Acute intracranial subdural hematoma following incidentally durotomy after revisional lumbar spinal surgery; case report

Sung Hwa Paeng1* and Sae Hoon Im2

Abstract
A fifty-four-year-old man was admitted with left leg severe radiating pain with a history of previous lumbar spinal fusion (L5/S1) presented with upper adjacent level spinal stenosis. He underwent a revision spinal arthrodesis from L4 to L5. Dissection at adhesion lesion in L4/5 level resulted in a unintended 3-mm dural tear with CSF loss. The durotomy was primarily repaired during the surgery. Total perioperative blood loss was 1900 mL and fluid replacement was about 4000 mL. On the fourth postoperative day, the patient had slightly headache. He was treated conservatively (Hydration and NSAIDs). On the tenth day, the patient complained of severe headache with nausea, vomiting. A brain CT was obtained and revealed acute subdural hematoma in left frontoparietal area. We did not have surgery because of no intracerebral mass effect, but follow up brain CT. The hematoma was not increased and resolved. The patient made an uneventful recovery. This case reminds that subdural hematoma can complicate durotomy during spine surgery. Therefore, neurological deterioration in the postoperative period rule out the diagnosis, especially in complicated by dural tears on revisional spinal operation.

Keywords: Acute subdural hematoma, headache, spinal stenosis, spinal arthrodesis

Introduction
Cerebrospinal fluid (CSF) leakage induced headache is often after spinal acupuncture and unintended durotomy and results from on pain sensitive structures within the cranium [1-3]. Moreover, it may tear the intracranial subdural veins and form intracranial subdural hematoma, as a result of traction on cerebral veins after unintended dural tear.

We report a case of intracranial subdural hematoma occurring after a spinal dural tear that was made unintentionally during the course of a posterior laminectomy and facetectomy in adjacent spinal lesion (L4/5 level). The possible pathophysiological mechanisms are discussed.

Case report
A fifty-four-year-old man was admitted with lower back pain and right leg severe radiating pain with a history of previous lumbar spinal fusion (L5/S1) presented with upper adjacent level spinal stenosis. The MRI and CT was shown in Figures 1A and 1B. We planned a revision spinal arthrodesis from L4 to L5. Hypertension and diabetes mellitus had been noted for years and were under regular medical treatment. All laboratory data including coagulation status were within normal limits on admission. Preoperative blood pressure was 150/80 mmHg with heart rate around 85 beats min. Anesthesia was induced with thiopentone and fentanyl. Tracheal intubation was facilitated by succinylcholine. Anesthesia was maintained with isoflurane 4–5% in O2 (300 mL/min). The arterial hemoglobin O2 saturation remained between 96–100% during surgery. The patient was ventilated mechanically and end-tidal CO2 maintained between 36–40 mmHg. The patient underwent posterior laminectomy for L4/5 under the microscoppy and posterior spinal fusion. Dissection at adhesion lesion in L4/5 level resulted in a unintended 3-mm dural tear with CSF loss. The durotomy was primarily repaired with 6-0 prolene watertightly. We reinforced with Tachacomb and fibrin selant (TISSEEL).

A subfascial hemovac drainage system was placed and set to positive pressure. The wound closed layer by layer watertightly. Total Perioperative blood loss was 1900 mL and total intravascular replacement included two unit packed blood cell and about 4000 mL of crystalloids during the surgery. There was continuous CSF-like drainage from the hemovac until seven days postoperatively. Fortunately, the patient did not have leaking fluid from lumbar incision and the patient did not have wound problem after removal of hemovac, which MRI was shown in Figure 2A. On the physical examination, the patient was no neurologic problem. The patient complained of severe headache with the vomiting. Moreover, general weakness
and loss of appetite were noted. He became drowsy. A brain CT and MRI was obtained and revealed subdural hematoma in left frontoparietal area. It was shown in Figures 2B and 2C. We did not operation because of no intracerebral mass effect and follow up brain CT. The hematoma was not increased and resolved. The patient made an uneventful recovery without any neurological sequale.

