Introduction

Subarachnoid hemorrhage (SAH) affects an estimated 25,000 to 30,000 individuals in the USA each year, and is a significant source of death and disability. Nearly 35% of patients do not survive the initial insult. Subsequent to SAH, many survivors experience constriction of cerebral arteries, commonly described as vasospasm. The exact mechanism by which constriction of cerebral arteries occurs following SAH is unclear, and may involve a number of factors. Arterial constrictions typically occur within 3–21 days after SAH and may last for 12–16 days. At 7 days after SAH, angiographic evidence indicating vascular constriction, commonly termed ‘arterial vasospasm’, is present in approximately 30–70% of patients. However, only 20–30% display signs of neurological deficit, termed ‘symptomatic vasospasm’. Constriction of cerebral vessels leads to the restriction of blood flow, causing severe cerebral ischemia in some patients, and is associated with an increase in mortality of 1.5–3-fold during the first 2 weeks after SAH. The term ‘cerebral vasospasm’ is commonly used to describe both the actual arterial constriction and the symptomatic deficits. The effects of SAH are not limited to vascular constriction, however, and may include damage to vascular tissue caused by inflammation and oxidative stress, indicating the presence of a broader pathologic condition. The purpose of this review is to suggest that these additional aspects should be taken into account during formulation of a treatment strategy in order to maximize benefit and minimize neurological damage following SAH.

Vascular changes after SAH

Many of the vascular effects of vasospasm have been evaluated using animal models of SAH. Animal models have the benefit of high reproducibility and the ability to evaluate changes at specific time points after SAH, which cannot be achieved with post-mortem human tissue. Models of SAH have been developed in a number of species, including dogs, cats, rats, and even primates. The double-hemorrhage model is most commonly used and typically consists of two injections of autologous blood into the cisterna magna. A single-hemorrhage model consisting of only one injection of blood may also be used. In rats, an endovascular puncture model involving puncture of the internal carotid artery at its bifurcation point is...
used as well. Other models may involve the implantation of preformed clots into the subarachnoid space, such as in primates. In a comparison of models used in rats, the double-hemorrhage model was found to best represent the time course of SAH in humans and produce the greatest severity of vasospasm.

Within 3–7 days after SAH in dogs, there is a significant decrease in the internal diameter of arterioles in the brain that is associated with an increase in wall thickness. Within the tunica media, there is hypertrophy and hyperplasia of smooth muscle cells. There are also signs of cell damage and death such as pallor of the smooth muscle and pale, pyknotic nuclei with indistinct cell membranes. Inflammation secondary to SAH can also lead to an increase in the thickness of the tunica adventitia. Collagen fibers within the cerebral vascular wall are increased after SAH in cats, particularly on the adventitial side. Connective tissue within the vessels is increased, and there is increased fibrosis in the tunica media, intima, and subintima. Moreover, there are obvious changes in the internal elastic lamina such as corrugations and increased thickness. Increased wall thickness is accompanied by narrowing of the lumen due to fibrotic and fibroproliferative processes. Changes in human vessels are similar to those reported in experimental animal models and include thickening of the tunica adventitia due to edema with associated accumulation of inflammatory cells, necrosis and atrophy of smooth muscle cells in the tunica media, and thickening of the tunica intima due to increased connective tissue and fibrosis leading to a decrease in lumen diameter.

The endothelium also exhibits changes in response to SAH. Endothelial cells appear swollen, detached from the basement membrane, and necrotic in a rat model of SAH. There is a vacuolization in their cytoplasm and nuclei, disruption of tight junctions, and widening of the inter-endothelial space. Endothelial cells show signs of apoptosis beginning at 3 days after SAH in a canine model. The use of caspase inhibitors to block apoptosis has been shown to reduce angiographic vasospasm in dogs. There are also changes in the endothelial response to vasoactive agents in the cerebral vessels after SAH. The effects of acetylcarnitine and the endothelium-dependent vasodilator calcimycin on cerebral blood flow and cerebrovascular resistance are altered after SAH in a primate model. Endothelial dysfunction as a result of SAH-induced damage may therefore play an important role in the occurrence of vasospasm.

