Case Report

A Case of Acute Myocardial Infarction in a Patient Whose Initial Complaints Were Hematemesis and Epigastric Discomfort

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The patient was a 64-year-old woman with systemic lupus erythematosus, thrombophlebitis of the lower legs, cerebral infarction with left hemiparesis, and colostomy after perforation of the sigmoid colon. On the morning of her presentation, the patient felt epigastric abnormality. Thereafter, hematemesis occurred twice, leading her to call an ambulance in the afternoon. Upon arrival, electrocardiography before securing a venous route and obtaining blood samples revealed ST segment elevation in leads II, III, and aVF. As her vital signs were stable and her hemoglobin level had decreased by just 1.1 g/dl in comparison to the previous day, emergency coronary angiography (CAG) was performed. CAG revealed complete occlusion at section #4. She underwent right coronary angioplasty with stent placement. The patient's course after angioplasty was uneventful. On the 15th hospital day, esophagogastroduodenoscopy revealed esophageal erosion and superficial gastritis. She was discharged on foot the following day.

When physicians treat patients with hematemesis, electrocardiography and the measurement of troponin are essential before esophagogastroduodenoscopy.

1. Introduction

Complication between gastroduodenal bleeding and acute coronary syndrome is not uncommon [1–4]. We report the case of a patient with acute myocardial infarction whose initial complaints were hematemesis and epigastric discomfort.

2. Case Report

The patient was a 64-year-old woman with systemic lupus erythematosus, thrombophlebitis of the lower legs, cerebral infarction with left hemiparesis, and colostomy after perforation of the sigmoid colon. She was treated with prednisolone, tacrolimus, mizoribine, edoxaban, limaprost, famotidine, sulfamethoxazole-trimethoprim, sertraline, eszopiclone, and minodronic. On the morning of her presentation, the patient felt epigastric abnormality. Thereafter, hematemesis occurred twice, leading her to call an ambulance in the afternoon. Upon arrival, her vital signs were as follows: Glasgow Coma Scale, E4V5M6; blood pressure, 110/76 mmHg; pulse rate, 78 beats per minute; and her peripheral oxygen saturation on 6 liters of oxygen per minute, 98%. A physiological examination revealed preexisting bilateral leg edema with pigmentation and left hemiparesis. Electrocardiography before securing venous route and blood examination revealed ST segment elevation in leads II, III, and aVF (Figure 1). Chest roentgenography showed cardiomegaly and cardiac ultrasound showed hypokinesis at the inferior wall. The results of a biochemical blood analysis on arrival were as follows: white blood cell count, 11,500/µL; hemoglobin, 9.6 g/dL; platelet count, 16.8 × 10^4/µL; total protein, 6.1 g/dL; total bilirubin, 0.5 mg/dL; aspartate aminotransferase, 86 IU/L; alanine aminotransferase, 8 IU/L; blood urea nitrogen, 13.7 mg/dL; creatinine, 0.49 mg/dL; sodium, 143 mEq/L; potassium, 3.6 mEq/L; chloride, 106 mEq/L; creatine phosphokinase, 1000 IU/L; troponin T, 13250 (26.2 >) pg/mL; prothrombin time, 12.7 (11.7) s; activated partial thromboplastin time, 30.1 (30.2) s; fibrinogen, 326 mg/dL; and D-dimer, 0.79 µg/mL. She was diagnosed with acute myocardial infarction with upper esophagogastroduodenal bleeding. As her vital signs were stable and her level of hemoglobin decreased by just 1.1 g/dl in comparison to the previous day when she had visited.
the dermatology department of Numazu City Hospital, she underwent emergency coronary angiography (CAG). CAG demonstrated 99% stenosis at section #2, complete occlusion at section #4 (Figure 2), and 75% stenosis at section #6. She underwent right coronary angioplasty with stent placement. The prescriptions of edoxaban and sertraline were stopped, famotidine was switched to lansoprazole, and treatment with aspirin, clopidogrel, rosuvastatin, and carvedilol was initiated. After angioplasty, her course was uneventful. Her creatinine kinase level peaked at 5655 IU/L on the 2nd hospital day, and her minimum level of hemoglobin was 8.3 g/dl on the 7th hospital day without transfusion. On the 15th hospital day, esophagogastroduodenoscopy revealed esophageal erosion and superficial gastritis (Figure 3). She was discharged on foot the following day.

3. Discussion

Esophagogastroduodenoscopy revealed esophageal erosion and gastritis; thus, we hypothesized that the patient’s initial epigastric discomfort was induced by acute myocardial infarction, following stress-induced esophageal erosion and gastritis. However, it is possible that preexisting bleeding from esophagogastric lesions induced acute myocardial infarction due to a mismatch from decreasing oxygen transport ability and increased cardiac oxygen consumption capacity due to anemia.

The typical complaint of patients with acute myocardial infarction is chest pain; however, acute myocardial infarction is also associated with atypical complaints or signs such as left shoulder pain, pharyngeal pain, earache, headache,
back pain, syncope, and dyspnea. In cases involving atypical complaints, signs, or symptoms, the diagnosis and treatment tend to be delayed, resulting in an unfavorable outcome [5, 6]. It is not uncommon for patients with acute myocardial infarction to be complicated by esophagogastrduodenal lesions or patients with esophagogastrduodenal lesions to be complicated by acute myocardial infarction. Accordingly, when physicians treat such patients, both possibilities should be excluded as soon as possible. For example, in case of patients with hematemesis, it is essential to perform electrocardiography and measure the troponin level before esophagagogastroduodenoscopy, similarly to our case.

The optimal treatment strategy for the combined acute myocardial infarction and hematemesis is controversial because the standard treatments for these diseases are totally different [7]. Acute myocardial infarction requires antiplatelet therapy, which may lead to the deterioration of hematemesis. In contrast, hematemesis requires treatment to achieve hemostasis, which may lead to the deterioration of acute myocardial infarction. Accordingly, these complications can lead to unfavorable outcomes. Our treatment strategy fortunately resulted in a favorable outcome; however, further examinations are needed to select appropriate treatments for patients with the combination of acute myocardial infarction and hematemesis.

Conflicts of Interest

The authors declare no conflicts of interest in association with the present study.

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