A case of juvenile dermatomyositis with esophago-pleural fistula caused by spontaneous lower esophageal perforation

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(Received November 28, 2012: Revised December 5, 2012: Accepted December 12, 2012)

Abstract

Juvenile dermatomyositis (JDM) is a multisystem disease of uncertain origin that results in nonsuppurative inflammation of striated muscle and skin. Gastrointestinal perforation including esophagus is a well-recognized complication of JDM. A one-year and nine-months-old girl with JDM was recently seen with esophago-pleural fistula. The fistula was caused by spontaneous lower esophageal perforation near the gastro-esophageal junction. The treatment consisted of adequate drainage of perforation site, total parenteral nutrition, long-term intravenous hydrocortisone, and broad-spectrum antibiotics. Although the duration of the fistula closing was long, the fistula spontaneously obliterated without other surgical management. This is a rare case of JDM with esophago-pleural fistula caused by spontaneous lower esophageal perforation. (J Med Life Sci 2012;9(2):110-113)

Key Words : Juvenile dermatomyositis, fistula, esophageal perforation

Introduction

Juvenile dermatomyositis (JDM) is a chronic multisystem disease characterized by nonsuppurative inflammation of striated muscle and skin. Gastrointestinal perforation including esophagus is a well-recognized complication of JDM. Although the overall prognosis of JDM is relatively good without any adverse factor, gastrointestinal perforation indicates a poor prognosis. When it sometimes rapidly leads to death, it is frequently due to underlying vasculopathy of the gastrointestinal perforation.

This report was described that a 1-year-old JDM girl with esophago-pleural fistula by spontaneous lower esophageal perforation was successfully treated with adequate drainage of perforation site and long-term hydrocortisone therapy.

Case Report

A girl, aged one year and nine months, was referred for further evaluation of respiratory distress. Three months before admission, she began to unable to stand herself and thereafter she gradually developed muscle weakness, skin ulcerations. Two months later, she was admitted to other hospital because she developed dysphasia to solid food, hypotonia, and aggregated skin ulcerations. At that time, physical examination showed decreased gag reflex, decreased muscle power, and skin ulcerations over left upper eyelid, right axilla, and perianal region. Laboratory findings were elevated muscle enzymes (CPK 487 IU/L, LDH 1,541 IU/L, aldolase 16 IU/L, AST 110 IU/L, ALT 48 IU/L), negative antinuclear antibody and anti-PM-Scl antibody, and negative viral markers (Coxsackie B, rubella, EBV, and measles). Muscle biopsy showed small groups of necrotic fibers, perifascicular and perivascular inflammatory cell infiltration, endomysial and perilysial fibrosis. Because of the proximal muscle weakness, elevated muscle enzymes levels, and histologic findings, a diagnosis of JDM was established according to the criteria of Bohan and Peter.

After about one month, she abruptly developed respiratory difficulty, and decreased oxygen saturation. Chest plain film revealed the fluid collection of left lung field and pneumothorax (Fig 1). Chest tube inserted under the impression of pyopneumothorax and she was referred to our hospital.
On admission, she was acutely ill with marked respiratory distress: respiration, 50/min and body temperature, 38°C. Findings of physical examination were decreased breath sound at left lung field, a chest tube in left hemithorax with continuous drainage, and skin ulcerations over left upper eyelid, both axilla, right post-auricular and perianal region. Muscle mass, muscle tone, and muscle power were also decreased. Laboratory studies revealed a leukocyte count of 4,860/mm³, hemoglobin of 14.1 g/dL, platelet count of 265,000/mm³, erythrocyte sedimentation rate (ESR) of 35 mm/hr (Westergren method) and positive CRP. Electrolytes and renal function were normal, with albumin of 3.7 g/dL. The laboratory findings included WBC 10,550/mm³, hematocrit 35.9%, total protein 5.0 g/dL, albumin 2.3 g/dL, AST 45 IU/L, ALT 33 IU/L, CPK 70 IU/L, LDH 336 IU/L, aldolase 8.5 IU/L. The erythrocyte sedimentation rate and the levels of C–reactive protein were increased. She was treated with stress–dose intravenous hydrocortisone and Gram positive bacteria–directed antibiotics. The liquid diet fed via nasogastric tube. Until second hospital day, chest tube drained 1,500–1,800 mL/day. Esophagography revealed the contrast dye leaked to left pleural space through the esophagopleural fistula near the gastroesophageal junction (Fig 2). However, she could not take surgical intervention because her general condition was poor. She received total parenteral nutrition, intravenous hydrocortisone, and antibiotics with adequate drainage of perforation site. Esophagography was followed up one month later and showed obliteration of the fistular tract. She began a liquid diet and later progressed to a regular diet with no subsequent problem. She was discharged after one month. On follow-up 2 months later, she was tolerating oral intake without other problems.
Although the precise pathogenesis of JMD is unknown, recent studies suggested autoimmune play an important role in pathogenesis. Immune complex-mediated vasculitis may be an initiating factor in JMD, and the vasculopathy affects striated muscles, skin, subcutaneous tissues, and gastrointestinal tract. Gastrointestinal perforation in JDM caused by this vasculopathy can occur in any part of the gastrointestinal tract. Wedgewood and et al. reported an underlying vasculopathy as the cause of gastrointestinal ulceration in JDM. Banker and Victor noted a multifocal, submucosal, and noninflammatory endarteropathy with superimposed thrombosis was responsible for the infarcted tissues of gastrointestinal perforation.

Corticosteroid therapy in JMD may be the cause of the ulceration and perforation of the gastrointestinal tract regardless of the disease course. However, previous reported cases presented that gastrointestinal ulcerations in dermatomyositis developed before the corticosteroid therapy or after a very short time of the therapy. Moreover, gastrointestinal ulcerations have described long before the introduction of the corticosteroid therapy. The corticosteroid therapy in JDM is not regarded as an essential factor in the gastrointestinal ulceration and perforation. In this case, she was diagnosed with JDM two months before. During admission of other hospital, she was given antacid medication for the prevention of gastrointestinal ulceration and the perforation developed one month later. No definitely precipitating factors in her hospital course were attributed to esophageal perforation except disease itself.

The surgical management of gastrointestinal perforation including esophagus is dependent on location and extent of the perforation. In this case, the perforation was lower esophagus near the gastroesophageal junction with the esophago-pleural fistula. The fistula connected with left pleural throughout left subphrenic space. However, she could not take surgical management for poor condition, we chose to manage our patient by adequate drainage of perforation site, TPN, low dose hydrocortisone, and broad-spectrum antibiotics. After one month, the fistula was spontaneously obliterated without any surgical management. She was tolerating oral intake without other problems on follow-up 2 months later.