies upon the mechanisms arising out of, and treatment of SAH, would appear to be relevant provided the observed differences in the CBF response compared to that obtained in the clinical situation remains recognised.
Fig. 17. Percentage changes in rCBF following SAH, saline-injection, or sham-operation. Values are the means ± SEM, (—●—, parietal cortex; —○—, occipital cortex; —▲—, cerebellar cortex). The x-axis is the log scale of time in hours.
Fig. 18. Representative arterial (upper trace) and intracranial pressure (lower trace) recordings from experimental animals. (A) During and immediately following an 0.3 ml (6 X 0.05 ml aliquots) cisternal saline-injection. (B) During and immediately following an 0.3 ml (6 X 0.05 ml aliquots) SAH. (C) During induction of a SAH in an animal that demonstrates well the pressure-wave phenomena frequently observed.
Fig. 19. Sequential changes in MABP, CPP and ICP in the three experimental groups of animals. values are means ± SEM, (–•–, SAH animals; –○–, saline-injected animals; –▲–, sham-operated animals). The x-axis is the log scale of time in hours. (C - pre-injection values, P - values at time of peak ICP).
Table 2. Global CBF* as a percentage of control flow following cisternal injection of 0.3 ml blood, buffered saline or sham-operation.

<table>
<thead>
<tr>
<th>t</th>
<th>sham(1)</th>
<th>saline(2)</th>
<th>P</th>
<th>SAH (3)</th>
<th>P</th>
<th>P</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td>(1-2)</td>
<td></td>
<td>(1-3)</td>
<td>(2-3)</td>
</tr>
<tr>
<td>control</td>
<td>100 ± 0</td>
<td>100 ± 0</td>
<td>-</td>
<td>100 ± 0</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>3 min</td>
<td>105 ± 3</td>
<td>101 ± 6</td>
<td>ns</td>
<td>58 ± 2</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>15 min</td>
<td>103 ± 3</td>
<td>100 ± 4</td>
<td>ns</td>
<td>50 ± 2</td>
<td>***</td>
<td>***</td>
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<td>30 min</td>
<td>93 ± 3</td>
<td>95 ± 4</td>
<td>ns</td>
<td>51 ± 2</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>60 min</td>
<td>89 ± 3</td>
<td>87 ± 3</td>
<td>ns</td>
<td>55 ± 3</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>120 min</td>
<td>87 ± 4</td>
<td>80 ± 3</td>
<td>ns</td>
<td>55 ± 3</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>180 min</td>
<td>84 ± 5</td>
<td>79 ± 5</td>
<td>ns</td>
<td>57 ± 4</td>
<td>**</td>
<td>**</td>
</tr>
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<td>1 day</td>
<td>102 ± 3</td>
<td>104 ± 4</td>
<td>ns</td>
<td>85 ± 7</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>2 days</td>
<td>104 ± 4</td>
<td>111 ± 2</td>
<td>ns</td>
<td>98 ± 4</td>
<td>ns</td>
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<td>102 ± 3</td>
<td>106 ± 4</td>
<td>ns</td>
<td>108 ± 3</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>5 days</td>
<td>104 ± 3</td>
<td>112 ± 1</td>
<td>ns</td>
<td>107 ± 3</td>
<td>ns</td>
<td>ns</td>
</tr>
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</table>

Values are means ± SEM, n = 4 - 6, (\(^* \) pooled regional flow percentages).

***  P < 0.001
**   P < 0.01
*    P < 0.05
ns   nonsignificant
Table 3. Physiological measurements in the experimental groups (mean ± SEM).

<table>
<thead>
<tr>
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<th>Arterial blood gases</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Haematocrit (%):</td>
<td>PCO₂ (mmHg):</td>
<td>PO₂ (mmHg)</td>
<td>pH</td>
</tr>
<tr>
<td>sham-operated</td>
<td></td>
<td></td>
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<tr>
<td>control</td>
<td>45 ± 1</td>
<td>38.4 ± 1.6</td>
<td>136 ± 2</td>
<td>7.38 ± 0.02</td>
</tr>
<tr>
<td>3 min</td>
<td>37.5 ± 1.8</td>
<td>139 ± 4</td>
<td>7.43 ± 0.02</td>
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</tr>
<tr>
<td>15 min</td>
<td>36.6 ± 2.2</td>
<td>141 ± 4</td>
<td>7.42 ± 0.03</td>
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<tr>
<td>30 min</td>
<td>36.1 ± 1.6</td>
<td>139 ± 6</td>
<td>7.44 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>60 min</td>
<td>38.4 ± 1.1</td>
<td>139 ± 3</td>
<td>7.43 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>120 min</td>
<td>36.3 ± 0.8</td>
<td>140 ± 3</td>
<td>7.41 ± 0.02</td>
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</tr>
<tr>
<td>180 min</td>
<td>43 ± 1</td>
<td>35.9 ± 2.2</td>
<td>141 ± 7</td>
<td>7.39 ± 0.02</td>
</tr>
<tr>
<td>1 day</td>
<td>45 ± 1</td>
<td>37.6 ± 1.5</td>
<td>138 ± 4</td>
<td>7.39 ± 0.03</td>
</tr>
<tr>
<td>3 days</td>
<td>44 ± 1</td>
<td>37.1 ± 2.0</td>
<td>139 ± 3</td>
<td>7.37 ± 0.02</td>
</tr>
<tr>
<td>5 days</td>
<td>45 ± 1</td>
<td>38.4 ± 1.7</td>
<td>143 ± 5</td>
<td>7.42 ± 0.02</td>
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<tr>
<td>saline-injected</td>
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</tr>
<tr>
<td>control</td>
<td>46 ± 1</td>
<td>38.9 ± 1.0</td>
<td>141 ± 3</td>
<td>7.43 ± 0.02</td>
</tr>
<tr>
<td>3 min</td>
<td>37.4 ± 1.4</td>
<td>132 ± 3</td>
<td>7.45 ± 0.02</td>
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8. EXPERIMENTAL CHAPTER VI:
ULTRASTRUCTURAL CHANGES IN MAJOR CEREBRAL ARTERIES AND THEIR PERIVASCULAR CSF SPACES FOLLOWING EXPERIMENTAL SUBARACHNOID HAEMORRHAGE IN THE RAT
Summary: The ultrastructural changes that develop in major cerebral arteries, and the mechanism by which the perivascular subarachnoid CSF-containing spaces are cleared of blood following subarachnoid haemorrhage (SAH), has been examined in the rat. Scanning and transmission electron microscopy studies were performed upon vessels taken from animals 15 min to several days following experimental SAH induced by the injection of autologous blood into the cerebrospinal fluid space of the cisterna magna. At 48 h, and to a lesser extent 24 h post-SAH, the development of myonecrosis affecting a number of smooth muscle cells of the media was observed, largely confined to the immediate subintimal layer. However, no significant pathological changes in the endothelial layer or other components of the intima were seen, and the structure of the perivascular nerve plexus of the adventitia was likewise unaffected. Within 24 h of haemorrhage only a limited degree of phagocytosis of erythrocytes, by pial lining cells, took place. Early on the second day post-SAH however, a dramatic increase in the numbers of subarachnoid macrophages occurred, that derived primarily from a transformation of cells of the pia-arachnoid. This period was accompanied by a burst of intense phagocytotic activity; erythrocytes, fibrin and other debris being largely cleared over the next 24 h. At 5 days post-SAH, the subarachnoid macrophage population declined, with cells losing their mobile active characteristics and flattening out to assume a typical pia-arachnoid cell appearance once more. This chapter reveals that a mild vasculopathy afflicting cerebral
arteries, and a phase of pronounced macrophage phagocytic activity, are both most marked at around 48 h following SAH in the rat. These two findings may be of further interest, inasmuch as it is also around this time that a phase of delayed cerebral arterial narrowing has been previously documented.
8.1 INTRODUCTION

