Spontaneous retroclival hematoma: a case series

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OBJECTIVE Retroclival hematomas are rare, appearing mostly as posttraumatic phenomena in children. Spontaneous retroclival hematoma (SRH) in the absence of trauma also has few descriptions in the literature. None of the reported clinical cases features the combination of an SRH and intraventricular hemorrhage (IVH). Nevertheless, despite extensive cases of idiopathic or angiographically negative subarachnoid hemorrhage (SAH) of the posterior fossa, only a single case report of a patient with a unique spontaneous retroclival hematoma has been identified. In this study, the authors reviewed the presentation, management, and clinical outcome of this rare entity.

METHODS The authors performed a retrospective analysis of all patients with diagnosed SRH at their institution over a 3-year period. Collected data included clinical history, laboratory results, treatment, and review of all imaging studies performed.

RESULTS Four patients had SRH. All were appropriately evaluated for coagulopathic and/or traumatic etiologies of hemorrhage, though no etiology could be found. Moreover, all of the patients demonstrated SRH that both clearly crossed the basioccipital synchondrosis and was contained within a nondependent configuration along the retroclival dura mater.

CONCLUSIONS Spontaneous retroclival hematoma, often associated with IVH, is a rare subtype of intracranial hemorrhage frequently recognized only when MRI demonstrates compartmentalization of the posterior fossa hemorrhage. When angiography fails to reveal an underlying lesion, SRH patients, like patients with traditional angiographically negative SAH, enjoy a remarkably good prognosis.

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KEY WORDS clivus; intracranial hematoma; subarachnoid hemorrhage; vascular disorders

Retroclival hematomas are rare and appear mostly as posttraumatic phenomena in children.10 Spontaneous retroclival hematoma (SRH) in the absence of trauma also has few descriptions in the literature.3,9,11 None of the reported clinical cases features the combination of an SRH and intraventricular hemorrhage (IVH), although Schievink et al. have described a patient with CSF xanthochromia and SRH, suggesting a combination of subarachnoid hemorrhage (SAH) and SRH.3 Nevertheless, despite extensive cases of idiopathic or angiographically negative SAH and IVH,2,2 the vascular or etiological source of SRH remains unexplained, and the relationship among SAH, IVH, and SRH remains unexamined.

Current clinical knowledge regarding patients with SRH is restricted to the aforementioned case report. We describe the first case series of patients with SRH.

Methods

We performed a retrospective analysis of all patients with diagnosed SRH at our institution over a 3-year period from January 2012 to January 2015. Collected data
Illustrative Cases

Case 1

A 58-year-old diabetic male experienced a bilateral frontal thunderclap headache while watching television. He had no recent trauma or injury, nor did he have a history of substance use. On examination, no focal neurological signs were found. Noncontrast CT demonstrated posterior fossa hemorrhage and IVH in a configuration typical of retroclival extraaxial blood (Fig. 1A). Magnetic resonance imaging confirmed the predominant subdural location of retroclival blood (Fig. 1B and C). Diagnostic digital subtraction angiograms obtained on Days 1 and 5 of admission revealed no source of the bleed and no evidence of vasospasm (Fig. 1D). Total spine MRI also revealed no abnormality (not shown). Transcranial Doppler ultrasound showed no evidence of vasospasm. The patient’s hospital course included typical measures for intracranial pressure management and cerebral salt wasting followed by an uncomplicated discharge home. At the 30- and 90-day posthospitalization outpatient clinical follow-ups, the patient denied having any symptoms or complaints.

Case 2

A 64-year-old female experienced the sudden onset of a progressive worsening headache over the course of 1 day. She presented without focal neurological signs. Noncontrast CT demonstrated a retroclival hematoma (Fig. 2A and B). Subsequently, MRI, MR angiography (MRA), and catheter angiography were performed, demonstrating no structural, tumoral, or angiographic cause of the hemorrhage (Fig. 2C and D). An uncomplicated 5-day hospital course with subsequent discharge followed. At the 90-day posthospitalization outpatient clinical follow-up, the patient denied having any symptoms or complaints.