Inflammatory responses to SAH

During the first hour after SAH, white blood cells (WBCs) begin to associate with the endothelium. Adhesion of WBCs to the endothelium and their subsequent migration into the periadventitial space may play an important role in vascular constriction. Adhesion of WBCs to the endothelium is mediated by lymphocyte function-associated antigen-1 (CD11a/CD18) and macrophage antigen-1 (CD11b/CD18) on neutrophils and macrophages, and by intercellular adhesion molecule-1 (CD54) on endothelial cells. Monoclonal antibodies directed against CD11/CD18 inhibit the decrease in lumen diameter following SAH in rabbits. One potential role for WBCs in the response to SAH may be the secretion of cytokines. Activation of interleukin-1 beta is increased in lipopolysaccharide-stimulated monocytes from SAH patients with symptomatic cerebral vasospasm. Peak leukocyte counts greater than 1.5 × 10^9/l are associated with a greater than threefold increase in vasospasm after SAH.

Phospholipase activity, which involves the release of arachidonic acid and other free fatty acids from membrane phospholipids, is enhanced after many types of brain injury. The production of inflammatory cytokines after SAH results in the induction of cyclooxygenase 2, which is involved in the breakdown of arachidonic acid. The lipoxygenase pathway of arachidonic acid breakdown is also stimulated by SAH. The products of these enzymes have roles in vasoconstriction of blood vessels, neutrophil and eosinophil chemotaxis, activation of lymphocytes, increased vascular permeability, and increased reactive oxygen species production. ABT-761 and 1,2-bis(nicotinamido)propane, inhibitors of 5-lipoxygenase, attenuate the vasospastic response in experimental animals.

Platelet-activating factor (PAF) may also play an important role in the inflammatory response and vasospasm after SAH. Plasma concentrations of PAF are higher in SAH patients with cerebral infarction due to vasospasm. Animal studies have shown that the administration of the PAF antagonist E5880 can prevent vasospasm following SAH. Inhibition of poly(ADP-ribose), another important molecule in modulating inflammation, also attenuates cerebral vasospasm. Inflammation therefore seems to play multiple roles in the development of vasospasm, and presents a number of possible targets for the treatment of SAH.

The influence of molecular mechanisms on vasospasm after SAH

Nitric oxide, hemoglobin, and endothelin

The constant release of nitric oxide (NO) from the endothelium helps to maintain basal cerebrovascular tone by inducing vascular relaxation. The release of NO by endothelial cells has been shown to be decreased after SAH. Nitric oxide is an inhibitor of the vasoconstrictor endothelin-1, and inhibition of NO synthesis leads to vessel constriction and reduced blood flow that can be alleviated by endothelin receptor blockade.
Endothelin results in the influx of calcium into vascular smooth muscle cells, leading to their contraction and vessel constriction. Endothelin levels are increased in the cerebrospinal fluid (CSF), plasma, and hypothalamus after SAH. Increased plasma endothelin is associated with delayed cerebral ischemia and vasospasm after SAH. Lysis of red blood cells after SAH also leads to the release of hemoglobin, resulting in increased concentrations within the perivascular space. Hemoglobin binds and inhibits the activity of NO, thereby enhancing vascular constriction.

In rodent models of SAH, intracranial administration of nitric oxide donors increases levels of NO and prevents vasospasm. The NO donor sodium nitroprusside has also been shown to prevent or even reverse vasospasm when administered intraventricularly in human SAH patients. Inhibition of endothelin-induced calcium influx reduces the magnitude of vasospasm in animal models of SAH. Metabolism of hemoglobin by hemoxygenase-1 or chelation of its ferrous component also prevents vasospasm in animals models of SAH. These results suggest that a reduction in NO availability may play a role in the pathogenesis of vasospasm, and that the administration of NO donors may be beneficial for the prevention and treatment of vasospasm. The use of polymers that allow a controlled release of NO donors has shown considerable promise in animal models. To date, however, only local injection of these polymers directly into the brain has been tested. This route of administration may be a limiting factor in the treatment of human patients due to the specialized facilities that would be required. The effects of these compounds when administered via more traditional routes should be explored to determine whether they are practical and safe for widespread use.