A major cause of potentially preventable mortality and morbidity in patients following aneurysmal subarachnoid haemorrhage (SAH) is the cerebral ischaemia that has been associated with the delayed development of cerebral arterial luminal narrowing (Heilbrun et al., 1972; Ischii, 1979; Mickey et al., 1984). Studies of human cerebral vessels obtained at operation and post-mortem in such patients, have revealed the early occurrence in narrowed arterial segments, of myonecrosis affecting a number of the smooth muscle cells of the media. This is followed in some later cases by a phase of intimal thickening and subintimal proliferation (Crompton, 1964; Conway and McDonald, 1972; Hughes and Schianchi, 1978; Peerless et al., 1980; Smith et al., 1985). Pathological studies in experimental SAH, have taken place mainly in larger animals such as the monkey and dog, where some evidence for similar changes arising within the vessel wall have also been described (Tanabe et al., 1978; Peerless et al., 1980; Espinosa et al., 1984b). SAH both in man and from experimental studies has additionally been shown to result in the development post-haemorrhage of a dysfunction of the cerebrovascular sympathetic nerves, with alterations in the levels of their contained neurotransmitters (Lobato et al., 1980; Edvinsson et al., 1982; Tsukahara et al., 1988; Jackowski et al., 1989b). Although whether these latter findings represented the development of actual structural damage to the neural plexus itself, or merely changes in neurochemical expression, has not hitherto been established.
In humans, the release of blood into the subarachnoid spaces is commonly associated with a reduced cerebrospinal fluid (CSF) resorption, and may lead in some cases, to the development of symptomatic communicating hydrocephalus with raised intracranial pressure. The mechanism by which released blood is cleared from the cisternal, perivascular and other CSF spaces following SAH has been uncertain. In the remainder of the body, macrophages play a major role in the removal of extravasated blood, but in the subarachnoid spaces of normal animals, cells resembling macrophages have only rarely been described (Wislocki, 1932). Following the introduction of bacteria or foreign proteins in the CSF spaces, clearance occurs by phagocytosis, initially by leptomeningeal cells and subsequently by cells that resemble activated macrophages (Nelson et al., 1962; Shabo and Maxwell, 1971). The origin of these subarachnoid macrophages has been the subject of much controversy. Essick (1920) first proposed that such cells arose directly from the pia-arachnoid itself, a suggestion supported by experimental findings in the dog, (Merchant and Low, 1979). Others however, have presented evidence for a haematogenous source of origin of leptomeningeal macrophages (Koningsmark and Sidman, 1963; Morse and Low, 1972).

This study was undertaken, firstly to investigate if similar changes to those previously seen in the cerebral vessels of man and some larger animals post-SAH, also developed following experimental haemorrhage in a small animal model such as the rat. Secondly, it was hoped to establish the mechanism by which released blood is
subsequently cleared from the perivascular and cisternal subarachnoid spaces surrounding the major cerebral arteries.
8.2 MATERIALS AND METHODS

Animal preparations

Adult male Sprague-Dawley rats weighing 350-450 g were used for the experiments. Subarachnoid heamorrhage was induced in 36 animals anaesthetised with a mixture of 70% nitrous oxide, 30% oxygen, to which 1.5% halothane was added. Animals were allowed to breathe spontaneously at all times. Autologous arterial blood (0.3 ml) freshly withdrawn from a femoral arterial catheter and replaced with an equal volume of normal saline was injected via a 30 gauge needle into the cisterna magna. Control animals comprised a group of 24 animals in which an equal volume of buffered normal saline (25 mEq/l of NaHCO₃ added and equilibrated at 37°C with a gas mixture of 5% CO₂/95% O₂ to pH 7.4) was injected in place of blood, and a further group of 24 animals in which a sham operation only was performed, with no injections given. Animals were subsequently killed and examined at times 15 min, 1 h, 3 h, 1 day, 2 days, 3 days, 5 days and 7 days post-SAH.

Perfusion-fixation and tissue examination

Under inhalational anaesthesia, the heart was exposed and a cannula inserted through the left ventricle into the ascending aorta. After incising the right atrium, the whole animal was perfused with approximately 200 ml of fixative composed of 4% paraformaldehyde, 0.1% glutaraldehyde, in 0.1 M phosphate buffer, pH 7.3, and followed by 500 ml of the same fixative minus glutaraldehyde. The brains were removed shortly after perfusion-fixation and immersed in the same fixative for 2 h, then placed in 0.1 M phosphate buffer
overnight. Under the operating microscope, the circle of Willis was then dissected from the brain and the proximal segment of the basilar artery and a distal segment of the internal carotid artery were taken and prepared for transmission (TEM) and scanning (SEM) electron microscopy. After washing in 0.1 M phosphate buffer, the specimens were post-fixed in 1% osmium tetroxide in 0.1 M phosphate buffer for 30 min then rinsed in distilled water. Specimens for TEM were next stained in 1% aqueous uranyl acetate for 30 min, then dehydrated through graded alcohols into propylene oxide and flat embedded in araldite. Ultrathin sections were cut and mounted on 200-mesh grids to be stained with aqueous uranyl acetate followed by lead citrate and examined with a Philips-300 electron microscope. Specimens for SEM, after post-fixation in osmium tetroxide, were further prepared, both conventionally and following an acid-etch treatment using 8N HCl (Fujiwara and Uehara, 1984), the latter process allowing an improved visualisation of the perivascular neural plexus and vessel wall three-dimensional structure. Specimens, after dehydration in graded alcohols, were taken through amyl acetate, critical-point dried in a Balzers Union CO$_2$ bomb, mounted on SEM stubs, sputter coated with gold using a Polaron E 5000 unit and viewed with a Hitachi S-530 microscope.
8.3 RESULTS

Endothelium

Scanning electron microscopy revealed the intimal endothelium of arteries taken from animals following experimental SAH, as being largely similar in appearance to the endothelium taken from control animals (Fig. 20). A well defined polygonal pavement-like layer of endothelial cells was always present on the luminal surface, with the long axis of the cells being orientated in the direction of blood flow. Acutely (up to 3 h) following SAH, the endothelial cells tended to be more rounded and possess an increased number of bleb-like and microvillus surface projections compared to those from control animals, although no other evidence of epithelial cell loss or damage was apparent at any time. By 1 day following experimental SAH, the endothelial cell appearance was once again identical to controls.

