Thrombotic Thrombocytopenic Purpura: A Rare Case Presenting with Splenic Infarction

Abstract
Microangiopathic hemolytic anemia (MAHA), thrombocytopenia, fever, renal failure, and neurologic symptoms comprise the cardinal features of thrombotic thrombocytopenic purpura (TTP). Etiologies can include medications, infections, cancers, or transplantation. However, recognition of thrombotic thrombocytopenic purpura can be difficult because of the variety of presentations and lack of specific diagnostic criteria. We are presenting a case of acute TTP following a bout of splenic infarction.

This rare case reminds us that splenic infarction can be an atypical presentation of TTP. Prompt recognition of TTP is important because the disease responds well to plasma-exchange treatment which improves patient’s prognosis, and is associated with a high mortality rate when untreated.

Keywords: Thrombotic thrombocytopenic purpura; Splenic infarction; Microangiopathic hemolytic anemia (MAHA)

Abbreviations: TTP: Thrombotic Thrombocytopenic Purpura; HUS: Hemolytic Uremic Syndrome; ITP: Idiopathic Thrombocytopenic Purpura; DIC: Disseminated Intravascular Coagulation; FFP: Fresh Frozen Plasma; vWF: VonWillibrand’s Factor; MRI: Magnetic Resonance Imaging

Introduction
Thrombotic Thrombocytopenic Purpura (TTP) is a hypercoagulable state, in which platelets aggregate and clot in the microvasculature. The majority of cases can be linked to ADAMST13 deficiencies, a protein responsible for cleaving VonWillibrand’s Factor (vWF). Hypo functional ADAMST13 results in un-mitigated vWF activity, leading to excessive platelet aggregation. Increased clotting leads to decreased circulating platelets, hemolysis and anemia. Microvascular infarctions cause non-blanching purpura, renal damage, and cerebral ischemia resulting in a myriad of Central Nervous System manifestations from altered mental status to seizures. While some cases are idiopathic, 10% of TTP diagnoses are associated with sepsis [1] or malignancy [2] as well as enterocolitis, especially from E. coli [3].

Originally, TTP was characterized by a classic pentad of microangiopathic anemia, thrombocytopenia, fever, neural manifestations, and renal damage [4]. However, too many cases were missed so a more liberal diagnostic triad was created: microangiopathic anemia associated fragmented erythrocytes, and thrombocytopenia [5].

Case Description
79 year old Hispanic male with past medical history of HTN and osteoarthritis presented to the emergency room complaining of left side abdominal pain for three days and intermittent confusion. Physical examination was unremarkable apart from fluctuating mental status and left lower quadrant tenderness on palpation of abdomen. Initial laboratory investigations revealed anemia, thrombocytopenia, elevated bilirubin but normal liver enzymes and lipase. The peripheral blood smear showed schistocytes. Imaging studies including abdominal Computed Tomography (CT) scan demonstrated a wedge-shaped hypo-enhancing lesions in the spleen typical for splenic infarcts as shown in Figure 1 and a brain CT scan was unremarkable.

Plasma exchange was initiated immediately along with systemic steroids. Platelets counts improved, the patient remains afebrile and hemodynamically stable but his altered mental status persisted. Further workup for his persistent change in mental status revealed multiple bilateral acute lacunar infarcts on Magnetic Resonance Imaging (MRI) of the Brain as shown in Figure 2, which is likely embolic in etiology.

Figure 1: Computed Tomography scan of the Abdomen showing wedge-shaped hypo-enhancing lesion in the spleen.

After six days, the patient became lethargic, febrile, tachycardia and the platelets count dropped again. Plasma exchange subsequently stopped when positive blood cultures grew a gram-
negative bacteria (E. coli). The patient progressively improved following therapy with cefepime and gentamycin. Plasma exchange resumed and he maintained acceptable platelet count until being discharged from the hospital.