Free radicals and oxidative damage
Reactive oxygen species, particularly superoxide, are believed to play a role in the pathogenesis of vasospasm after SAH. Effects of superoxide include destruction of NO, DNA and protein dysfunction, lipid peroxidation, and elevation in the levels of hydrogen peroxide. Lipid peroxides in the CSF are increased following SAH in the dog and can be blocked by the administration of methylprednisolone. Enhanced production of superoxide is seen in the subarachnoid space after SAH and can be blocked by the intracisternal administration of superoxide dismutase, which metabolizes superoxide into less toxic products, in a canine SAH model. Mice overexpressing superoxide dismutase also show reduced vasospasm after SAH.

Oxidative damage is believed to play a role in a number of neurological diseases, and a considerable amount of effort is being put into the investigation of antioxidants as potential treatment. It is possible that one of these compounds may prove to be beneficial for the treatment of vasospasm as well. The free radical scavenger tirilazad mesylate, although showing initial signs of promise, has failed in human trials upon further study. Among four large multicenter trials, only one has shown a reduction in the frequency and severity of vasospasm. The differences between trials can potentially be explained by methodological differences, particularly the use of other potentially beneficial treatments. Numerous other potential compounds remain to be tested, however, many of which can be easily administered and have already been shown to be safe for use in humans.

Prevention and treatment of vasospasm
In some cases, the occurrence of hemorrhage can be avoided with proper treatment. Endovascular coiling, for example, may restrict blood from entering an aneurysm and prevent it from rupturing (Figure 1). Once hemorrhage has occurred, however, efforts should focus on...
preventing its pathological effects. It is therefore important to predict and detect the occurrence of vasospasm in SAH patients as early as possible in order to maximize the effects of preventative treatment. Clinical grade is commonly used as a predictor for the risk of vasospasm, but other criteria may also be used for the evaluation of patient risk such as the amount of blood in the subarachnoid space, the size and location of the hemorrhage, age, sex, hypertension, and tobacco or drug use. Currently, however, the presence of a large amount of blood in the subarachnoid space is the only consistent risk factor for vasospasm. Transcranial Doppler ultrasonography can also be effective for the prediction of vasospasm, but only after the vessel has begun to constrict, thus limiting the amount of time for preventative treatment before the onset of ischemia.

The most common form of prevention of vasospasm after SAH is the administration of calcium antagonists such as nimodipine. By preventing the calcium-induced contraction of vascular smooth muscle, nimodipine and other antagonists may be useful to block the occurrence of arterial constriction after SAH. Oral nimodipine treatment has been shown to reduce the incidence of cerebral infarction after subarachnoid hemorrhage. Intra-arterial nimodipine has also been shown to be beneficial for the treatment of vasospasm. Among a group of nine patients treated with intra-arterial nimodipine, vessel diameter in the vasospastic segment increased by an average of 66.6% with no alteration of blood pressure. Intravenous nimodipine, however, appears to have no benefit, and other calcium antagonists have failed to consistently prevent vasospasm when tested. For example, the calcium channel blocker verapamil, commonly used for the treatment of coronary vasospasm, has shown improvement in some studies but no effect in others. Evaluation of 12 clinical trials with a total of 2844 patients, 1396 of which received calcium antagonists, found an overall risk reduction of 5.1% with calcium antagonist treatment. This reduction, however, could be attributed primarily to oral nimodipine treatment, and the results of treatment with nicardipine, AT877, or magnesium were considered inconclusive.