Media

A striking feature was the development 1 day post-SAH of a small but significant number of smooth muscle cells, that displayed an overall increased electron density, contained dense-bodies, degenerate mitochondria, and showed loss of their normal internal structure (Fig. 21A). By 2 days the sarcolemma of such necrotic cells had broken down, with the appearance of aggregated myofilament complexes, free ribosomes and membrane debris, lying freely in the extracellular space between smooth muscle cells (Fig. 21B). These changes were most commonly observed in, although not wholly confined to, the medial layers closest to the
Perivascular nerves and adventitia

The schwann-cell/neural plexus was most readily visualised in the acid-etched SEM preparations (Fig. 22). No differences were apparent between experimental animals and controls, either in the structure of the plexus itself or in the extent of its distribution within the adventitial layer, at any of the times studied.

Subarachnoid spaces

Immediately following SAH, the perivascular and cisternal subarachnoid spaces adjacent to the basilar and internal carotid arteries were occupied by a dense blood-clot that consisted largely of erythrocytes trapped within a fibrin mesh (Fig. 23A). Little change was detected on the first day, although occasionally erythrocytes were seen that appeared to have been directly phagocytosed by cells of the pial membrane. Early on the second day post-SAH, large numbers of rounded cells became evident. Fully matured forms displayed a variety of ruffled membranes and microappendages, and possessed the characteristics of subarachnoid macrophages. This emergent cell population appeared to originate principally from the pial, and to a lesser extent the arachnoid membranes. Within these structures, transition zones appeared (Fig. 23B), from which intermediate cell types could be seen in various stages of transformation, initially rounding up then gaining microappendages upon their cell surface. Defects in the normally continuous sheet of overlapping pial cells became
evident, as maturing cells migrated out into the surrounding subarachnoid blood-clot. These cells behaved subsequently in a manner characteristic of activated macrophages, and could be seen avidly phagocytosing erythrocytes and other debris from the subarachnoid spaces (Fig. 24). Already by 3 days post-SAH, the perivascular CSF spaces were largely cleared of blood cells and fibrin. At 5 days post-SAH the macrophage population declined and sites were increasingly observed upon the pia-arachnoid membranes and arachnoid trabeculations, where subarachnoid macrophages appeared to be losing their mobile active characteristics, flattening out to resemble typical pia-arachnoid cells once more (Fig. 25).
8.4 DISCUSSION

The development of a delayed narrowing of major cerebral arteries following SAH due to aneurysmal rupture, has been known since the original angiographic description by Ecker and Riemenschneider (1951). The time course of this constriction in man has been detailed (Kwak et al., 1979), its onset is usually delayed until the fourth or fifth day post-bleed and has a maximum incidence between 10 and 14 days. Experimental studies in the dog (Brawley et al., 1967; Kuwayama et al., 1972; Nagai et al., 1974) revealed that cerebral arteries respond to SAH in a biphasic manner, with an acute reversible phase of narrowing immediately post bleed, followed in some instances by a more persistent phase that develops after a variable delay.

Study of such angiographically observed narrowed segments of cerebral arteries, taken either at operation or at autopsy from patients dying following SAH, have demonstrated the occurrence of varying degrees of medial necrosis, subintimal proliferation and other pathological changes (Crompton, 1964; Conway and McDonald, 1972; Hughes and Schianchi, 1978; Peerless et al., 1980; Smith et al., 1985). Whilst investigators of experimental SAH in the monkey have documented similar morphological changes (Peerless et al., 1980; Espinosa et al., 1984b), studies in dogs have been more variable, some authors observing pathological changes in the walls of cerebral vessels (Tanabe et al., 1978; Tani et al., 1978) while others have found few or none at all (Pickard et al., 1985).

Recently the laboratory rat has been increasingly
employed for studies of experimental SAH (Barry et al., 1979; Lacy and Earle, 1982; Svendgaard et al., 1985; Solomon et al., 1987), its relative cheapness and greater availability allowing its use in larger animal numbers. A biphasic time-course of narrowing in cerebral vessels post-SA has been reported in the rat (Delgado et al., 1985), but no pathological changes of rat cerebral vessels after haemorrhage described. In the present study, we have now found evidence for similar changes to those previously described in larger animal models. Myonecrosis of limited but significant numbers of smooth muscle cells of the media of cerebral vessels was observed, that commenced the day following haemorrhage. A number of studies, both of man and in a variety of animals including the rat, have now clearly demonstrated that a dysfunction of both endothelial cell and perivascular nerve related cerebrovascular reactivity develops post-SA (Pickard and Perry, 1983; Sahlin et al., 1985; Nakagomi et al., 1987; Kim et al., 1988a). In the present study, the ultrastructural morphology of both the endothelium and neural plexus appeared to be largely unaffected following haemorrhage. This would suggest therefore, that these alterations possibly reflect either impaired smooth muscle responsiveness, or qualitative changes in endothelial cell and nerve function, rather than any physical damage to the latter structures themselves.

Clearance of released blood from the subarachnoid spaces in man can take from 6 to 30 days (Tourtellotte et al., 1964), the continued presence of blood clot within these spaces predisposes to impaired CSF resorption,
hydrocephalus, raised intracranial pressure and delayed cerebral ischaemia (Ellington and Margolis, 1969; Vassilouthis and Richardson, 1979; Butler et al., 1980; Mohsen et al., 1984). In man, most erythrocytes rapidly become emeshed within the arachnoid trabeculae of the subarachnoid spaces and it was naturally hypothesised that they were then removed by lysis and phagocytosis (Sprang, 1934; Hammes, 1944). Early studies of erythrocyte clearance in animals however, seemed to suggest that cells might instead be largely removed intact via perineural lymphatic-type channels (Kennady, 1967; McQueen et al., 1974). Separate studies had suggested a system of tubular spaces and pores across the arachnoid villi (Welch and Friedman, 1960) and it was also proposed that whole erythrocytes might pass directly into the blood stream via this route (Simmond, 1953). Ultrastructural studies have since shown the endothelium of the arachnoid villi to be continuous, thereby preventing such a mode of clearance (Alksne and White, 1965). More recently, studies in the dog (Alksne and Lovings, 1972) have suggested that removal of erythrocytes from the subarachnoid spaces occurred mainly by lysis and possibly by phagocytosis of resultant debris by cells of the arachnoid villi. Again in the dog, an increase in the numbers of free subarachnoid macrophages was also shown to occur after SAH by Julow et al. (1979), who observed limited erythrophagocytosis to take place.

