Abstract
An overview of the main features of cerebral vasospasm is provided in this report, highlighting the possible future direction of development in the diagnosis and management of this severe complication of aneurysmal subarachnoid hemorrhage.

Introduction and context
Vasospasm is one of the most dreaded acute complications, developing 4–15 days after aneurysmal subarachnoid hemorrhage (SAH). It is characterized by a pathological, diffuse, and long-lasting narrowing of the vessel lumen of large-capacity cerebral arteries at the base of the brain either close or distal to the site of bleeding, and is associated with a reduced perfusion of the territories distal to the affected vessel. Angiographic vasospasm occurs in 30–70% of patients with SAH, but it leads to clinically evident signs and symptoms in 20–30% of patients who experience delayed ischemic neurological deficits. About half of this latter group of patients suffer severe permanent neurological dysfunction or death [1].

Vasospasm affects all layers of the involved arterial wall of the cerebral vessels. A proliferative inflammatory arteriopathy is the pathological feature of cerebral vasospasm. In fact, the adventitia is infiltrated with inflammatory cells and the neuronal endings are damaged. The media is thickened and fibrotic, with an increased proliferation of smooth muscle cells. The intima shows a disruption of the internal elastic lamina [2].

An important predictor of the occurrence of vasospasm after SAH is the amount of blood present around the cerebral arteries of the circle of Willis. The Fisher computed-tomography rating scale of SAH, and recent modified versions, have demonstrated a strong clinical correlation with the development of clinically significant vasospasm [3–5]. Patients with thick basal cistern blood and the presence of intraventricular blood in the lateral ventricles carry the highest risk. Other risk factors include young age, hypertension, smoking, and cocaine use [6].

It has been clearly demonstrated that prolonged exposure of cerebral arteries to perivascular blood is necessary for the development of vasospasm. However, it has been impossible until now to identify a single causative molecule as the culprit of vasospasm. Nonetheless, there is evidence that a few agents, such as oxyhemoglobin, nitric oxide, and endothelin-1, may be contributors to this pathological event.

Oxyhemoglobin, a product of auto-oxidation of hemoglobin, can directly or indirectly induce arterial vasoconstriction, especially if the oxygen-free radical scavenging systems are insufficient. Oxyhemoglobin can also exert a scavenging effect toward nitric oxide (a potent vasodilator whose depletion has been demonstrated during vasospasm) and can stimulate endothelial cells to produce endothelin-1. Endothelin-1 causes the most potent and long-lasting vasoconstrictor effect, which is also associated with morphological changes, mimicking the delayed cerebral vasospasm. It has been demonstrated that endothelin-1 levels are increased, not only in the cerebrospinal fluid during SAH, but also during severe neuronal injury (when caused independently from vasospasm or the primary bleeding event).
Furthermore, endothelin levels change in parallel with neurological symptoms, but are not predictive of vasospasm as assessed by transcranial Doppler (TCD). These observations do not exclude a causative role of endothelin-1 for vasospasm but rather suggest that endothelin-1 acts as a marker of cerebral ischemic injury [7–10].

**Recent advances**

**Diagnosis**

Angiography of the vessels of the brain is the gold standard for the diagnosis of cerebral vasospasm. However, this procedure is invasive, requires the availability of significant resources, and may cause vessel dissection or thrombosis. Alternative diagnostic tests, such as computed tomographic angiography and TCD, have now been clinically validated [11]. Magnetic resonance imaging, radionuclide imaging, and electroencephalography have also been investigated as diagnostic tools.

TCD is not invasive and can be performed at the bedside. For the middle cerebral artery, TCD has a high specificity with a threshold value ranging between 160 and 200 cm/s [12]. TCD evaluation is recommended as a screening tool in high-grade WFNS (World Federation of Neurological Surgeons) scale patients in whom a neurological examination cannot be readily followed to identify those at higher risk [13].

In the most severe cases needing monitoring of intracranial pressure and cerebral perfusion pressure, the use of cerebral microdialysis has been proposed to identify the threshold of anaerobic metabolism (expressed by the lactate/pyruvate ratio as an indirect sign of hypoperfusion). Cerebral microdialysis in association with other brain-monitoring techniques may assist in the delivery of targeted therapy for prevention of secondary ischemic injury [14].

**Treatment**

Critical care management of patients with aneurysmal SAH aims at improving neurological outcome, and includes the treatment of non-neurological systems affecting the brain; a multi-organ clinical approach instead of a single-organ approach probably represents the optimal way to reach this goal. Indeed, recent studies showed that strategies directed at maintaining normothermia, normoglycemia, and prevention of anemia may improve outcome after SAH. In fact, fever, anemia, and hyperglycemia affect 30–54% of patients with SAH and are significantly associated with mortality and poor functional outcome [15].

