Delayed Positivization of Cerebral Angiography in Reversible Cerebral Vasoconstriction Syndrome (RCVS) Presenting with Recurrent Subarachnoid Haemorrhage

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Abstract: Benign angiopathy of the central nervous system is a clinical syndrome with evidence of reversible cerebral vasoconstriction (RCVS). Haemorrhagic stroke, either subarachnoid or intracerebral is an unusual presentation of RCVS. We describe a case of RCVS presenting with a subarachnoid haemorrhage (SAH), with rebleeding and onset of hydrocephalus during the first week, and, notably, delayed evidence of typical angiographic features after two negative prior exams. Normalization of the angiographic vasculitic-like lesions was documented at month +6. Repeated cerebral angiograms are mandatory to exclude this kind of disease, and the uncommon presentation of this case reinforces this concept.

Keywords: Reversible cerebral vasoconstriction syndrome, angiography, hydrocephalus, subarachnoid haemorrhage, vasculitis.

INTRODUCTION

Vasculitis of the central nervous system (CNS) is classified as primary angiitis or as secondary vasculitis related to a variety of diseases [1]. Reversible cerebral vasoconstriction syndromes (RCVS) represent a critical differential diagnosis of CNS vasculitis and they comprise a group of disorders characterized by prolonged but reversible vasoconstriction of the cerebral arteries, usually associated with acute-onset, severe, recurrent headaches, with or without additional neurologic signs and symptoms [2]. We report a case of a female patient with initial suspicion of primary angiitis of the cerebral nervous system (PACNS) presenting with a subarachnoid haemorrhage (SAH). Delayed evidence of typical angiographic features after negative prior angiograms, and the disappearance of the abnormalities six month later led to revise the initial diagnosis of PACNS towards the diagnosis of RCVS. The occurrence of delayed positivization of the cerebral angiograms must be considered for the diagnosis of this kind of the disease and it may differentiate RCVS from PACNS.

CASE REPORT

A 47-year-old woman was admitted to the Emergency Department of Udine Hospital on June 2006 for a sudden onset of thunder-clap headache following by loss of consciousness. Cerebral computed tomography scan (CT) revealed a subarachnoid haemorrhage, thus the patient was soon transferred to the Neurosurgery Unit. At admission in neurosurgical ward the patient was awake, alert, apyretic and with mild arterial hypertension (150/90 mmHg). No intracranial aneurysms or arterio-venous malformations (AVM) were found on cerebral four vessels angiography performed the first day after admission (Fig. 1). The patient was treated with nifedipine, obtaining normalization of blood pressure values. At the entrance inflammatory markers, such as erythrocyte sedimentation rate (ESR), C-reactive protein (CRP), serum ferritin and fibrinogen were normal. Hemoglobin level, leukocyte and platelet count, hepatic and renal function tests, protein electrophoresis, C3 and C4 levels were normal.

Seven days later the patient experienced a worsening of her neurological status, she complained of a crushing headache and nausea, and she vomited several times. A CT scan was performed and showed a new perimesenchepalic bleeding and acute hydrocephalus. An external ventricular drainage was performed. After this surgical procedure the neurological status temporally improved.

The second cerebral four vessels angiography performed fourteen days after the first one as a control after the second cerebral haemorrhage, did not show any arterial abnormalities. In particular, no aneurysms or AVM were documented. In the following days the patient referred mild headaches, myalgias and even muscle weakness, and developed a slight right hemiparesis. Serial CT scans documented the progressive reabsorption of the subarachnoid haemorrhage.
The third cerebral four vessels control angiography performed one month after the first one, (Fig. 2) showed marked focal narrowing and stenosis of both carotid siphons (supraclinoid tract), of the terminal tract of the basilar artery, of both P1-P2 tracts, and A1 right tract. No venous or capillary abnormalities were noticed. These findings, very delayed after the onset, excluded a vasospastic complication of SAH [3]. A ventriculo-peritoneal shunt was performed.

Fig. (1). Conventional angiography, first one, of the vertebrobasilar system (A, B) and the left carotid system (C): no arterial abnormalities, in particular no aneurysms or A-V malformation.

Fig. (2). Conventional angiography, one month after bleeding: marked focal narrowing and stenosis of both carotid siphons (arrows), terminal tract of the basilar artery, both P1-P2 tracts, and A1 right tract.
Then, the patient was evaluated by the Rheumatologist consultant, who planned an intense workup to clarify the diagnosis of cerebral vasculitis. Cerebral vasculitis due to a systemic disease was excluded based on i) the patient’s history and physical examination, not indicating an underlying systemic disease, such as systemic vasculitis, or connective tissue diseases; ii) the absence of serum antinuclear antibodies, anticytoplasmic neutrophil antibodies, rheumatoid factor, antiphospholipid antibodies and cryoglobulins. Inflammatory markers, such as ESR, CRP, serum ferritin and fibrinogen were normal. Cerebrospinal fluid (CSF) analysis resulted abnormal, showing lymphocytic pleocytosis and evidence of haemorrhage. Leptomeningeal biopsy was not performed due to the critical condition of the patient at the time of her admission in the Neurosurgery Unit. Based on these findings the diagnosis of PACNS was made and the patient started high-dose corticosteroids (dexamethasone 8 mg x 3/day, for three weeks, then 8 mg x 2/day for one week, then 8 mg/day for one week, then methylprednisolone 32 mg/day, tapered to 6 mg/day in two weeks) and pulse intravenous cyclophosphamide (750 mg weekly for 3 weeks, followed by 500 mg weekly for 2 weeks). The treatment was well tolerated and no other cerebral haemorrhage episodes were observed. No maintenance immunosuppressive treatment was then employed. Subsequently, the patient completed her rehabilitation program. Steroids were tapered to 5 mg/day of prednisone at month +6 and then maintained at this dose due to secondary hypopituitarism. A repeated cerebral angiography at month +6 did not reveal any abnormality with a complete vessels size recovery (Fig. 3). At the last follow-up (month +61) no relapse occurred, and the patient was able to normally make her daily practice activities. The cerebral MRI showed ischemic lesions in left thalamus, left pons and mesencephalon at month +6 (Fig. 4) and at month +12 and then remained unchanged in the subsequent annually repeated MRI.