In this investigation of the mode of clearance of the perivascular subarachnoid spaces in the rat, erythrocytes immediately following haemorrhage, occasionally underwent
phagocytosis by cells of the pial membrane. With increased survival time, there followed a period characterised by a burst of intense phagocytic activity by mobile cells possessing the typical features of activated macrophages. This greatly increased population of subarachnoid macrophages appeared to be largely derived by the transformation of cells from the pial membrane. Upon clearance of blood-clot from the perivascular subarachnoid spaces, these activated macrophages flattened out, assuming the characteristics of typical pia-arachnoid cells once more.

The development of pathological changes affecting the media of the rat basilar artery and a period of intense macrophage erythrophagocytic activity within perivascular subarachnoid blood-clot, were both maximal at around 48 h post-haemorrhage. Interestingly, this period also coincides with the time at which a delayed onset of basilar arterial luminal narrowing has previously been described (Delgado et al 1985). Macrophage activity and oxyhaemoglobin autoxidation are both well known to be associated with the generation and release of free radicals (Sutton et al., 1976; Babior, 1978). The findings from this experimental chapter may be of further interest therefore, inasmuch as work by a number of authors now suggests that an excess of free radicals may be involved in the genesis of both arterial wall damage and the angiographically visible delayed cerebral arterial narrowing that may follow SAH (Sakaki et al., 1988; Steinke et al., 1989; Zuccarello et al., 1989)
Fig. 20. Scanning electron micrographs of the luminal aspect of basilar arteries of the rat following perfusion-fixation at physiological pressure. A: Sham-operated animal, 1 h post procedure. B: SAH animal, 1 h post-haemorrhage. C: SAH animal, 2 days post-haemorrhage. In all preparations the endothelial cell outlines and overlapping folds can be readily identified. At 1 h, many cells can be seen to possess bleb-like and microvillus projections. Scale bar = 10 um. All micrographs have the same magnification.
Fig. 21. Transmission electron micrographs of the rat basilar artery demonstrating myonecrotic changes in the media following experimental SAH. A: Artery from an animal killed 24 h post-SAH. sm, normal smooth muscle cell; dm, degenerating smooth muscle cell displaying increased electron density, dense bodies and mitochondrial necrosis. Scale bar = 3 μm. B: Artery from an animal killed 48 h post-SAH. el, elastic lamina; Nm, remains of a necrosed smooth muscle cell in which the sarcolemma has largely disappeared. Note the myofilament complexes (cpx) and other cellular debris lying freely in the intercellular space. Scale bar = 1.5 μm.
Fig. 22. Acid-etch scanning electron photomicrographs of rat cerebral arteries, prepared so as to demonstrate the neural plexus (Np), taken from animals killed A: at 1 day and B: at 60 min post-SAH. In (A) the adventitial layer has been lifted off the vessel together with the attached neural plexus and is viewed from its inner aspect. Sh, Schwann-cell body with nuclear outline; Er, erythrocytes. Scale bar = 20 um. In (B) the vessel has been microdissected to reveal the intimal elastic lamina (in), medial smooth muscle (me), and adventitial (ad) layers together. The schwann-cell/neural plexus is now seen lying upon the smooth muscle cells of the media. Scale bar = 30 um.
Fig. 23. Scanning electron micrographs of perivascular subarachnoid spaces adjacent to the major cerebral arteries. A: At 60 min post-SAH the subarachnoid space is entirely filled with blood-clot (Bc). Pm, pia mater; Tr, arachnoid trabeculae; Ar, arachnoid mater. Scale bar = 35 um. B: At 48 h post-SAH, large numbers of rounded cells (arrowed ma), some possessing microappendages, are now present within the blood-clot. Zones (Tz) within the pia have appeared, where defects in the membrane are occupied by cells of intermediate forms to these rounded cells. Scale bar = 40 um.
Fig. 24. Scanning electron micrograph of the perivascular subarachnoid space 3 days post-SAH. At higher magnification, the rounded cells seen previously, display the ruffled membranes and microappendages characteristic of subarachnoid macrophages (Ma) and are seen actively engaged in erythrophagocytosis. Er, erythrocyte. Scale bar = 10 µm. Inset: Transmission electron micrograph of one such cell. Internally, these transformed pial cells display increased perinuclear cytoplasm and clear vacuoles. A pseudopod process is in close contact with an erythrocyte. Scale bar = 5 µm.
Fig. 25. Scanning electron micrograph of the pia mater 6 days post-SAH. Cells possessing macrophage features are still present (Ma). However, new zones of transformation (Tz) have appeared, where intermediate cell forms are seen in various stages, flattening out and assuming more typical pial cell morphology. Scale bar = 25 um.
9. CONCLUSIONS FROM THE STUDY

The findings can be summarised in the following way:

(1) A significant discrete serotonergic innervation of the major cerebral arteries in the rat, and possibly other species including man, does not appear to exist, contrary to earlier reports in the literature.

(2) Serotonin, when observed within the perivascular nerves of the large cerebral arteries, is found to occur only in nerves also identified as catecholaminergic sympathetics. When present, it appears to arise largely as a result of a direct uptake of the indole from the surroundings.

(3) After an induced SAH, serotonin was rapidly and readily taken up by the cerebrovascular sympathetic nerves, while a coincident depletion of NPY from these same nerves took place. Once blood clot had been largely resorbed from the CSF, the neurotransmitter content of these nerves was then restored once more to normal.

(4) Experimental SAH in the rat was found to result in an immediate 50% global reduction in cortical CBF that persisted for up to three hours post-haemorrhage. There was a smaller but significant reduction in flow 24 hours post SAH, while CBF was normal between day 2 and 5. The decrease in CBF was clearly most marked at about the same time at which a maximal acute spasm has been observed by other authors, using the same animal model. However, no later reduction in flow was seen at 48 hours, the time at which a delayed phase of vasospasm has previously been described in
the rat. The ICP was increased and the CPP decreased during the first 24 hours after the SAH. The recorded reduction in CPP of 15% was found not to be in accordance with a fall in CBF of 50%. It is suggested that early vasospasm secondary to released blood rather than pressure changes is responsible for the initial cerebral ischaemia observed.

(5) Ultrastructural studies of the cerebral arteries from rats subjected to SAH, revealed some, but not all of the morphological changes found in the vasculopathy associated with delayed cerebral arterial narrowing in man and large animal models after SAH. Subintimal myonecrosis was observed, especially at 48 hours after the SAH, while no intimal or subintimal proliferation developed subsequently. The findings of intact perivascular nerves post SAH are of interest, and suggest that the described disappearance of neurotransmitter content is due to a functional change in the nerve rather than to a chemical neurectomy.