Discussion

Epidemiology

TTP is estimated to have prevalence between 4-11 cases per million [6]. When untreated, TTP has a mortality rate of 90% [7]. However, treated TTP has a survival rate of up to 80% [7]. As TTP is a thrombotic disease, the most common and feared causes of death are cerebral, renal and myocardial infarction [4]. Fifty % of cases will have central nervous complications, such as seizures [8], and some cases have even reported neural sequelae preceding hematological findings [9]. Renal injury (indicated by elevated creatinine) occurs in 33% of patients [7]. While TTP is thought of as an acute process, 25% of cases can be indolent, progressing over weeks [7].

With regards to laboratory findings, 95% of TTP cases will have thrombocytopenia with platelet counts <60,000 [9]. Evidence of hemolysis should be found in the form of hyperbilirubinemia, elevated LDH, schistocytes on peripheral smear, and anemia.

Diagnostic work-up

The differential diagnosis for TTP includes Hemolytic Uremic Syndrome (HUS), Idiopathic Thrombocytopenic Purpura (ITP), Disseminated Intravascular Coagulation (DIC), HELLP syndrome in pregnant patients, as well as secondary causes of TTP. Some drugs are known to cause TTP, such as mitomycin, gemcitabine, and of course anti-platelet drugs such as clopidogrel and ticlopidine [10]. HUS more commonly has dramatic renal involvement, is linked to GI infections, and is not treated by plasma exchange [11]. DIC will have prolonged PT and INR when compared to TTP. ITP will often lack the neurological sequelae indicative of TTP.

Treatment

The first line treatment for TTP is plasma exchange. The objective is to remove the auto antibodies, and replace them with Fresh Frozen Plasma (FFP) [12]. Complete plasma exchange, replacing 1-1.5x the patient’s calculated volume occurs daily until platelets and organ function normalize [13]. This can often take weeks, as the neurological sequelae resolve, then the anemia, and finally renal normalization. LDH is also a keen marker for treatment response, as it indicates hemolysis and tissue damage [14]. If plasma exchange is not possible, FFP infusions are an acceptable bridging therapy until the patient can be transferred [15].

Twenty percent of patients will not respond to plasma exchange [16], making our first line treatment significantly flawed. On a side note, it is postulated that the majority of patients who do not respond to plasma exchange, developed TTP from atypical etiologies such as malignancy [17]. As a second line treatment and for severe cases, immuno suppressants are used. The mainstays of this treatment are steroids and rituximab [17]. Response to treatment with rituximab is usually seen within fourteen days [18]. After platelets have normalized, treatment should be continued for at least 2 more days to insure efficacy [6, 13].

Given the intensity of the treatment, it is not surprising that 26% of patients develop major complications, including infections, venous clots, and hypotension. Two percent develop life-threatening conditions such as a catheter-site hemorrhage, and catheter-related sepsis [19].

TTP is a rare but potentially fatal disorder, prompt recognition is important because the disease responds well to plasma-exchange treatment, but is associated with a high mortality rate when untreated. In the era before effective treatment with plasma exchange, 90 percent of patients with thrombotic thrombocytopenic purpura died from systemic microvascular thrombosis that caused cerebral and myocardial infarctions and renal failure. However, recognition of thrombotic thrombocytopenic purpura can be difficult because of the variety of presentations and lack of specific diagnostic criteria [7].

Conclusion

Our patient presented with abdominal pain secondary to a splenic infarct. Eighty-eight percent of splenic infarcts are caused by hematological problems such as blood cancers, hypercoagulable states, atrial fibrillation and vasculitides [20]. However, very few have reported TTP as a precipitant. Indeed, this case is remarkable as a rare disease with an even more rare presentation. Given the severity, the patient was started immediately on steroids and plasma exchange. The brain MRI revealed one of the dreaded complications listed above: multiple, bilateral lacunar infarcts. Finally, our patient also developed a feared, predictable life threatening complication to the therapy-related E. coli sepsis. Despite the myriad of set-backs, our patient improved and was discharged home.

This case stands apart as a rare presentation of a rare disease. Furthermore, it is a classic example of the importance of vigilance in treating a complex condition, with complications from both the disease, and the treatment. Early identification of TTP and anticipation of the adverse effects of treatment are vital in avoiding the significant morbidity and mortality of this potentially debilitating disease.
References