The specific treatment of cerebral vasospasm aims at improving cerebral blood flow with one of two possible approaches: indirect pharmacological protection of the brain tissue or direct mechanical dilation of the vasospastic vessel.

Though not proven by any randomized clinical trial, induced hypertension, hypervolemia, and hemodilution (triple H therapy) are considered the mainstay of the treatment of vasospasm. This strategy is associated with a high rate of complications however, limiting its usefulness, and has been demonstrated to be ineffective in many patients [16]. A recent investigation, focusing on the effect of each component of triple H therapy on cerebral blood flow, showed that only vasopressor-induced elevation of mean arterial pressure caused a significant increase of regional cerebral blood flow and brain tissue oxygenation, confirming that cerebral pressure autoregulation is often impaired in these patients [17]. Although there is no proven role for hypertovolemia, hypovolemia must be avoided because it increases the risk of delayed infarction [18]. Regarding the hemodilution component, recent studies tried to identify the optimal hemoglobin range and concluded that extreme hemodilution should be avoided, particularly in patients with a cardiac condition [19].

The calcium channel blocker nimodipine is considered the standard of care in aneurysmal SAH patients. Although experimental studies failed to show its ability to prevent angiographic vasospasm, its administration immediately after SAH diagnosis and for 10–15 days thereafter has been shown to improve outcome and reduce cerebral infarction [20].

Several experimental treatments have recently been proposed for the management of cerebral vasospasm (such as, cisternal thrombolysis, surgical removal of the clot, lipid peroxidation inhibitors, and the use of scavengers of hydroxyl radicals), but further evidence is needed to prove their efficacy [21]. Currently, the most promising experimental treatment is the administration of magnesium sulfate, statins, nitric oxide donors, or endothelin-1 antagonists. High-dose magnesium therapy might be efficient as a prophylactic adjacent therapy after SAH to reduce the risk for poor outcome. Nevertheless, because of the high frequency of side effects, patients should be observed in an intensive or intermediate care setting and hypomagnesemia should always be avoided [22].

A meta-analysis based on three small clinical trials showing reduced incidences of vasospasm, delayed ischemic deficits, and mortality, advocates the routine...
therapeutic use of statins after aneurysmal SAH [23]. A phase II clinical trial with an endothelin receptor A antagonist, clazosentan, has demonstrated a reduction in the incidence and severity of angiographic vasospasm from 66 to 23%, while adverse events were comparable to those of placebo [24]. Moreover, another phase II trial, testing an antagonist against the endothelin receptors A and B, showed an improvement in delayed ischemic deficits, occurring in 30% of patients receiving active treatment and 37% of patients on placebo [25].

Mechanical dilation of the vasospastic vessel by transluminal cerebral angioplasty has been demonstrated to effectively reverse radiographically confirmed vasospasm [26]. Moreover, with this technique, it is possible to selectively inject vasodilators, such as papaverine and more recently nicardipine, milrinone, and verapamil [27]. Nevertheless, since few long-term outcome studies are available, its superiority to medical management for symptomatic cerebral vasospasm is questionable [28]. Therefore, the use of angioplasty should be reserved for patients who failed to improve with conventional hemodynamic management.

Recently, the effect of prophylactic transluminal balloon angioplasty on cerebral vasospasm in patients with SAH (Fisher grade III) has been investigated in a randomized clinical trial [29]. Fewer patients developed vasospasm after treatment with angioplasty, and there was a significant decrease in the need for therapeutic angioplasty, however, the use of angioplasty did not improve the outcome of these patients.

Implications for clinical practice
Although knowledge of the pathophysiology of vasospasm after SAH has advanced significantly over the past years, it continues to be a major cause of mortality and morbidity without a known specific treatment.

Recent studies have validated the use of computed-tomographic angiography and TCD in the diagnosis of vasospasm, and it has been shown that cerebral microdialysis in association with other brain-monitoring techniques may assist in the delivery of targeted therapy to prevent secondary ischemic injury.

A multi-organ clinical approach to the treatment of cerebral vasospasm is recommended, including the maintenance of normothermia, normoglycemia, and the prevention of anemia.

The efficacy of triple H therapy remains uncertain and its limitations need to be recognized. Studies into the administration of magnesium sulfate, statins, nitric oxide donors, or endothelin-1 antagonists have, however, proved promising, but further efforts are needed to translate these results into clinical practice in order to improve the neurointensive care management of this severe complication.

Some improvement was seen in preventing vasospasm using prophylactic transluminal balloon angioplasty, but the use of angioplasty should be reserved for those patients who failed to improve with conventional hemodynamic management.

Abbreviations
SAH, subarachnoid hemorrhage; TCD, transcranial Doppler; WFNS, World Federation of Neurological Surgeons.

Competing interests
The authors declare that they have no competing interests.

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Changes Clinical Practice
Factor 6.0 Must Read
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