(6) The primary mechanism by which the perivascular subarachnoid spaces were cleared of adjacent blood clot following haemorrhage was by phagocytosis. This took place largely on the second day post-SAH, a period characterised by intense macrophage phagocytic activity. The origin of these subarachnoid macrophages was studied, and it appears that they are derived by a transformation from cells of the pia-arachnoid membranes. In the context of these findings, it is perhaps relevant to note that such phagocytic activity is well known to be associated with the local release of free radicals. These have been strongly implicated by some
authors in one mechanism of cellular damage that may subsequently lead to delayed cerebrovasculopathy and arterial narrowing.

Secondary deterioration after SAH in man, other than from proven episodes of rebleeding, is commonly attributable to the onset of delayed cerebral ischaemia. The causes of which are probably multifactorial, but most certainly include: a reduced CBF, impaired cerebrovascular autoregulation, and raised intracranial pressure. The first of these three, has been closely linked with the development of arterial luminal narrowing affecting one or more of the major cerebral arteries. Pathological studies have suggested that such vessel narrowing may represent a reactive vasculopathy induced either by blood-clot directly, or in response to secondarily released products.

The results obtained from this present study indicate, that a simple single-haemorrhage rat model displays a number at least of the features that can arise following SAH either in man or in larger animals after SAH. In particular, the development of a global reduction in CBF, an elevated ICP with reduced intracranial compliance, alterations in neurotransmitter balance within the perivascular sympathetic nerves that normally contribute to autoregulation, and ultrastructural evidence of a delayed cerebrovasculopathy were noted. Although not responding in an entirely identical manner to that of the human response after SAH, the rat model of SAH is revealed to be a worthwhile one for future studies to further investigate and determine how best to manage the disordered pathophysiology that commonly arises
following SAH.
10. FINAL DISCUSSION

10.1 EXPERIMENTAL MODELS AND SAH

In vivo animal studies:

A wide variety of animal species have been utilised in experimental studies of subarachnoid haemorrhage. Baboons, monkeys, dogs, cats, rabbits and rats having all been variously employed for this purpose. The in vivo SAH created in these studies, has largely been simulated either by the subarachnoid injection of the animals own autologous arterial blood, or by inducing rupture of a vessel of the circle of Willis. A considerable number of such animal studies, in which the primary aim was purportedly to investigate the mechanisms causing delayed cerebral vasospasm, have in fact instead been largely concerned with examining the acutely vasoconstrictive effects of subarachnoid blood. This caveat aside, a wealth of useful information has now emerged from such laboratory studies.

Cerebral arterial narrowing - Vasospasm

Echlin (1965), was one of the first to examine the effects of experimental SAH, produced in monkeys by the single direct application of arterial blood close to a major cerebral vessel. He demonstrated, that when subarachnoid blood came into contact with large arteries, that a marked vasospasm (30 to 60 percent of prehaemorrhage diameter) developed almost immediately; an effect that he noted generally lasted 10 to 40 minutes but sometimes longer. Brawley et al (1968) subsequently made the important observation in his studies on the dog, that cerebral
arterial narrowing following experimental SAH generally resulted in a biphasic response. In adult mongrel dogs in which the SAH was produced by rupture of an anterior cerebral artery, resultant haemorrhage was initially associated with a directly observed acute phase of arterial constriction that lasted for less than 1 hour. This was followed by a delayed recurrence of vessel narrowing, that developed 3 to 24 hours later and became maximal on the third day post-SA. A similarly biphasic time-course was also observed, when SAH was instead produced in the dog by a single injection of autologous arterial blood into the cisterna magna (Kuwayama et al., 1972). These latter authors followed their animals by means of serial angiograms, and demonstrated a response pattern that consisted of an acute phase of constriction that had largely recovered by 2 hours. This was followed by a delayed onset of recurrent narrowing at 24 hours with the affected vessels subsequently recovering to their normal diameters by 7 days. A persistant arterial narrowing following experimental SAH, has subsequently been reported as occurring in other smaller animal species. In the rabbit (Edvinsson et al., 1982) and in the rat (Barry et al., 1979), reductions in basilar artery diameters have been shown to be present for periods of up to 9 days and 48 hours post-haemorrhage respectively. Delgado and coworkers (1985) have performed angiographic studies in the rat, both acutely and at different time points up to one week following the SAH produced by a single intracisternal blood injection. In this small animal study, a biphasic time-course, similar to that previously reported
in the dog, was also found. An early phase of marked luminal narrowing (40 percent of prehaemorrhage diameter) was seen at 10 minutes, and had largely recovered by 90 minutes. This was then followed by a second but less marked constriction (25 percent of prehaemorrhage diameter) at 24 and 48 hours, which subsequently resolved fully by 72 hours.

**CBF, Autoregulation and Cerebral Metabolism**

The acute effects of subarachnoid haemorrhage upon cerebral blood flow, autoregulation, and reactivity to pCO₂ have been widely investigated in experimental models, whilst delayed effects have received somewhat less detailed attention. In adult rhesus monkeys, Petruk et al. (1972) measured the CBF by a Xenon¹³³ clearance technique and showed a significant acute (up to 3 hours) reduction in CBF in 12 out of 14 monkeys subjected to SAH. They also reported finding a close correlation between the degree of impaired flow and severity of neurological deficit. A number of authors have similarly studied the acute changes in CBF that followed experimental SAH in baboons (Jakubowski et al., 1982; Kamiya et al., 1983; Kuyama et al., 1984). Haemorrhage being produced by snare transection of the posterior communicating artery. In these studies, CBF generally became rapidly reduced to 20% - 50% of normal and showed a variable pattern of recovery, depending upon the neurological grade of the animal. In some instances, persistant reductions in CBF of up to 18% at 3 days were observed (Jakubowski et al., 1982; Sahlin et al., 1987). In the cat, experimentally induced SAH, provoked either by a cisternal blood injection (Umansky et al., 1983) or by internal carotid artery
puncture (Trojanowski, 1984) gave rise to similar results. Both regional and global CBF's were found to fall at 10 minutes by 15 – 50% of control values, and these remained depressed for up to 3 hours posthaemorrhage. Soloman (1985) has examined the immediate effects only, of experimental SAH upon CBF in a smaller animal model, the rat, induced by a single intracisternal injection of blood. Measuring blood flow acutely using a labelled microsphere technique, he demonstrated a 40% decrease in CBF over a 60 minute period.

Assessment of autoregulatory function and CO₂ reactivity after experimental SAH has been carried out predominantly using the baboon (Jakubowski et al., 1982; Kamiya et al., 1983; Sahlin et al., 1987). In such studies it was found that autoregulation and cerebrovascular reactivity to CO₂ became depressed in a global fashion, both acutely and for up to 3 days following the induction of SAH.

