EXTREMELY DELAYED CEREBRAL VASOSPASM AFTER SUBARACHNOID HEMORRHAGE

Angelo Daros Cecon1, Eberval Gadelha Figueiredo2, Edson Bor-Seng-Shu3, Milberto Scaff4, Manoel Jacobsen Teixeira5

Delayed cerebral ischemia is an important cause of poor outcome in patients with aneurysmal subarachnoid hemorrhage (SAH). This clinical syndrome is caused by a critical reduction of cerebral blood flow brought on by a pathologic luminal narrowing of the cerebral arteries after SAH (vasospasm). It usually presents within 12 days after SAH and its pathogenesis probably depends upon inflammatory mechanisms. Its occurrence after 15 days is quite uncommon. Late diagnosis is often difficult and many times overlooked. Considering the role of triple H therapy in the treatment, differential diagnosis acquires paramount importance.

We present a rare case of extremely delayed cerebral vasospasm after SAH.

CASE
A 43 years-old female attended to the emergency room (ER) complaining of sudden, intense headache. Neurological examination disclosed meningismus. She was alert, presenting score 15 in the Glasgow Coma Scale. Head CT revealed a Fischer grade III SAH (Fig 1). Angiogram depicted an anterior communicating artery aneurysm, without signs of radiological vasoespasm (Fig 2). Aneurysm clipping was uneventfully performed three days after SAH, without need for temporary clipping. Postoperative

Fig 1. Axial head CT scan depicting a subarachnoid hemorrhage Fisher grade III.
CT scan did not reveal abnormalities. There was no clinical vasospasm during the first twenty-one days of SAH, and patient was neurologically intact when discharged.

After 45 days, she returned to the ER presenting a left-sided hemiparesis. CT scan revealed a capsular ischemic area on the right cerebral hemisphere (Fig 3). No antecedents of hypertension and diabetes were observed. Arterial pressure levels were normal at admission. Serum levels of glucose and cholesterol were normal. Cardiac ultrasonography revealed no abnormalities as well as carotid and vertebral ultrasonography. Transcranial doppler (TCD) ultrasonography disclosed higher flow velocities suggestive of vasospasm. Triple H therapy was installed and patient completely recovered. Clinical improvement coincides with normalization of the TCD initial findings. Patient was discharged clinically intact.

**DISCUSSION**

Symptomatic vasospasm has been reported in 22% to 40% of patients suffering SAH and leads to high rates of morbidity (34%) and mortality (30%)\(^2\). The diagnosis of vasospasm is based upon clinical findings and evidence of luminal narrowing on angiogram or flow alterations on TDU\(^3\). Angiographic evidence of arterial spasm is seen in up to 70% of patients, and clinical manifestations are witnessed in 20 to 30% of patients\(^5,6\). Despite the introduction of nimodipine and the use of induced hypertension, symptomatic vasospasm still remains the major cause of morbidity and mortality in patients with SAH\(^7-11\).

Symptomatic vasospasm is defined by clinical criteria such as: 1) symptoms occurring between days 5 and 12 after SAH, including headache, stiff neck, low-grade fever, insidious confusion, decline in level of consciousness and focal neurological deficit; 2) a head CT scan excluding other causes of worsening; and 3) no other identifiable cause of neurological deterioration\(^8,9\). Vasospasm can occur within 30 days after SAH, and clinical vasospasm after one month aneurysm bleeding is extremely rare and may difficult differential diagnosis. The case of patient who presented clinical deterioration after 45 days of SAH confirmed by TCD and clinical response to spasms therapy. In the present case, the age of the patient, absence of risk factors, normal carotid, vertebral and cardiac ultrasonography, associated with the TCD findings, coupled with the clinical improvement after triple H therapy and normalization of the TCD, practically excludes other causes of ischemic damage (embolism, aterothrombosis, etc) points to the vasospasm as the causal factor.

Delayed cerebral vasospasm, which is the most critical clinical complication that occurs after SAH, seems to be
associated with both impaired dilator and increased constrictor mechanisms in cerebral arteries \(^3\)\(^4\)\(^5\)\(^6\)\(^8\)\(^9\)\(^10\). Mechanisms contributing to development of vasospasm and abnormal reactivity of cerebral arteries after SAH have been intensively investigated in recent years \(^10\)\(^11\)\(^12\)\(^13\). The studies in the pathogenesis of vasospasm have focused on the inflammatory response after SAH, which may play a relevant role in its development and maintenance \(^14\), secondary to a sudden abnormal contact of the extraluminal walls of arteries with all of the components of blood \(^2\). The inflammatory factors induced by SAH seem to be a great factor causing arterial spasm.

In the instance of SAH, a complex series of cellular and molecular events is elicited by the presence of blood clot in the subarachnoid space, culminating in a robust inflammatory response. Erythrocytes are dissolved, releasing oxyhemoglobin, which diffuses through the arterial wall of the endothelium \(^13\). A cascade of inflammation occurs, including infiltration and activation and recruitment of leukocytes, cytokine production, immunoglobulin and complement activation, and transcription factor activation, resulting in prolonged smooth muscle contraction and arterial spasm \(^13\). These processes are auto-limited in the majority of the cases, however, delayed manifestations may occur.

It is possible that the mass effect of a large hematoma remaining in the sylvian fissure and an intracerebral hematoma after surgery may justify delayed development of spasm in some cases. Nonetheless, in the current case there was no hematoma left to explain late vasospasm. It is possible that the continuous inflammatory disease could bring to a progressive nip of brain arteries, and this was only revealed later in the course.

In conclusion, even though a very rare occurrence, extremely delayed vasospasm may occur after the first month after SAH. The neurological follow-up should be judicious and include frequent medical consultation in the first two months. Its occurrence must always be considered in the differential diagnosis of late neurological deterioration.

REFERENCES