A Case of Cerebral Venous Sinus Thrombosis in a Patient with Sickle Cell Disease

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Abstract
We report a 30 year old woman with sickle cell disease (SCD) who presented with a severe right side headache and neck pain of 1 week duration. Results of a peripheral smear showed sickle cells and sickling test was positive. Magnetic resonance venography showed dural venous sinus thrombosis involving right transverse and sigmoid sinuses with venous infarcts involving right temporal & occipital lobes. The patient was treated with low molecular weight heparin, mannitol and her symptoms and signs resolved.

Keywords: Sickle cell disease, Dural venous sinus thrombosis, Magnetic resonance venography, Low molecular weight heparin.

Key Message: Involvement of Transverse sinus and Sigmoid sinus without involvement of Sagittal sinus is rare.

Introduction
Sickle cell disease (SCD) is a well-known hereditary hematologic disorder of hemoglobin. SCD has been reported as a cause of dural venous sinus thrombosis (DVST) events. The involvement of transverse and sigmoid sinus without involvement of sagittal sinus is rare.

Case Report
A 30 year old woman was admitted with severe right side headache and neck pain of 1 week duration. The character of the pain does not suggest migraine or raised Intracranial tension though the patient gave history of migraine in the past. Patient is a known case of sickle cell disease with two other family members were affected, one of the younger brother died of sickle cell crisis. Patient is recently married and not on any contraceptive. On examination patient was conscious and oriented. She was pale with peri orbital puffiness (Right>Left) with stable vitals. Her neurological examination, cardiovascular examination and respiratory examination did not reveal any abnormality. Her haematological investigation revealed a haemoglobin of 10.1 g/dl with hematocrit of 30.5%, reticulocyte count 0.5% and marked neutrophilic leukocytosis. Peripheral smear showed sickle cells and target cells (figure 1 & 2), sickling test was positive. Her urine examination,
renal function tests, liver function tests were within normal limits. Serology for retrovirus, hepatitis and ANA were negative. Her workup for procoagulant state indicated negative for Antiphospholipid antibodies, antithrombin-3, factor V leiden mutation and Protein C levels. The only abnormality was reduced protein S levels which were low.

MRI, MRV of the patient showed dural venous sinus thrombosis involving right transverse and sigmoid sinuses (figure 3,4,5) with venous infarcts involving right temporal & occipital lobes. A diagnosis of Aseptic dural venous sinus thrombosis due to sickle cell disease was made, possibly due to protein S deficiency and patient was started on low molecular weight heparin and patient improved.

Discussion
Dural venous sinus thrombosis can affect any of the superficial or deep cerebral veins, but most commonly involved one is the sagittal sinus. In our patient right transverse sinus and sigmoid sinus were involved without involvement of
sagittal sinus. Clinical manifestations are nonspecific, they can vary from isolated headache to coma. It may be associated with trauma, infections, renal failure and hematological disorders. In SCD there is an increased adherence of sickle cells to the vascular endothelia. Abnormal platelet activation, endothelial cell damage and activation of cell adhesion molecules, low levels of proteins C and S, and increased antiphospholipids contributes further to thrombogenesis and vascular occlusion in SCD.

In our patient hypercoagulation profile were negative, except for the low levels of protein S. Diagnosis is confirmed by detection of the thrombus in the cerebral venous sinus system and changes in brain parenchyma by imaging. Treatment with Unfractionated heparin and LMWH have been reported to be effective in DVST. Our patient improved clinically with LMWH.

Conclusion
This case is reported for its rare anatomical site and isolated neurological involvement without other systemic complications and successfully treated with Low molecular weight heparin.

References