Whilst in man it has been found that SAH is clearly associated with global reductions in cerebral metabolism, particularly in those patients in the poorest clinical grades, the few results obtained from animal studies are extremely inconsistent. Studies have shown either no change (Sahlin et al., 1987), or a diffuse increase (Delgado et al., 1986). Although, both Fein (1975) and Soloman (1987) separately found that metabolic activity in the monkey and in rats was globally decreased following experimental SAH.

**Blood Brain Barrier / Brain oedema**

Experimental studies in animal models, have revealed that the BBB breakdown observed following SAH, predominantly affects the blood arterial-wall barrier of major cerebral
arteries. Although the resultant changes may partly be explained by the sudden rise in ICP and arterial pressure that occurs post-SAH, they can also be provoked equally well by the presence of blood clot in the subarachnoid space alone (Peterson et al., 1983; Sasaki et al., 1985; Sasaki et al., 1986; Zuccarello et al., 1987).

An increased brain water content, with an accompanying increase in sodium content, is known to develop early following experimental SAH (Kamiya et al., 1983; Kuyama et al., 1984; Doczi, 1985). Interestingly, evidence has emerged to suggest that such brain swelling may be modified by vasopressin (Doczi, 1984), implying that brain water permability and electrolyte balance may in part be regulated by this centrally released neuropeptide.

Peripheral autonomic and Central nervous system changes

Following experimentally induced SAH in in vivo animal models, there is now both direct and indirect evidence that a comparable degree of peripheral sympathetic overactivity to that seen in man, similarly takes place. Elevated levels of plasma catecholamines develop, also an increased release of adrenal medullary catecholamines and enkephalins can be demonstrated (Offerhaus and Van Gool, 1969; Klein et al., 1986). Electrocardiographic abnormalities, similar in variety to those that are seen clinically in patients, have also been observed after experimental SAH (Estanol et al., 1977; Lacy and Earle, 1983). These appear to occur, as a direct result of the effects of a sustained high level of circulating catecholamines upon the myocardium and conducting tissues of the heart (Schenk and Moss, 1966).
As shown in man, again evidence exists from animal studies, performed in this instance in the rat, to show that an elevation of brain noradrenaline concentration develops after experimental SAH (Solomon et al., 1986). Further studies have shown that this elevation reflects an increased synthesis and release of the neurotransmitter, presumably arising from projection axons that originate from neurons of the locus coeruleus (McCormack et al., 1986).

A number of authors have examined the effects of experimental SAH upon the periarterial nerve plexus that surrounds and innervates the major intracranial arteries. These studies have consistently demonstrated the development of a marked reduction, to complete exhaustion, of noradrenaline stores in the nerve terminals about such vessels (Fraser et al., 1970; Peerless and Kendall, 1975; Svendgaard et al., 1977a). Another finding of significance has been that a reduced uptake of noradrenaline by cerebrovascular perivascular nerves, arises post SAH (Lobato et al., 1980; Edvinsson et al., 1982). Such vessels also display a state of denervation supersensitivity to exogenously applied noradrenaline (Duckles et al., 1977; Svendgaard et al., 1977b). A reduction in the neurotransmitter content does not appear to be solely confined to the noradrenergic system. More recently, in studies both upon the dog and in the monkey, it has been reported that a reduction in the numbers of intracranial perivascular nerves displaying VIP, SP and NPY immunoreactivity may also occur following haemorrhage (Hara et al., 1986; Uemura et al., 1987).
Of interest therefore, in view of these observed changes in cerebral noradrenaline content and perivascular neurotransmitter status, have been a series of experiments performed by Svendgaard and his coworkers. They have attempted to elucidate if any role is played by the centrally projecting serotonergic, dopaminergic and catecholaminergic neural pathways, in the development of vasospasm affecting the vertebrobasilar arteries following experimental SAH. Chemical destruction of the central serotonergic and dopaminergic pathways prior to the intracisternal injection of blood, apparently does not affect the occurrence of such vasospasm after SAH (Svengaard et al., 1986). These same authors report however, that the lesioning of ascending catecholaminergic fibres in the mesencephalon or medulla either by a stereotactic lesioning with 6-OHDA, or by neonatal systemic 6-OHDA administration, effectively prevents the development of vertebrobasilar artery spasm (Svendgaard et al., 1985; Delgado et al., 1987). These findings however, are difficult to reconcile with the fact that there is no known evidence for any central catecholaminergic innervation of the vertebrobasilar system. They have instead therefore proposed, an alternative neurohumoral mechanism, in which overactivity in central catecholaminergic projections to the hypothalamus might cause vasospasm of large cerebral arteries through secondary release of vasopressin. In support of this hypothesis, they have conducted experiments demonstrating that arginine-vasopressin deficient Brattleboro rats fail to develop acute vasospasm after SAH (Delgado et al., 1988).
Against this argument however, must be considered the fact that several other authors have been unable to demonstrate any consistent vasoconstrictor response of the cerebral vessels in response to vasopressin (Allen et al., 1974a; Lassoff and Alturs, 1980; Katusic et al., 1984).

In vitro studies:

These studies, employing vessels and CSF obtained either from patients or in animals following SAH, are all characterised by the investigation of any resultant changes in cerebrovascular responsiveness or pharmacology in an in vitro environment.

Such investigations, have been largely focused upon attempts to unmask a specific substance or series of substances, putatively contained within the CSF in increased amounts following haemorrhage, and which are held responsible for the production of vasospasm leading to cerebral ischaemia. It had been noted early on, that the application of CSF obtained from patients with angiographic evidence of vasospasm following SAH, caused isolated cerebral arteries to contract, as well as sensitising them to many other vasoconstrictor agents (Boullin et al., 1976). It was also found that such vasoconstrictor activity was highest in those patients who had evidence of severe angiographic vasospasm or who developed delayed ischaemic deficits and subsequently died (Boullin, 1980; Kaye et al., 1984).

