Case Report

Spontaneous hemorrhage after external ventricular drain placement in the setting of low factor VII secondary to liver cirrhosis

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ABSTRACT

Background: Alterations in normal coagulation and hemostasis are critical issues that require special attention in the neurosurgical patient. These disorders pose unique challenges in the management of these patients who often have concurrent acute ischemic and hemorrhagic injuries. Although neurosurgical intervention in such cases may be unavoidable and potentially life-saving, these patients should be closely observed after instrumentation.

Case Description: A 57-year-old male with liver cirrhosis secondary to amyloid light-chain amyloidosis was admitted to the intensive care unit for the management of delayed hydrocephalus. An external ventricular drain (EVD) was placed for the treatment and monitoring of hydrocephalus. Five days after EVD placement, a head computed tomography scan revealed a tract hemorrhage. However, on repeated imaging, the size of the hemorrhage continued to increase despite aggressive blood pressure control and several doses of phytonadione. Extensive coagulopathy workup was remarkable for low factor VII levels. In that setting, recombinant activated factor VII was administered to normalize factor VII levels, and the tract hemorrhage stabilized.

Conclusion: To the best of our knowledge, this is the first case of spontaneous hemorrhage after EVD placement in the setting of liver cirrhosis-associated factor VII deficiency. Our case highlights the importance of identifying coagulation disorders in neurosurgical patients at high risk for coagulopathy and closely monitoring them postoperatively.

Keywords: Coagulopathy, Factor VII, Liver cirrhosis

INTRODUCTION

Hematoma is particularly important in neurosurgical patients as minor abnormalities can pose significant bleeding risks leading to worsened morbidity and mortality. As such, coagulopathy disorders should be diagnosed and managed appropriately, especially after instrumentation. Common causes of acquired coagulopathies in neurosurgical patients include antithrombotic use, thrombocytopenia, sepsis, disseminated intravascular coagulation, uremia, and liver disease.¹² These disorders pose unique challenges in the management of neurosurgical patients, who often have other ischemic and hemorrhagic comorbidities. Although surgical intervention in such cases may be unavoidable and potentially lifesaving, these patients should be closely observed after any procedure, including those performed at bedside. Herein, we present a...
A 57-year-old Caucasian male with atrial fibrillation (not on anticoagulation), hypertension, monoclonal gammopathy of undetermined significance, and AL amyloidosis was admitted to the hospital with right-sided weakness and headache. His blood pressure on presentation was 164/78 mmHg, and he was found to have a left parietal intraparenchymal hemorrhage and an incidental right parietal arteriovenous malformation [Figure 1]. He was medicated with aggressive antihypertensive therapy and was discharged to an acute rehabilitation center with stable right-sided weakness (strength diffusely 4/5) within a few days.

He was readmitted 1 month later with severe headaches and worsening right-sided weakness (3-/5 strength) with a blood pressure of 148/73 mmHg. A head computed tomography (CT) revealed an increase in size of the left parietal hemorrhage. At that time, he also experienced epistaxis as well as catheter site bleeding that resolved spontaneously. Laboratory workup showed a stable platelet count of 127 × 10^9/L, normal partial thromboplastin time (PTT) of 30.8, and an INR of 1.4. He was given intravenous (IV) Vitamin K (phytonadione); however, his INR did not normalize. Given the expansion of the tract hemorrhage and his underlying AL amyloidosis with cirrhosis, further coagulopathy testing was pursued. This revealed a low factor VII level (38%), a normal factor II level (69%), a normal factor IX level (69%), a normal factor X level (65%), and elevated factor VIII (166%) and von Willebrand factor (421%) levels. Recombinant activated factor VII (rFVIIa) (20 mcg/kg every 2 h for 3 doses) was administered to promote clot formation and achieve hemostasis. He also had normalization of factor VII levels to 122%. A head CT performed a day later revealed stability of the tract hemorrhage [Figure 2a-d]. Two days later, he developed an acute worsening of examination with fixed pupils and extensor posturing. Repeat imaging revealed development of a new acute subdural hemorrhage [Figure 2e]. He was found to have low factor VII level to 31%. He underwent a left hemicraniectomy and was given rFVII (20 mcg/kg every 2 h for 3 doses) and placed on a continuous infusion of aminocaproic acid at 1 g/h intraoperatively which stabilized the bleeding. He was continued on the aminocaproic acid infusion postoperatively throughout his hospital stay. Despite evacuation of the hematoma and bleeding stabilization, his mental status did not improve. He was transitioned to comfort care measures and passed away back to rehabilitation with an antiepileptic drug regimen of levetiracetam (2000 mg twice daily), clobazam (20 mg twice daily), and lacosamide (50 mg daily).

