Superior sagittal sinus thrombosis after spinal anesthesia

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Abstract

Cerebral venous sinus thrombosis (CVT) is an uncommon alarming pathological process. Moreover, women affected more commonly than men and the incidence in annual basis is about 0.9 per 100,000.1,2 It commonly occurs secondarily to a genetic or acquired hypercoagulable state such as pregnancy, malignancy, infection, trauma, mechanical heart valve, and oral contraceptive use.3 Cerebral venous sinus thrombosis has variable clinical presentations. The most common presenting symptom is headache that can be accompanied with vomiting, papilledema, visual disturbance, and/or focal neurological syndrome such as focal neurologic deficit with or without seizures. It can also associate with mental status changes such as stupor or with coma in some occasions.2,3

The positional headache considered as the common presenting neurological symptoms after spinal anesthesia, and it occurs in 10 to 30% of patients. The mechanism behind it, is secondary to persistent cerebrospinal fluid leakage after epidural anesthesia. The course of the headache is usually benign in nature with almost complete recovery a few weeks. A rare complication can occur, which is secondary to brain stem compression such as subdural hygroma and neurological deficits.4 It has been reported that CVT rarely occurs in post-partum women after epidural anesthesia who have a tendency for thrombosis secondary to hypercoagulable state.4 Herein, we report a rare occurrence of superior sagittal sinus thrombosis in a healthy young man without prothrombotic status who underwent a day surgery procedure with microscopic varicocele ligation, which was performed under spinal anesthesia.

Case Report

Patient information and clinical findings. A young 29-year-old healthy man who came
to the Neurology Clinics, United Clinics in Madinah Al-Munawarah, at day three with a persistent positional headache after microscopic varicocelectomy for primary infertility. It was associated with nausea and low back pain at the site of spinal puncture. The headache was global type and improved temporarily with hydration and oral caffeine intake. During follow-up, he presented to the Emergency Department with still recurrent headache and new onset of vomiting and blurred vision at day 10. There was neither history of trauma nor relevant past or family medical history for thromboembolic disorders; he had not been on any medical therapy before. On examination, he was conscious and oriented with a GCS of 15/15. His body temperature was 37.1°C, with a heart rate of 86 beats/min. His blood pressure was 133/85 mmHg, and his respiratory rate was 17 breaths/min. Fundoscopic examination showed bilateral papilledema, and no focal neurological deficits were recognized (Table 1).

**Diagnostic assessment.** Laboratory investigations revealed a sodium level of 131 mEq/L, blood urea and nitrogen levels of 4.5 mg/dL, a creatinine level of 55 micromole/L, and a level of glucose was 109 mg/dL. A complete blood count revealed a normal white blood cell count of 8×10^9/L, a hemoglobin level of 12.9 g/dL, and a platelet level of 356×10^9/L. The liver function test was normal, with an alanine aminotransferase (ALT) level of 33 U/L, aspartate aminotransferase (AST) level of 16 U/L, and an alkaline phosphatase (ALP) level of 103 U/L. Further hematological workup revealed normal levels of protein C and S. His D-dimer level was 2.1 nmol/L, with a fibrinogen level of 3.1 g/L. His partial thromboplastin time was 35 seconds and high prothrombin time was 11.1 seconds. Anticardiolipin and anti-lupus antibodies were negative. Fasting plasma homocysteine level is within normal range (13.4 µmol/L). Antithrombin III activity level is 114% which is within normal limit. The sickle cell screening was negative. The total cholesterol is 6.4 mmol/l, triglyceride 1.85 mmol/l, high density lipoprotein is 1.14 mmol/l, and low-density lipoprotein is 5.1 mmol/l. Further genetic tests that include factor V Leiden and prothrombin gene mutation were negative.

Magnetic resonance (MR) imaging showed increase signal intensity in both T1 weighted images (WI) and T2WI as well as bilateral superficial cortical veins with filling defect as seen in the magnetic resonance venography (MRV); there was no diffusion restriction suggesting venous infarction (Figure 1). Findings are consistent with superior sagittal sinus thrombosis.

**Therapeutic intervention.** He required admission to the hospital to observe and stabilize him. He was started on heparin infusion and warfarin until the target INR level reached 2-3. On day 8 after admission, the patient showed complete resolution of headache and vomiting. He still had a blurred vision and he was started on acetazolamide 250 mg daily dose for 3 months only. His symptoms gradually improving over time until complete recovery by the end of the second month after discharge.

**Follow-up and outcome.** His last follow-up was 4 months after surgery and was asymptomatic and his further genetic tests were unremarkable. His INR was maintained between 2 to 3 through all visits to coagulation clinic. He is going to undergo MR imaging follow-up by the end of the 6th month prior to stop anticoagulation.

| Table 1 - Timetable demonstrating the sequence of events and case progression |
|--------------------------|---------------------------------|-----------------------------|
| **Dates**               | **Relevant past medical history and interventions** | **Interventions** |
| 29-9-2019               | The patient had history of primary infertility with azospermia. |                      |
|                        | He underwent left side microscopic varicocelectomy under spinal anesthesia as day surgical procedure. |                      |
|                        | Unremarkable family and social history. |                      |
| Dates                  | **Summaries from Initial and Follow-up Visits** | **Diagnostic Testing (including dates)** | **Interventions** |
| 2-10-2019              | Admitted to the hospital with drowsines and severe persistent headach for conservative measures. | Complete blood count | Intravenous hydration |
|                        | He was discharged home in days 2 in a good condition. | Serum electrolyte | Intravenous Paracetamol |
| 10-10-2019             | Readmitted to the hospital with Sev headach, vomiting, and papilledema. | MRI and MRV | Intravenous metoclopramide |
|                        | His symptoms was improved and target INR was achieved | Thromboembolic blood work | Non-steroidal anti-inflammatory drug |
| 25-12-2019             |                      | Repeated Thromboembolic work-up | Heparin infusion |
|                        |                      | Continue on Warfarin | Warfarin was started |