Serotonin

One of the first substances to be extensively
investigated in this way was 5-hydroxytryptamine. In 1974 in a series of classic experiments, Allen (Allen et al., 1974a; Allen et al., 1974b; Allen et al., 1974c) using the canine basilar artery, examined the contractile activity of human CSF obtained following haemorrhage. Largely on the basis of these studies, it was at that time concluded that serotonin was the main agent responsible for the immediate arterial spasm that arose following vessel exposure to bloody CSF obtained post SAH. The subsequent finding that perianeurysmal CSF obtained at the time of operation for ruptured intracranial aneurysms (and shown to contain 5-HT) could be prevented from causing the constriction of isolated human intracranial arteries by concomitant application of the 5-HT antagonist ketaserin (Tagari et al., 1983), would appear to strongly support this argument. Although it remains a prime candidate for the cause of the acute vasospasm commonly seen immediately after experimental SAH in animal models, 5-HT is however, unlikely to alone be responsible for a more chronic vasospasm, or the delayed vasospasm seen in man. In vivo administration of serotonin into the CSF space in animals, indeed produces spasm, but this is short-lived, persisting for some 2-3 hours only (Allen et al., 1974c; Boullin et al., 1978). A point that has been made in man is that although 5-HT concentrations in the CSF will be considerably elevated immediately following release of blood into the subarachnoid space, that there is no consistently found elevation of levels subsequently, during the period of delayed arterial narrowing. These measurements have however, been made predominantly upon
ventricular CSF samples (Vapalahti et al., 1978; Voldby et al., 1982). More recently, evidence has emerged that suggests that significantly elevated concentrations of 5-HT may in fact persist for considerably longer in the CSF contained locally within periarterial cisterns (Tagari et al., 1983; Chehrazi et al., 1989).

Most in vitro studies upon the role of serotonin released from platelets following experimental SAH, have naturally focused upon its known vasoconstrictor actions. Of perhaps more relevance to the aetiology of the chronic arterial narrowing found in human SAH, may be its less well known, stimulatory actions upon vascular smooth muscle (VSM) mitogenesis and deoxyribonucleic acid synthesis. The mitogenic effects of serotonin and for that matter human platelet-derived growth factor (PDGF), have to date, primarily been studied in bovine aortic smooth muscle cells in culture but may well be relevant for VSM cells of the intracranial vessels following SAH. PDGF is another potent mitogen for VSM cells. It is released from the alpha-granules of platelets that have been exposed to vascular connective tissue, and its mitogenic response is synergistically affected by the concommitant presence of serotonin (Nemecek et al., 1986). Serotonin-induced VSM mitogenesis is mediated by specific receptors and is critically dependent upon guanine nucleotide binding proteins while that of the PDGF response is not (Kavanaugh et al., 1988).

Erythrocytes / Oxyhaemoglobin

Erythocyte products appear to be a necessary factor in
the development of a cerebral vasculopathy arising following experimental SAH (Duff et al., 1988). In vitro, the directly vasoconstrictor effects of either haemoglobin or haemolysate have been well described (Handa et al., 1980; Wellum et al., 1980; Boullin et al., 1983). Prolonged exposure to oxyhaemoglobin is known to modify the response of isolated cerebral vessels from various species, to a variety of agents. Relaxations induced by substance P released from vasodilator nerves, the electrogenic sodium pump, endothelium-derived relaxing factor, prostaglandin I$_2$, nitroglycerin, and angiotensin-II are all impaired following exposure to oxyhaemoglobin (Lee et al., 1984; Toda, 1988; Onoue et al., 1989). Conversely, oxyhaemoglobin potentiates both neurogenic vasoconstriction and 5-HT induced vasoconstriction of cerebral vessels (Lee et al., 1984; Tanaka and Chiba, 1988).

**Oxygen Free Radicals**

Active species of oxygen are liberated upon the conversion of oxyhaemoglobin to methaemoglobin (Misra and Fridovich, 1972; Sutton et al., 1976). They are also generated by the phagocytic activity of polymorphonuclear leucocytes, monocytes and macrophages (Babior, 1978). Both of these processes will naturally arise in intimate proximity to the walls of the major cerebral vessels, following a SAH. Any excessive formation of oxygen free radicals will tend to overwhelm the naturally occurring enzyme-defence mechanisms subserved by superoxide dismutase (SOD), catalase and glutathione peroxidase. Released free radicals will then be available to participate in harmful
oxidation reactions involving nearby cell structures. Unsaturated membrane fatty acids undergo lipid peroxidation, while DNA and proteins may also be irreversibly damaged, such changes if severe, leading to tissue death (Joyce, 1987). After induction of experimental SAH, an elevation in the concentration of lipid peroxides, both in the CSF and walls of major cerebral arteries, has been clearly demonstrated (Sakaki et al., 1988). Early attempts at protection against free radicals using treatment with a 21-aminosteroid with antioxidant properties, have suggested that the degree of vasospasm seen following experimental SAH may be moderated by such agents (Steinke et al., 1989; Zuccarello et al., 1989).

EDRF

Endothelium-dependent relaxation in response to ATP, thrombin, acetylcholine, and other substances, is now known to be significantly impaired in isolated arteries taken from animals following induced SAH (Nagakomi et al., 1987; Hongo et al., 1988; Kim et al., 1988a). In contrast however, the endothelium-dependent contractions in response to arachidonic acid, calicium ionophore A23187, hypoxia and mechanical stretch apparently remain intact (Kim et al., 1988a; Kim et al., 1988b). Measurements, both of normal basal release and of increased release in response to stimuli, have shown that endothelial cell production and release of EDRF is in fact unchanged following SAH (Kim et al., 1989). This would imply that somehow a reduced cerebrovascular smooth muscle responsiveness to EDRF develops, and is itself the primary cause of the impaired
relaxation responses that develop posthaemorrhage.

**Arachidonic acid metabolites**

Elevations in the levels of prostaglandins and leukotrienes within the CSF, have been reported in patients following aneurysmal SAH (Paoletti et al., 1988; Chehrazi et al., 1989). Both compounds are the products of arachidonic acid metabolism. Prostaglandins being derived via the cyclo-oxygenase pathway and acting primarily as vasoactive agents; while the leukotrienes formed via the lipoxygenase pathway modulate membrane permability, brain oedema formation and local inflammatory responses (Piper, 1983; Dempsey et al., 1986). Prostaglandin concentrations are also significantly raised in CSF and cerebral blood vessels taken from animals following an experimentally induced SAH (Maeda et al., 1981; Walker et al., 1983). Cyclo-oxygenase inhibitors are known to decrease the directly vasoconstrictive effects of whole blood and haemolystate upon isolated vessels (Okamoto et al., 1984; Lang and Maron, 1988). It has been proposed therefore, that an increased synthesis of vasoconstrictor prostaglandins by these vessels may be partly responsible for the blood-induced constrictor response observed in vitro. It has also been reported that cyclo-oxygenase inhibitors may attenuate cerebral vasospasm after experimental SAH (White et al., 1979; White and Robertson, 1983). Paradoxically however, the vasoconstrictor effect of CSF obtained from patients following recent SAH, upon isolated vessels, is enhanced in vessels preincubated with indomethacin (Brandt et al., 1981) suggesting that vasodilator prostacyclin synthesis tends to
predominate in the latter situation. Overall, the results obtained do suggest that an imbalance in the synthesis of contractile and relaxant arachidonic acid metabolites can arise in vessels following SAH, and that this may contribute to the genesis of arterial narrowing.
10.2 TOWARDS AN UNDERSTANDING OF CEREBROVASOSPASM AND DELAYED CEREBRAL ISCHAEMIA FOLLOWING SAH

What then, can we now say regarding the mechanisms that are likely to be responsible for the production of delayed cerebral ischaemia, and of the role of cerebral arterial narrowing or chronic 'vasospasm' in its genesis? Firstly, let us reconsider what is known both of acute and chronic cerebrovasospasm.