Case 3

A 64-year-old man experienced the acute onset of severe occipital headache with subsequent diplopia. He had a history of hypertension. Admission CT scanning revealed SRH and IVH (Fig. 3A–C). Diagnostic digital subtraction angiograms on Day 2 of admission revealed no source of the bleeding and no evidence of vasospasm (Fig. 3D). Brain MRI, MRA, and spine MRI revealed no source of the bleed but did confirm the location of blood confined to the retroclival subdural space. The patient demonstrated no clinical or ultrasonographic signs of vasospasm during a week of hospitalization and was discharged. At the 90-day and 1-year outpatient clinical follow-ups, the patient complained of occasional mild headaches without accompanying neurological signs or symptoms.

Case 4

A 67-year-old man developed acute-onset severe headache and neck pain. He became unresponsive in the emergency department. Admission CT scanning revealed SRH and punctate hemorrhage within the fourth ventricle (Fig. 4A and B). Subsequently, the patient underwent cerebral angiography, which showed no cause for the anatomical abnormality (Fig. 4C). On Day 2 of hospitalization, the patient underwent brain and total spine MRI (Fig. 4D–F),
which clarified the location of blood within the retroclival subdural space but did not reveal a vascular cause for the SAH. At the 15-month follow-up, the patient described no neurological complaints.

Discussion

Each patient presented with a sudden-onset severe headache. None of them demonstrated sellar or pituitary abnormalities to suggest pituitary apoplexy. The differential diagnosis of thunderclap headache remains broad, including aneurysmal and nonaneurysmal causes. Schievink et al. have reported the only case within the extensive literature on angiographically negative CSF xanthochromia to include an SRH. With our cases, we add to this singular description by clarifying several points. First, 3 of the 4 patients did indeed have IVH in addition to the retroclival subdural hemorrhage. All patients were also appropriately evaluated for coagulopathic and/or traumatic etiologies of the hemorrhage, but no etiology was found. Moreover, all patients demonstrated a contained collection of retroclival blood that both clearly crossed the basioccipital synchondrosis and remained within a nondependent configuration along the retroclival dura.

One limitation of this study is the lack of pathological specimens to definitively evaluate the histo-anatomical location of retroclival blood in our patients. Some authors have argued for a more nuanced nomenclature when describing the meningeal coverings in this region given that the basilar plexus exists within the interdural (subdural) space between the 2-layered dura in this region and that a separately named layer, designated the “anterior pontine membrane” (free arachnoid membrane), separates the clival dura from the subarachnoid space. Although MRI, especially 3D high-resolution 3T MRI, helps identify retroclival blood within the extraaxial epidural or subdural
space rather than the subarachnoid space, no current imaging modality specifically allows evaluation of the dural border cell layer, which defines the meningeal dura.

Finally, although follow-up in our patients is limited to less than 2 years, none of the patients suffered the sequelae, namely vasospasm and delayed ischemia, typically associated with aneurysmal SAH. Just the opposite, patients had a remarkably benign hospitalization and convalescence, entirely consistent with the natural history of nonaneurysmal subarachnoid bleeds and prior literature on SRH.

Conclusions

Spontaneous retroclival hematoma, often associated with IVH, is a rare subtype of intracranial hemorrhage frequently recognized only when MRI demonstrates compartmentalization of the posterior fossa hemorrhage. Within the context of posterior fossa hemorrhage, catheter angiography remains the most important diagnostic investigation. Furthermore, when angiography fails to reveal an underlying lesion, SRH patients, like patients with traditional angiographically negative SAH, enjoy a remarkably good prognosis.

References


Author Contributions

Conception and design: Narvid. Acquisition of data: Narvid. Analysis and interpretation of data: Narvid. Drafting the article: Narvid. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Narvid. Study supervision: Narvid.

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