A report of disseminated adenocarcinoma presenting as thrombotic thrombocytopenic purpura

Joaquín Valle Alonso, Javier Fonseca, Elisa Lopera Lopera, Miguel Ángel Aguayo, Yelda Hernandez Montes, Jose Carlos Llamas
Departament of Emergency and Critical care Medicine, Hospital Valle de los Pedroches, Pozoblanco, Córdoba, Spain

Abstract

Thrombotic microangiopathies (TMAs) represent a heterogeneous group of diseases characterized by a microangiopathic hemolytic anemia, peripheral thrombocytopenia, and organ failure of variable severity. TMAs encompass thrombotic thrombocytopenic purpura (TTP), typically characterized by fever, central nervous system manifestations and hemolytic uremic syndrome (HUS), in which renal failure is the prominent abnormality. In patients with cancer TMAs may be related to various antineoplastic drugs or to the malignant disease itself. The reported series of patients with TMAs directly related to cancer are usually heterogeneous, retrospective, and encompass patients with hematologic malignancies with solid tumors or receiving chemotherapy, each of which may have distinct presentations and pathophysiological mechanisms. Patients with disseminated malignancy who present with microangiopathic hemolytic anemia and thrombocytopenia may be misdiagnosed as thrombotic thrombocytopenic purpura (TTP). Only a few cases of TTP secondary to metastatic adenocarcinoma are known in the literature. We present a case of a 34-year-old man with TTP syndrome secondary to metastatic small-bowel adenocarcinoma. Patients with disseminated malignancy had a longer duration of symptoms, more frequent presence of respiratory symptoms, higher lactate dehydrogenase levels, and more often failed to respond to plasma exchange treatment. A search for systemic malignancy, including a bone marrow biopsy, is appropriate when patients with TTP have atypical clinical features or fail to respond to plasma exchange.

Introduction

The initial diagnosis of thrombotic thrombocytopenic purpura (TTP) may be uncertain because other disorders that can cause microangiopathic hemolytic anemia and thrombocytopenia, the principal diagnostic criteria, may not be initially apparent. Disseminated malignancy is an important consideration in the differential diagnosis of TTP since cancer has been well described for many years as a cause of microangiopathic hemolytic anemia and thrombocytopenia.7,8 Although in most patients the disseminated malignancy that causes microangiopathic hemolytic anemia and thrombocytopenia is easily recognized, in some patients, the malignancy is not clinically apparent, and therefore TTP is diagnosed and plasma exchange treatment is begun. Failure to diagnose disseminated malignancy exposes the patient to the major risks of plasma exchange and causes delay of appropriate chemotherapy.8 However, failure to urgently initiate plasma exchange treatment in a patient with TTP may result in death. The thrombotic microangiopathies are a group of syndromes characterized by small-vessel thrombosis, microangiopathic hemolytic anemia, thrombocytopenia and organ failure. The most common of these disorders, hemolytic uremic syndrome (HUS) and TTP, were once considered 2 distinct disease etiologies. In consideration of their extensive pathophysiological overlap, they are now termed the HUS/TTP syndrome. The classic clinical pentad of TTP includes i) fever, ii) neurologic symptoms, iii) microangiopathic hemolytic anemia, iv) thrombocytopenic purpura, and v) the presence of thrombi in glomerular capillaries and afferent arterioles. Although hemolytic anemia is a universal feature, all 5 classic findins are present in only 40% of cases. Thrombotic thrombocytopenic purpura is seen more often in women, with most of the patients younger than 40 years, the diagnosis rests on evidence of microangiopathic hemolytic anemia and thrombocytopenia in the absence of DIC or other known causes of thrombotic microangiopathy. There are no clinical features or laboratory tests that can confirm the diagnosis of TTP-HUS; the most important diagnostic criterion, but also the most difficult, is the exclusion of alternative etiologies.9,10 ADAMTS13 (A Disintegrin And Metallo-protease with ThromboSpondin-1-like domains), an enzyme required for normal proteolytic processing of von Willebrand factor (VWF), is important in the pathophysiology of TTP, but patients may have characteristic presenting features and clinical courses without severe ADAMTS13 deficiency.11,12 Even after a diagnosis of thrombotic thrombocytopenic purpura is made, continuing evaluation is important. In the Oklahoma Thrombotic Thrombocytopenic Purpura-Hemolytic Uremic Syndrome (TTP-HUS) Registry, 10 percent of patients with an initial diagnosis of idiopathic thrombotic thrombocytopenic purpura were subsequently found to have sepsis or systemic cancer (Table 1).
serum sodium 139 mg/dL, serum potassium 6.2 mg/dL, total bilirubin 6.8 mg/dL, aspartate aminotransferase 5318 U/L, alanine aminotransferase 1947 U/L, lactate dehydrogenase 19,758 U/L. Blood coagulation tests; prothrombin time; 23.1 sec, activated partial thromboplastin time 53.2 sec.; Fibrinogen 3.9 g/L (2.0-6.0) all within normal values; D-dimer 5250 ng/dL. A Coomb's tests showed a negative result. Toxicologic screening of serum was negative for ethanol, acetonaphenin, and salicylates, and toxicologic screening of urine was also negative. Blood cultures failed to grow any organism. Antibody titer for human immunodeficiency virus was normal. A chest radiograph was normal, and a bedside echocardiogram showed no pericardial fluid. Specimens of blood and urine were taken for bacterial and viral cultures and testing for viral antigens. Toraco-abdominal computed tomography (CT) revealed enlarged esophageal and para-cardiac lymph nodes, bilateral posteriorabdominal lung consolidations, hepatosplenomegaly, multiple enlarged retroperitoneal and paraeaortic lymph nodes (Figure 1).

