A Rare Complications of Sickle Cell Disease: Empyema and Chronic Subdural Hematoma

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ABSTRACT

Sickle cell disease is a hemoglobinopathy responsible for many complications that are sometimes frightening. Brain damage is part of one of the complications that can affect the vital and/or functional prognosis of patients. We report two cases of neurological complications, patients with known homozygous sickle cell disease, admitted to the Resuscitation Service of the JRA University Hospital for neurological deficit of recent appearance. One presented a spontaneous bilateral chronic bilateral subdural hematoma, and the other an extradural empyema diagnosed on brain scan. The patients benefited from a neurosurgical intervention with favorable postoperative outcomes, marked by an improvement in neurological symptoms. This form of complication is relatively rare in patients with sickle cell disease. It is favoured by repeated vaso-occlusive crises responsible for vascular fragility following repeated ischemia and inflammatory phenomena. The surgical indication depends on the clinical condition and the radiological aspect.

Keywords: empyema, sickle cell disease, subdural hematoma, surgery

I. INTRODUCTION

Sickle cell disease is an autosomal recessive genetic hemoglobin disorder characterized by the presence of an abnormal hemoglobin (hemoglobin S) [1]. African countries are the most affected countries. In Madagascar, it affects more than 10% of the population [2]. The prevalence found for AS and SS forms was 9.4% and 0.2% respectively [3].

It is a major public health problem and the occurrence of complications jeopardizes the vital and functional prognosis of patients. Its complications are numerous; those of neurological complications are rare and are represented by ischemic stroke, spontaneous hemorrhagic complications and intracranial infections.

II. OBSERVATIONS

A. Case 1

This was a 17-year-old patient, known homozygous sickle cell disease, who presented with a headache of sudden onset with rapidly progressive aggravation. He was treated with PARACETAMOL, but without improvement with worsening of the symptomatology and progressive onset of right frontal swelling (Figure 1).

On physical examination the patient was febrile at 39°C, icteric, with a Glasgow score of 15/15 and there was no neurological deficit. The frontal swelling was painful, of soft consistency associated with homolateral palpebral edema.

Brain CT scan showed an image in favor of a left frontal empyma associated with frontal sinusitis that had intracranial communication (Figure 2) and fluid content of frontal swelling.

Biological examination showed anemia at 5.1g/dL, hematocrit at 13.8%, white blood cells at 9 Giga/L, platelets at 670Giga/L, and Prothrombin at 78%.

An evacuating puncture of the frontal subcutaneous collection was performed and sent for bacteriological examination and antibiotic susceptibility testing; the bacteriological examination identified Providenciacrettgeri infection. A probabilistic antibiotic therapy was started and then secondarily adapted to the antibiogram with a favorable evolution of the clinical symptomatology.
Neurological complications are one of the major causes of morbidity and mortality in sickle cell disease, represented by ischemic stroke, intracranial hematomas and infections [4].

In our first case, it was an extradural empyema type infection and a frontal subcutaneous collection due to the Providenciacattergeri germ, which is a Gram-negative bacterium belonging to the Enterobacteriaceae family. According to the literature, pneumococcus is the first suspected germ before Haemophilus influenzae B in cases of intracranial infection [5]. Infections are common in sickle cell disease and can be severe. This susceptibility to infections can be promoted by [5]: functional asplenism, alteration of the complement pathway, which are proteins that participate in the phagocytosis of intruding germs, repeated vaso-occlusive crises that lead to venous stasis and promote infections, especially of the skin.

In the second case, it was a rare intracranial complication of a type of bilateral spontaneous chronic subdural hematoma in a patient with a history of ischemic stroke. Ischemic stroke accounts for 50-80% of cases and increases with age [7]. The mechanism of a chronic subdural hematoma may be secondary to cerebral atrophy, leading to stretching of the cortico-dural vein, which is responsible for bleeding (found in the reported case). According to the literature, risk factors for bleeding in children with sickle cell disease are favoured by recent blood transfusion, high blood pressure and corticosteroid or non-steroidal anti-inflammatory drug use [8]. In the reported case, this patient has a history of recent repeated transfusion. On the other hand, a subdural hematoma may be secondary to a vaso-occlusive mechanism but often associated with bone damage and extradural hematoma [9]. The inflammatory phenomenon secondary to the adhesion of sickle cell adhesion to the vascular endothelium, which will be responsible for vascular fragility, is also responsible for the occurrence of this hemorrhagic complication.

IV. CONCLUSION

Sickle cell disease is a challenge in terms of public health, both because of its complications, but also and especially because of the prolonged disabilities it is likely to cause. Neurological complications are rare and often serious but should be evoked in front of neurological signs and should benefit from an emergency brain scan.

REFERENCES