Treatment for vasospasm may include the induction of hypervolemia, hypertension, and hemodilution, commonly known as “Triple-H therapy.” The goal of this treatment is to maintain adequate blood flow through affected vessels. Triple-H therapy has been shown to maintain or improve neurologic grade in a majority of patients, resulting in minor or no neurological deficits. The efficacy of this treatment has been questioned, however, and in some cases it may be harmful. Endovascular treatments such as balloon angioplasty or administration of vasodilators or calcium antagonists may also be used to increase the luminal diameter of constricted vessels during vasospasm (Figure 2). Angioplasty has been shown to result in clinical improvement, particularly when administered early after the onset of symptoms. Other groups, however, have found no improvement in outcome after angioplasty despite effective reversal of vasospasm. The effectiveness of this procedure, therefore, is questionable. The non-specific smooth muscle relaxant papaverine can also be used to increase the lumen size in vasospastic vessels, either alone or in combination with angioplasty.

Several other compounds have also shown promise for the treatment of vasospasm. Magnesium sulfate has been shown to reverse vasospasm after SAH in rats. Although further studies are needed, it has also been shown to reduce the occurrence of delayed cerebral ischemia after SAH in human trials. Out of 283 patients receiving either magnesium or placebo, magnesium reduced the risk of new lesion development by 34%. Potential mechanisms of magnesium

![Figure 2](image_url)
include inhibition of excitatory amino acid release and antagonism of NMDA receptors and calcium channels. Statins, or 3-hydroxy-3-methylglutaryl coenzyme A reductase (HMGR) inhibitors, may represent another potential treatment. In addition to their known role in blocking cholesterol synthesis, statins also regulate nitric oxide levels and have antioxidant and anti-inflammatory properties. Pravastatin has been shown to reduce the incidence and duration of vasospasm after SAH in clinical trials. Among a group of 80 patients given pravastatin or placebo, pravastatin treatment reduced the incidence of vasospasm by 32% and shortened the duration by 0.8 days. More importantly, delayed ischemic deficits were reduced by 83% and mortality by 75%. Studies with a larger number of patients are currently in progress.

The American Heart Association has established guidelines for the management of subarachnoid hemorrhage. However, these guidelines were formulated in 1994, and before endovascular therapy was commonly used. These guidelines include recommendations for the treatment of vasospasm. The recommended treatment is clipping of the aneurysm, Triple-H therapy for the management of ischemic complications, and oral nimodipine for the improvement of outcome. Patients should be closely monitored during treatment, and transluminal angioplasty is recommended if the standard treatment fails. Although a number of promising treatments are currently under evaluation, further study is still needed to determine their safety and efficacy compared with what is currently being used.

Summary

The exact mechanism by which vasospasm is induced following SAH is unknown. One potential pathway emphasizing the roles of nitric oxide and endothelin is shown in Figure 3. In this pathway, the level of NO is decreased by a number of factors, leading to an increase in endothelin levels that induces calcium influx and vascular constriction. This is but one potential mechanism for the induction of vasospasm, and it is likely that many other pathways are involved. Further research is required to determine which pathways, if any, play a dominant role. From the studies described above, it is clear that vasospasm can be influenced by a number of factors. What is not clear, however, is the magnitude of the role that each of these factors plays in the development of vasospasm after SAH. It is possible that only one or a few of these processes are directly responsible for the induction of vasospasm and that the remainders serve only to modulate the degree of vascular constriction. Furthermore, many of these processes have other effects on tissue that are unrelated to their potential role in vasospasm. Endothelial cell death within blood vessels would be expected to have additional effects beyond constriction of blood vessels. Inflammation and reactive oxygen

![Figure 3](image_url)
species are expected to be damaging not only to vessels but to the surrounding tissue. In addition, many of the molecules and enzymes mentioned in this review function in multiple biochemical pathways. Vasospasm may therefore be considered as merely one component of a broader condition involving multiple damaging effects. The term vasospasm specifically refers to only a single phenomenon, the constriction of blood vessels, and is therefore inadequate to encompass the full range of effects seen. We therefore propose the use of a broader term, post-subarachnoid hemorrhage vasculopathy, to better describe the resulting condition after SAH.

**References**