Acute vasospasm:

In vivo studies of SAH, in which experimental haemorrhage has been induced by various means in animal models, reveal under such circumstances the universal occurrence of an immediate but transitory phase of reversible physiological cerebral arterial narrowing. Similarly, the application of fresh arterial whole-blood or oxyhaemoglobin in vitro, causes an immediate contractile response in isolated cerebral vessels taken from many species, including it should be noted those of man. Although an immediate phase of short-lived vasoconstriction has on occasions been observed to arise in humans acutely after SAH, a truly biphasic timecourse of vasospasm as described from animal studies, does not appear to be a common occurrence in man.

The major factors thought responsible for the generation of an acute physiological constriction of cerebral arteries after haemorrhage have been discussed earlier, and their interactions are now summarised in Figure 26.

Might it be possible either that a marked acute physiological contraction or state of intense sympathetic
ACUTE VASOSPASM (physiopharmacological)

Arterial blood release into CSF

Platelets
- Dense granule release
  - 5-HT
    - Perivascular sympathetic release of NA / NPY / ATP
      - Direct 5-HT₂ effect

Leucocytes

Erythrocytes
- Oxyhaemoglobin

Potentiates

Direct effect (? prostaglandin dependent)

Inhibits EDRF + vasodilator nerves

CEREBRAL ARTERIAL CONTRACTION
overactivity, could in any way contribute to later vessel wall damage and delayed chronic cerebral arterial narrowing; mediated perhaps via the mechanism of excitotoxic neurotransmitter release? High levels, both of CSF and circulating catecholamines are quite commonly observed in man after SAH, with concommitant depletion of neurotransmitters from within the terminals of cerebrovascular nerve fibers. It has certainly been observed experimentally that production of an intense arterial constriction by the application of such catecholamines in vitro, can result subsequently in the development of recognisable vessel wall damage (Alksne and Greenhoot, 1974; Joris and Majno, 1981).

**Chronic vasospasm:**

Chronic cerebrovasospasm is usually only manifested at about the fourth day after haemorrhage, it is maximal in incidence in the second week and largely gone by the end of the third week. Despite the application of a quite extensive variety of pharmacological agents, such delayed vasospasm, unlike acute vasospasm, has proven remarkably resistant to all attempts aimed either at preventing or producing lasting reversal of luminal narrowing. In the introduction to experimental chapter VI, evidence was presented obtained both from human studies and from animal models including this one, that strongly suggests that delayed cerebral arterial luminal narrowing represents the development of structural pathological changes in the vessel wall itself, rather than any contractile activity of the cerebrovascular smooth muscle per-se. The occurrence of vessel wall changes
in such cases, is evidenced by the development in narrowed arteries of an initial relatively early phase of patchy necrosis of smooth muscle cells involving the subintimal media. This is followed in more severe instances, by one in which proliferative intimal and subintimal changes predominate, typically, with subendothelial intimal fibrosis, medial necrosis and myofibroblast proliferation. A number of causal mechanisms, that are considered highly likely to be involved in the aetiology of this cerebral arterial vasculopathy have been examined earlier, and are again detailed in Figure 27.

Any hypothesis, that seeks to answer how the mechanism of injury to the vessel wall takes place, must account for the known period of delay before its appearance, and be satisfactorily explained by release of factors known either to already be present in or that can be generated by the subarachnoid blood clot. Current theories would appear to implicate cellular damage induced by oxygen free radicals, released into the subarachnoid spaces as a result of oxyhaemoglobin autoxidation and macrophage phagocytosis. A largely unexplored avenue of research at present, is the possibility of a direct involvement of vascular smooth muscle mitogenic factors released by platelets such as PDGF and serotonin

Delayed cerebral ischaemia:

One thing that now seems fairly certain, is that chronic cerebrovasospasm represents only one component, amongst a multitude of other interacting factors arising posthaemorrhage, that can lead directly or indirectly to a
CHRONIC VASOSPASM (pathological)

Arterial blood release into CSF

Platelets
- $\alpha$ granule PDGF release
- Dense granule serotonin release
  - Smooth muscle mitogenesis & hypertrophy

Leucocytes
- Intense contraction
- Perivascular sympathetic excitotoxic transmitter release

Macrophage phagocytosis
- Free radical generation
  - Lipid peroxidation, DNA & protein damage
    - Vessel wall injury, cell death, inflammation & fibrosis

Erythrocytes
- Oxyhaemoglobin autoxidation
delayed onset of reduced cerebral blood flow with its attendant risk of cerebral ischaemia (Figure 28). Autoregulation of cerebrovascular blood flow is commonly disturbed after SAH, and it becomes increasingly impaired in those patients in the worst clinical grades. Such patients tolerate poorly, even quite modest falls in their arterial blood pressure. More recently, a greater awareness has emerged of the frequency with which patients who develop hyponatraemia after SAH also suffer from hypovolaemia. In these patients there is an urgent need to actually expand the circulating volume, rather than to fluid restrict them and thereby worsen the risk of ischaemia. Similarly, the role of communicating hydrocephalus and raised intracranial pressure developing subsequent to haemorrhage, has become more commonly recognised as one readily reversible cause of a compromised cerebral perfusion. Finally, an interesting finding that has recently emerged from experimental studies, is the suggestion that the sympathetic outflow from the brainstem locus coeruleus (LC) nucleus is also activated following SAH, along with the more well known activation of the peripheral sympathetic system (McCormack et al., 1986; Soloman et al., 1986). Potentially, this is of considerable interest, as it is known from experimental studies that stimulation of the LC nucleus can result in a global reduction in CBF (Raichle et al., 1975). It is possible therefore, although as yet only speculative, that in man a greatly increased activity of the LC nucleus after SAH may be one further mechanism in the pathophysiology of the reduced CBF following SAH.
CEREBRAL ISCHAEMIA FOLLOWING SAH

Arterial blood release into CSF

- Pulmonary oedema and hypoventilation
- Raised ICP and hydrocephalus
- Cerebral arterial luminal narrowing
- Locus coeruleus sympathetic overactivity
- Auto-regulatory paralysis
- Hyponatraemic hypovolaemia

Hypoxia

Reduced cerebral perfusion

- Arterial hypotension

NEURONAL ISCHAEMIA AND DEATH
11 ACKNOWLEDGEMENTS AND STATEMENT OF ORIGINALITY

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During the course of performing these studies I held the following appointments:

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13 PUBLICATIONS ARISING

The following papers, either arising out of, or related to the work performed in the preparation of this dissertation, have been published or accepted with a view to publication.