Figure 1. Preoperative sagittal. (A) and axial (B) lumbar CT showing Previous posterior fusion L5/S1 and foraminal stenosis and calcified hypertrophic ligamentum & facet lesion on left lumbar 4/5 level.

Figure 2. Postoperative axial (A) lumbar MRI reveling the well decompressed foramen but fluid collection in laminectomy site Brain axial (B) CT scan showing the intracranial subdural hematoma and coronal (C) brain MRI showing the acute stage on left frontotemproparietal lesion and no midline shifting.

Discussion
Dural tear is well-known complications of spine surgery, with a prevalence of 1 to 17% [4-7]. From 0.67% to 10% in primary
lumbar degenerative surgery [3], and from 13.2% to 21% in revision lumbar surgery [8]. The causes of dural tear during primary operation include eroded or thin dura, dura adhesion, and redundant dura in severe spinal stenosis patients.

It is particularly prevalent in patients who have epidural fibrosis and scar tissue adherent to the dura during revision of spine surgery [3]. In general, revision surgeries have a greater chance of dural tears than primary surgeries because of adhesions and scarring around the dura. If dura tear develop on spinal surgery, primary repair is still considered the gold standard. Various methods for treating identifiable durotomies exist, including primary repair, fat/muscle graft, collagen matrix onlay, and fibrin glue [9]. Moreover, The operative technique plays a key role in avoiding injury to the dural tissue.

Subdural hematoma on brain has been reported after dural tear with persistent leakage of CSF [10]. However, pathophysiological mechanism is unclear. One hypothesis is that CSF leakage reduces intraspinal as well as intracranial pressure. The CSF dynamics may result in a caudally-directed movement of the spinal cord and brain, which in turn stretches the pain-sensitive structures, dura, cranial nerves and bridging veins [10,11]. Cerebral veins drain into dural sinuses that are adherent to the inner table of the skull. Bridging veins are more fragile in the subdural portion than in the subarachnoid space [12]. Therefore, mechanical stress exerted on the bridging veins will rupture at their weakest point in the subdural space. It is presumed that CSF leaked via a dural fistula created by accidental duralotomy may cause a downward brain displacement, which in turn stretches and tears the bridging veins, and results in an intracranial subdural hematoma. The symptoms of such intracranial hematoma are persistent headache, sometimes accompanied with other symptoms and signs such as vomiting, somnolence, confusion and neurologic mental change. Brain CT will confirm the diagnosis and must be performed as quickly as it is suspected.

Another hypothesis is that dehydration may increase the risk of imbalance between CSF leakage and production [13]. Dehydration induced reduction of brain volume could lead to subdural hemorrhage. Low intracranial pressure, as a result of perioperative dehydration, promotes excessive blood congestion in the bridging veins with consequent dilatation and increasing tension of the vessels, which are further pulled by a caudal displacement of the brain, and consequently more vulnerable to the movement of the brain within its coverings due to the decreased cerebral volume [10]. In our case, total intraoperative blood loss was 1900 mL, fluid replacement totally was 4000mL. Therefore, we could rule out dehydration. Minor head injury may also cause to a subdural hematoma in elderly patients and spontaneous hematoma is also observed in anticoagulopathy patients. In our case, the patient did not have a history of head injury in preoperative period and he did not take anticoagulants and the postoperative coagulation status was within normal limits. During the postoperative hospital care, the patient did not injury of head trauma. If dura tear is detected during surgery, then they can be closed intraoperatively, after which the patients are treated with specialized postoperative care, such as bed rest and vigilance for positional headaches. However, if a dural tear is noted postoperatively, it may or may not be associated with persistent signs and symptoms. Many can result in asymptomatic pseudomeningoceles that can be observed or followed clinically. However, if signs or symptoms after surgery persist, then additional postoperative treatment and evaluation may be needed.

Conclusions
We reminds that subdural hematoma can complicate durotomy during spine surgery. Therefore, neurological deterioration in the postoperative period rule out the diagnosis, especially in complicated patients by dural tears on revisional spinal operations. Therefore, because intracranial hemorrhage can present with variable symptoms, it is important that surgeons recognize and aggressively evaluate early signs of neurologic deterioration to prevent a potentially life-threatening complication.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions

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References


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