A definite diagnosis could not be determined on admission because of the complexity of signs and symptoms, the diagnosis of TTP was suggested based on the presence of fever, neurologic symptoms, microangiopathic hemolytic anemia, thrombocytopenia, renal failure, increased serum levels of lactate dehydrogenase and indirect-reacting bilirubin, and negative direct Coomb's test. Plasma exchange therapy and supportive care was begun. It was critical to rule out disseminated cancer. The importance of prompt diagnosis of the systemic malignancy was to provide an opportunity for treatment with appropriate chemotherapy. In the Intensive Care Unit treatment with dopamine to maintain a systolic blood pressure of 140 was started, and aggressive fluid resuscitation was continued. Two hours later the blood pressure decreased to 90/45 mm Hg; Norepinephrine was added. After obtaining informed consent from his family bone marrow aspiration sample was obtained. Twenty-nine hours after presentation the axillary temperature was 40.4°C (104.8°F) and the blood pressure 82/55 mm Hg despite increasing doses of dopamine, epinephrine, and norepinephrine. Laboratory evaluation demonstrated severe anemia (hematocrit 23%) despite blood transfusion, thrombocytopenia (platelet count 20,000/mL), coagulation tests (international normalized ratio (INR) 1.2; activated partial prothrombin time, 25 s; fibrinogen concentration, 3.3 g/L (2.0-6.0) were normal. Results of Microbiologic and Serologic Tests were received, Blood culture negative, Urine culture negative, Cytomegalovirus antigenemia assay negative, Epstein-Barr virus anti-VCA IgG positive, Epstein-Barr virus anti-VCA IgM negative, Epstein-Barr virus early antigen negative, Epstein–Barr virus antinuclear antigen negative, Herpes simplex virus type 1 antibody IgG positive, Herpes simplex virus type 2 antibody IgG negative, Carcinoembryonic antigen (ng/mL) 1.4, Prostate-specific antigen (ng/mL) 4.6, CA-125 (U/mL) 36, CA 19-9 (U/mL) 6.1, Rheumatoid factor (IU/mL) <30, Antinuclear antibody negative, Antineutrophil cytoplasmic antibody Negative, Hepatitis B surface antigen Negative, Hepatitis C antibody negative, ADAMTS13 (< 20%) Thirty-one hours after presentation, bradycardia developed, followed rapidly by asystole, and cardiopulmonary resuscitation was initiated. The patient was pronounced dead 32 hours after arrival in the emergency room. The marrow aspirate results were received with occupation by neoplastic cells of large size and frequent syncytia. The autopsy revealed a tumor at the ileum 10 cm proximal from the ileocecal region. Peritoneal dissemination was recognized around the ileocecal region, pathological diagnosis of the specimen was adenocarcinoma with lymph nodes metastasis.

## Discussion

Microangiopathic hemolytic anemia and thrombocytopenia caused by systemic malignancies have been well described, but it is uncommon for microangiopathic hemolytic anemia and thrombocytopenia to be the predominant presenting clinical features in patients whose systemic malignancy is not ini-

### Table 1. Clinical features that may suggest disseminated malignancy as an alternative diagnosis in patients with assumed TTP-HUS.