**DISCUSSION**

The first step to diagnose CNS vasculitis is early suspicion, but all the non-vasculitic processes that might be involved should be excluded [1, 2]. In our case, the patient’s history, the physical exam, the laboratory findings and the instrumental tests excluded other non-vasculitic causes of CNS haemorrhage, i.e. infectious diseases, uncontrolled arterial hypertension, malignancy, aneurism rupture, metabolic disorders, or drug abuse. Furthermore, a CNS involvement either in the context of a systemic vasculitis, such as Wegener’s granulomatosis, giant cell arteritis, polyarteritis nodosa or cryoglobulinemia, or in the course of other rheumatic diseases, such as systemic lupus erythematosus, Sjögren’s syndrome, rheumatoid arthritis, Behçet syndrome, sarcoidosis, was considered and excluded through a careful workup. The initial diagnosis of isolated CNS vasculitis was suggested only by the angiographic study, since the diagnostic brain biopsy and a magnetic resonance imaging study were not performed due to the critical condition of the patient and the CSF was not informative. Thus, the acute onset and the rapid recurrence of cerebral haemorrhage justified an initial aggressive immunosuppressive approach. However, the favourable outcome characterized by a rapid and sustained response after a short course of immunosuppression, CSF slightly abnormal, angiographic changes compatible with vasculitis suggested a benign angiopathy of the CNS, possibly with a vasospasm-induced angiopathy rather than a vasculitis [2, 4, 5]. The disappearance of the abnormal findings in the repeated angiography at month +6 confirmed the benign nature of the disease. Based on a very favourable clinical response, the cyclophosphamide was stopped and steroids were minimized at the lowest dose, as soon as possible.

RCVS can be confused with cerebral vasculitis, and in unusual and acute presentation [2, 4, 5], mainly when intracranial haemorrhage occurred [6]. This disorder is characterized by reversible multifocal narrowing of the cerebral arteries heralded by sudden, severe headaches with or without neurologic deficits, and it seems to well respond to a short course of glucocorticoids and calcium-channel blockers [2]. Typically, angiographic abnormalities resolved within 12 weeks [2]. In our case, the angiographic vasculitic-like findings developed later, when haemorrhage had already occurred, and were absent in the first two examinations. The delayed onset of angiographic vessel abnormalities excluded a vasospastic complication of SAH [3]. However, this delayed appearance of typical vessel abnormalities was quite unusual in RCVS [7], but angiogram-negative SAH is a well recognized and open issue, and it accounts for about 15% of nontraumatic SAH [8]. Also, postpartum cerebral angiopathy

![Fig. (3)]. Conventional angiography, six months after bleeding and after immunosuppressive treatment.
represents a well known subgroup of the RCVS [9, 10], where there is an increasingly description of cases with a delay between clinical onset and angiographic changes [11]. It has been suggested a sequential involvement of arterial beds, initial involvement of small distal arteries (not visible on angiography) progressing over days or weeks to involve medium and large sized arteries (visible by angiography) [12].

Given the small size of the reported group and the absence of a prospective randomized controlled therapeutic trial, a diagnostic algorithm or largely approved treatment strategy is presently lacking in PACNS, the choice of a more or less aggressive immunosuppressive treatment (i.e. prolonged or short course of high-dose steroids with or without cyclophosphamide) should be tailored on the single case [13]. In addition, even if angiography by itself is not sufficient to make the specific diagnosis of vasculitis, biopsy results of angiographically abnormal vessels have been reported as normal [1], perhaps due to sampling error, patchy distribution of isolated CNS vasculitis, and a survey of European neurologists diagnosing CNS vasculitis showed that only a minority sought biopsy proof of the diagnosis [14].

In conclusion, RCVS is a relative uncommon cause of intracerebral haemorrhage [2, 7]. Its definitive diagnosis is based on the reversible nature of the angiographic vasculitic-like lesions. Therefore, repeated cerebral angiograms together with clinical data are mandatory for the diagnosis, since leptomeningeal biopsy is negative in this setting, but false negative results may be also observed in PACNS [15, 16], and magnetic resonance imaging study may be not the exam of choice in acute and critical conditions. Delayed appearance of vasculitic-like lesions on angiography could suggest RCVS rather than PACNS. Early immunosuppressive treatment and type of regimen (less or more aggressive) should be based on the level of suspicion of RCVS or PACNS, and then modulated on the basis of the clinical outcome and the imaging study results in the short term.

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CONFLICT OF INTEREST

Declared none.

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