A Signet Ring Cell Carcinoma Presented as Refractory Acquired Thrombotic Thrombocytopenic Purpura

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Keywords
Microangiopathic hemolytic anemia · Thrombotic thrombocytopenic purpura · Signet ring cell carcinoma

Abstract
Microangiopathic hemolytic anemia (MAHA) can be observed as a paraneoplastic syndrome (PS) in certain tumors. MAHA-related signet ring cell carcinoma (SRCC) of an unknown origin is very infrequent. Herein we present a SRCC case presented with refractory acquired thrombotic thrombocytopenic purpura (TTP). A 35-year-old man applied to the emergency service with fatigue and headache. His laboratory tests resulted as white blood cell 9,020/µL, hemoglobin 3.5 g/dL, platelet 18,000/µL. Schistocytes, micro-spherocytes, and thrombocytopenia were observed in his blood smear. MAHA was present and he was considered as having TTP. Plasma exchange treatment was initiated; however, he was refractory to this treatment. Thorax and abdomen computerized tomography revealed thickening of minor curvature in stomach corpus with hepatogastric and paraceliac lymphadenopathy. Bone marrow (BM) investigation by our clinic resulted as the metastasis of adenocarcinoma. Ulceration and necrosis were observed by gastric endoscopy procedure. Biopsy was taken during endoscopic intervention, which resulted as SRCC. MAHA may be seen as a PS in some tumors, especially gastric cancers. Tumor-related MAHA is generally accompanied by BM metastases. As a result, BM investigation may be used as the main diagnostic method to find the underlying cancer. The clinical course of cases with tumor-related MAHA is usually poor, and these cases are usually refractory to plasma exchange treatment. In conclusion, physicians should suspect a malignancy and BM involvement when faced with a case of refractory TTP.

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Introduction

Acquired thrombotic thrombocytopenic purpura (TTP) frequently presents as severe microangiopathic hemolytic anemia (MAHA) and thrombocytopenia in a previously healthy individual. Fatigue, dyspnea, petechiae, or other hemorrhagic events are usually the initial symptoms of MAHA and thrombocytopenia [1]. MAHA can appear as a paraneoplastic syndrome in certain tumors. MAHA is a late and mortal complication of metastatic cancer [2]. The most common tumors related with MAHA are gastric, breast, and lung cancers and malignancies of unknown origin [3, 4]. However, MAHA-related signet ring cell carcinoma of unknown origin is very infrequent [5, 6]. Herein we present a signet ring cell carcinoma case presented with refractory acquired TTP.

Case Report

A 35-year-old man applied to the emergency service with fatigue and headache on January 2020. In his anamnesis, he had a history of alcoholic pancreatitis. His physical examination was normal except for the neurological symptoms, which were temporary loss of consciousness and disorientation. His laboratory tests resulted as white blood cell 9,020/µL, hemoglobin 3.5 g/dL, platelet 18,000/µL, MCV 110.7 fl, urea 58 mg/dL, creatinine 0.84 mg/dL, AST 68 U/L, ALT 33 U/L, indirect bilirubin 1.88 mg/dL, LDH 2,257 U/L, reticulocyte %10.1, haptoglobin <8 mg/dL, INR 1.42, prothrombin time 13.2, fibrinogen 184 mg/dL, Coombs negative. He had consulted our clinic with bicytopenia and hemolysis. Schistocytes, micro-spherocytes, and thrombocytopenia were observed in his blood smear (Fig. 1). Microangiopathic hemolytic anemia was present and he was considered as having TTP. Plasma exchange treatment was initiated; however, he was refractory to this treatment. He had epistaxis and blurred vision during the follow-up. Superficial hemorrhages on the edges of the optic disc and Roth spots were detected. Pain had emerged in his right arm. Doppler ultrasonography revealed the occlusion of cephalic vein with non-recanalized thrombus in the subacute process from the antecubital level at the forearm level. Thorax and abdomen computerized tomography (CT) resulted as liver 220 cm, spleen 14 cm, minimal pleural effusion, thickening of minor curvature in stomach corpus with hepatogastric and paraceliac lymphadenopathy. As a result of CT, endoscopic examination was planned. Bone marrow

Fig. 1. Schistocytes, micro-spherocytes and thrombocytopenia were observed in the blood smear.

Fig. 2. Bone marrow biopsy under ×40 magnification showing signet ring cells next to bone marrow trabecular (H&E ×400).
investigation by our clinic resulted as the metastasis of adenocarcinoma (Fig. 2, 3). Ulceration and necrosis were observed by gastric endoscopy procedure. Biopsy was taken during endoscopic intervention, which resulted as signet ring cell carcinoma (Fig. 4, 5). He was transferred to oncology clinic for his treatment. Unfortunately, he died in 1 month after his transfer.

**Discussion**

It is an important aspect in the clinical practice that not all patients with TTP are critically ill. TTP patients may present with minor complaints of weakness and dizziness, abdominal pain, easy bruising, or nausea and vomiting. Moreover, in some patients, the diagnosis of TTP
may not be made until the laboratory test reveals severe thrombocytopenia and MAHA. Only 40% of TTP cases have the complete pentad and in 75% of the cases there is a triad of microangiopathic hemolytic anemia, thrombocytopenia, and neurological findings [7]. At present, the diagnosis of TTP is considered with microangiopathic hemolytic anemia and thrombocytopenia when any other cause is excluded [7]. In our case there was no acute kidney failure; however, all the other features favored TTP, and the diagnosis was made without kidney failure. MAHA can be observed as a paraneoplastic syndrome in some malignant conditions, particularly gastric tumors. The mechanisms of tumor-related MAHA include tumor-derived factors, procoagulants, immune complexes, some chemotherapeutic agents, fibrinoid necrosis of bone marrow, and tumor cell emboli of arteries, arterioles, and capillary [8]. Cancer-related MAHA is generally accompanied by multiple bone or bone marrow metastases [3]. Therefore, bone marrow investigation is useful as a primary diagnostic tool to examine the underlying cancer. Similarly, in our case, bone marrow investigation revealed the metastatic adenocarcinoma. Total plasma exchange is usually performed in the treatment of cancer-associated TTP; however, less than 20% of the cases respond to plasma exchange. Likely, our case did not respond to plasma exchange treatment either. The clinical course of cancer-associated TTP is generally poor. Patients are generally lost within 2 months due to multiorgan failure. Similarly, our patient had died soon after the diagnosis.

To summarize, MAHA may be seen as a paraneoplastic syndrome in some tumors, especially gastric cancer. Tumor-related MAHA is generally accompanied by bone marrow metastases. As a result, bone marrow investigation may be used as the main diagnostic method to find the underlying cancer. The clinical course of cases with tumor-related MAHA is usually poor, and these cases are usually refractory to plasma exchange treatment. In conclusion, physicians should suspect a malignancy and bone marrow involvement when faced with a case of refractory TTP.

Statement of Ethics

Written informed consent was obtained from the patient for the publication of the case report and any accompanying images.

Conflict of Interest Statement

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Author Contributions

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