<table>
<thead>
<tr>
<th>Clinical feature</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>History of cancer</td>
<td>Even when the clinical evaluation and results of imaging studies are normal metastatic cancer must be suspected</td>
</tr>
<tr>
<td>Pulmonary infiltrates</td>
<td>Rare in TTP-HUS</td>
</tr>
<tr>
<td>Disseminated intravascular coagulation (DIC)</td>
<td>Although DIC is commonly associated with metastatic carcinoma, coagulation tests may be normal.</td>
</tr>
<tr>
<td>Nucleated red blood cells and immature myeloid cells on peripheral blood smear</td>
<td>These abnormalities may accompany the marrow response to severe hemolysis, but they commonly indicate marrow infiltration by tumor.</td>
</tr>
<tr>
<td>Extreme elevation of lactate dehydrogenase (LDH) level</td>
<td>Although an elevated LDH level, caused by hemolysis and tissue ischemia, is characteristic of TTP-HUS, levels exceeding 5,000 U/L are more commonly cause by tumor lysis</td>
</tr>
<tr>
<td>No response to plasma exchange therapy</td>
<td>Patients with TTP-HUS typically respond promptly to plasma exchange. No response should cause concern about the diagnosis.</td>
</tr>
<tr>
<td>Extreme elevations of D-dimer</td>
<td>D-dimers were found severely increased in all cases of disseminated cancer presenting as TTP</td>
</tr>
<tr>
<td>ADAMTS13 activity</td>
<td>ADAMTS13 activity may be normal or lower than normal in cancer-associated TTP</td>
</tr>
</tbody>
</table>

### Figure 1. Abdominal CT scan demonstrates A) pathological hepatic lymph nodes and B) retroperitoneal lymph nodes.
tially apparent. Although occult malignancy causing microangiopathic hemolytic anemia and thrombocytopenia may be uncommon, it is an important consideration in the evaluation of patients for TTP. In the Oklahoma TTP-HUS Registry, 10 (3%) of 351 patients who were initially diagnosed as having TTP and treated with plasma exchange were subsequently and unexpectedly diagnosed with disseminated malignancy. Only systemic infections have been a more common cause of an incorrect initial diagnosis of TTP in the Oklahoma Registry. Many different malignancies may mimic TTP; the importance of prompt diagnosis of the systemic malignancy is to provide an opportunity for treatment with appropriate chemotherapy. Early recognition of cancer may not benefit many patients who present with microangiopathic hemolytic anemia and thrombocytopenia since these patients often have widely disseminated cancer. Even though treatment success may be limited, prompt diagnosis is important for appropriate management. Prompt diagnosis of systemic malignancy is also important in avoiding unnecessary risks of plasma exchange treatment for TTP. The difficulty of diagnosing a systemic malignancy presenting with microangiopathic hemolytic anemia and thrombocytopenia is evident in literature. The diagnosis of disseminated malignancy excludes the diagnosis of TTP or HUS; these patients should not be considered to have cancer-associated TTP. Although multiple etiologies may contribute to the syndromes recognized as TTP and HUS, disseminated malignancy is a pathologically and clinically distinct disorder. Disseminated malignancy can cause microangiopathic hemolytic anemia and thrombocytopenia, in the absence of DIC, by microvascular tumor emboli. This has been most frequently observed with diffuse microscopic pulmonary involvement. ADAMTS13 activity is not severely deficient but may be lower than normal in some patients with disseminated malignancy due to high plasma levels of von Willebrand factor. Plasma exchange has no role in management when a malignant disorder is recognized. Clues that may suggest the presence of an occult systemic malignancy include presenting symptoms of dyspnea, cough, and pain other than abdominal pain. Although increased serum LDH is characteristic of patients with TTP, extreme elevations are not typical and may suggest tumor lysis. Median LDH level was 4.5 times upper the normal values (range, 3.2–8.9). D-dimers were found severely increased in all cases of disseminated cancer presenting as TTP in a recent report of several cases of disseminated cancer. Perhaps the most convincing clue that a patient with presumed TTP may have a disseminated malignancy is failure to respond to plasma exchange. If systematic malignancy is suspected, bone marrow biopsy is appropriate.

### Conclusions

The evaluation and management of patients who present with an acute onset of microangiopathic hemolytic anemia and thrombocytopenia remain critical challenges for clinicians. Although the diagnosis of TTP and urgent treatment with plasma exchange must be considered in patients with microangiopathic hemolytic anemia and thrombocytopenia, the possibility that all of the clinical features of TTP may be caused by an occult disseminated malignancy must be appreciated. With increased awareness of the possible diagnosis of disseminated malignancy, the diagnosis can be made sooner, avoiding unnecessary plasma exchange treatment and allowing appropriate chemotherapy treatment.

### References