The patient was readmitted 2 months later to our institution for worsening encephalopathy. A head CT showed a mild increase in ventricular size in addition to the resolving left parietal intraparenchymal hemorrhage. An electroencephalogram and extensive metabolic and infectious workup were unremarkable. Given the possibility of delayed hydrocephalus as explanation for his worsening encephalopathy, the patient was trialed on cerebrospinal fluid diversion with an EVD [Figure 2a]. However, 5 days after EVD placement, routine interval imaging revealed a hemorrhage along the tract of the EVD. The tract hemorrhage continued to increase in size over the next few days [Figure 2b-d]. Laboratory workup revealed mildly low platelets to 110 × 10^9/L, a normal partial thromboplastin time (PTT) of 30.8, and an INR of 1.4. He was given 3 days of IV phytonadione; however, his INR did not normalize. Given the expansion of the tract hemorrhage and his underlying AL amyloidosis with cirrhosis, further coagulopathy testing was pursued. This revealed a low factor VII level (38%), a normal factor II level (69%), a normal factor IX level (69%), a normal factor X level (65%), and elevated factor VIII (166%) and von Willebrand factor (421%) levels. Recombinant activated factor VII (rFVIIa) (20 mcg/kg every 2 h for 3 doses) was administered to promote clot formation and achieve hemostasis. He also had normalization of factor VII levels to 122%. A head CT performed a day later revealed stability of the tract hemorrhage [Figure 2d and e]. Two days later, he developed an acute worsening of examination with fixed pupils and extensor posturing. Repeat imaging revealed development of a new acute subdural hemorrhage [Figure 2f]. He was found to have low factor VII level to 31%. He underwent a left hemicraniectomy and was given rFVII (20 mcg/kg every 2 h for 3 doses) and placed on a continuous infusion of aminocaproic acid at 1 g/h intraoperatively which stabilized the bleeding. He was continued on the aminocaproic acid infusion postoperatively throughout his hospital stay. Despite evacuation of the hematoma and bleeding stabilization, his mental status did not improve. He was transitioned to comfort care measures and passed away back to rehabilitation with an antiepileptic drug regimen of levetiracetam (2000 mg twice daily), clobazam (20 mg twice daily), and lacosamide (50 mg daily).

DISCUSSION

We report a rare case of worsening hemorrhage almost 1 week after EVD placement in the setting of an acquired VII deficiency from liver cirrhosis secondary to AL amyloidosis. Liver cirrhosis is a known cause of coagulopathy which can complicate the postoperative management of a neurosurgical patient. Studies have shown up to 2.4% incidence of major bleeding after invasive procedures in cirrhotic patients. [9,16,26,27] Because several hemostatic factors are produced in the
liver, liver dysfunction can result in several clotting factor deficiencies including II, V, VII, IX, X, XI, Protein C, and Protein S. Factor VII levels tend to decrease earlier than the other Vitamin K-dependent factors due to its extremely short half-life. These lead to elevated PT and INR. Elevation in INR may not actually correlate with bleeding risk, and attempts to normalize INR with IV Vitamin K or fresh frozen plasma may not be sufficient to mitigate bleeding risk.

Over the past decade, viscoelastic testing has been increasing used to more accurately assess coagulation status and guide transfusion therapies in patients with end-stage liver disease. Viscoelastic testing including rotational thromboelastometry (ROTEM), thrombelastography (TEG), and sonoheometry, assesses the properties of whole blood during clot formation to determine homeostatic function. However, its application to perioperative and critical care settings outside of trauma has not been robustly validated.

Lever disease can often lead to low platelet counts and platelet dysfunction. Platelet counts <30 × 10^9/L as well as fibrinogen <60 mg/dL and PTT >100 s have been associated with a high bleeding risk. Although there are no consensus guidelines for the management of cirrhotic patients in the periprocedural period, expert recommendations suggest maintaining hemoglobin >7 g/dL, platelet count >50 × 10^9/L, and fibrinogen >150 mg/dL. In addition, aminocaproic acid administration is also recommended in patients with any uncontrolled bleeding. Aminocaproic acid competitively binds to plasminogen and prevents its binding to fibrin, which prevents conversion of plasminogen to plasmin, thus inhibiting fibrinolysis; it is FDA approved for use in acute hemorrhage associated with cirrhosis.

rFVIIa is only approved for replacement therapy in congenital factor VII deficiency where the recommended dose is 15–30 mcg/kg every 4–6 h until hemostasis is attained. Although there have been reports of off-label use of rFVIIa in intracranial hemorrhage, liver cirrhosis, and trauma, clear indications for use in neurosurgery are limited. rFVIIa is thought to be useful as Factor VIIa binds to tissue factor at local sites of tissue and vascular wall injury, leading to generation of sufficient amounts of thrombin for platelet activation. At pharmacologic doses, rFVIIa can directly activate factor X on platelet surfaces which leads to thrombin burst and accelerated coagulation.

For acquired factor VII deficiency, one study in neurosurgical patients with concomitant liver failure recommended a dose of 70–90 mcg/kg every 2 h until hemostasis has been achieved. In another study, nine patients requiring urgent neurosurgical intervention with elevated INR and PTT received rFVIIa at doses ranging from 40 to 90 mcg/kg. Four patients had abnormal coagulopathy secondary to liver disease. After receiving rFVIIa, all nine patients had normalization of their coagulopathies without any peri- or post-procedural complications.

Randomized clinical trials using rFVIIa in patients presenting with intracranial hemorrhage have yielded mixed results. In one study of anticoagulant-related intracranial hemorrhage, rFVIIa was effective in normalizing INR without producing any thromboembolic complications. In another similar study, rFVIIa was also shown to effectively stabilize intracranial hemorrhage and limit hematoma growth. Contrary to these findings, another study showed that rFVIIa did not produce any significant benefit in terms of clot stabilization or clinical outcomes in patients with spot sign positive intracranial hemorrhage. Although thromboembolic adverse events from rFVIIa are rare, they can be devastating and fatal. Such complications include deep venous thrombosis, arterial thrombosis, myocardial infarction, pulmonary embolism, stroke, and clotted devices.

CONCLUSION

Our case highlights the importance of identifying coagulation disorders in neurosurgical patients at high risk for coagulopathy and closely monitoring them postoperatively. Particularly in cases where coagulopathies are unable to be
reversed, there should be a low threshold to check coagulation factor levels including factor VII. In cases where correction of low factor VII levels do not achieve adequate hemostasis, aminocaproic acid should be considered as it was shown to be effective in our patient. To the best of our knowledge, this is the first case of spontaneous hemorrhage after EVD placement in the setting of liver cirrhosis-associated factor VII deficiency.

Declaration of patient consent

Patient's consent not required as patients identity is not disclosed or compromised.

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Conflicts of interest

There are no conflicts of interest.

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