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Discussion. The current study reports a case of superior sagittal sinus thrombosis after spinal anesthesia in a patient without prothrombotic risk status, which describes this rare finding in a healthy patient. The current finding was diagnosed by brain MR and MRV imaging studies. Our patient was complaining of severe persistent headache in the first day after surgery. He was readmitted for conservative measures on day 4 after surgery and was started on intravenous fluid and analgesics, with temporary improvement. He was discharged from the hospital in an acceptable condition. Despite these conservative measures, he had progressive severe symptoms after 2 weeks that mandated further imaging studies of the brain, showing superior sagittal sinus thrombosis. He was admitted to the hospital for anticoagulation and further diagnostic thromboembolic workups, albeit with negative results.

Using a large gauge needle and/or multiple punctures during spinal anesthesia will predispose the CSF space to persistent drainage and leak. This condition causes loss of the spinal fluid which can lead to stretching and dilatation of the cerebral vascular systems. Consequently, this will lead to fragility of the venous endothelium secondary to stretching of the cerebral vessels due to negative pressure. Furthermore, reduction in the absorption of the fluid into the sinuses leads to increase the viscosity. It is also proved in the study done by Canhao P in 2005 that the velocity of the venous blood flow is markedly decreased after spinal anesthesia as it was measured by transcranial doppler ultrasound. So, many theoretical processes can contribute to the occurrence of the thrombosis that can explain the condition in our patient who has no prothrombotic risk.

Brain MR and MRV imaging was supportive of the diagnosis, showing classical features of superior sagittal sinus thrombosis and cortical venous thrombosis. The mechanism of the CVT in such patients were poorly understood. However, the mechanism behind thought to be secondary to thrombus in the cerebral veins, that can cause obstruction of the blood drainage from the brain, this in turn can lead to increase venous pressure and affection of the blood-brain barrier. Eventually, this leads to decrease cerebrospinal fluid absorption and elevation of intracranial pressure, which can explain the positional headache and the papilledema.

Different case reports addressed the occurrence of sinus thrombosis in anecdotal cases after spinal anesthesia. However, these reported cases occurred among patients with a high risk for thrombosis who underwent spinal anesthesia. Our patient, however, had no identifiable risk for thrombosis. Mahesh and his associates described a case of A caesarean section was done for a young 31-year-old post-partum woman under spinal anesthesia, in which she developed superior sagittal sinus thrombosis secondary to protein C and S deficiency. In their report, they described the pathophysiological aspect of venous thrombosis after intracranial low-pressure system. Furthermore, they studied 52 cases and categorized them into an obstetric-associated group and a non-obstetric-associated group. This considered in the literature as the largest series describing the natural history of this phenomenon that occurred in presence of at least a single risk factor for sagittal venous sinus thrombosis. Yildiz and his group reported a series of anticoagulation...
three cases with intracranial hypotension after spinal anesthesia followed by CVT. However, all of them had at least one thromboembolic risk factor. Accordingly, previous studies identified different risk factors for cerebral venous sinus thrombosis after spinal anesthesia. However, our study supports the approach to image the brain and to suspect CVT in case of persistent severe symptoms, even when no identifiable risk factor could be recognized.

The use of magnetic resonance imaging (MRI) sequences (such sequences like T1WI, T2WI, fluid-attenuated inversion recovery (FLAIR), and diffusion-weighted imaging (DWI)) in combination with magnetic resonance venography (MRV) have been largely replaced the use of cerebral angiography and computed tomography (CT), which allows the detection of venous occlusion along with any complications secondary to venous occlusion, such as cerebral edema and areas of venous infarction.1-9 The use of MRI allows localization of the thrombus within the venous systems by altering the signal intensity in the MRI sequences secondary to disruption in flow due to thrombus. Thrombus in acute stage can lead to a highly subtle abnormal signal intensity on non-contrast MRI, which can be confused with a normal flow void on T2WI. Although, the isointense signal intensity to brain parenchyma in T1WI and low signal intensity in T2WI indicates that the thrombus in acute stage. When the signal intensity is high in T1WI and low in T2WI this indicate that the thrombus in early subacute phase, while when the signal is high in both T1WI and T2WI this indicates that the thrombus in late subacute phase as seen in our case (Figure 1 a, b &c). The subacute phase takes between 7 to 14 days after that the thrombus become chronic and gives a low to isointense signal in T1WI and high signal in T2WI, which mainly related to vascularization of the connective tissue.1-9 Here, MRI and MRV are not only helpful diagnostic tools, but also allow to assess the improvement and recanalization of the venous system in follow-up patients.

Spinal anesthesia is increasingly used in clinical practice, e.g., in pelvic surgical procedures. The current report elaborates the possibility of such complication after spinal anesthesia in the era of advancement of regional anesthesia for different pelvic procedures, especially for day-surgical procedures on young patients with a high risk for headache after spinal anesthesia. This condition could be decreased with the use of a small gauge needle during spinal anesthesia, avoiding multiple punctures to the canal.

In conclusion, our findings indicate the serious effect of severe persistent spinal leak after spinal anesthesia, which could lead to sinus thrombosis even in a patient without risk for prothrombotic state. Persistent severe symptoms after spinal anesthesia should be recognized by the medical team to proceed for brain imaging to enable timely diagnosis and optimize the medical condition with less complications secondary to superior sagittal sinus thrombosis.

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