A Case of Dermatomyositis and Ulcerative Colitis

Pedro Rosales Torres, 1* Rafael Pila Pérez, 1 Rafael Pila Peláez, 1 Javier Artola González. 1

Abstract

Introduction: The incidence of idiopathic inflammatory myopathy ranges from 4 to 15 cases per million inhabitants while its prevalence is 60 cases per million inhabitants. Given the identification of clinical forms, it is possible that these figures will increase in the coming years. Incidence and prevalence of ulcerative colitis have been increasing, and it now affects 2% of the world population. Symptoms are determined by the extent and severity of lesions. It is chronic with recurring outbreaks, and there is no medical cure. Dermatomyositis in association with ulcerative colitis occurs extremely infrequently. Objective: This study presents the case of a patient diagnosed with dermatomyositis associated with ulcerative colitis. Clinical Case: The patient was a 70 years white woman, a housewife with no family or epidemiological history of interest. She reported that she had had diarrhea accompanied by abdominal pain, mucus and pus when she was fifty years old. About one year prior to diagnosis, she developed muscle weakness in her shoulders that made it difficult for her to lift her arms. Laboratory, imaging and histopathological tests showed dermatomyositis associated with ulcerative colitis. We point out clinical characteristics, diagnostic means and the therapy used to treat our patient. Conclusions: This case of a patient with clinical manifestations of the very infrequent combination of dermatomyositis and ulcerative colitis should be taken into account by doctors who care for these patients because both conditions have immunological bases and can be expressed in various ways.

Keywords
Dermatomyositis, ulcerative colitis.

INTRODUCTION

Polymyositis was described by Wagner in 1882, and in 1887 Ulverrick invented the name dermatomyositis (DM) for the association of muscle pain and weakness with inflammatory skin lesions. Hachulla has noted that the Bohan and Peter diagnostic criteria for DM are useful for differentiating the pathogenesis of these myopathies. (1, 2) These criteria include symmetric and progressive loss of proximal muscle strength of the waists with or without dysphagia or compromised respiratory muscles, elevation of muscular enzymes, muscle biopsy; electromyographic alterations and cutaneous exanthema.

DM most frequently occurs secondary to neoplasms located in the lung and digestive tract but is infrequently associated with ulcerative colitis (UC). This is the first case reported in our hospital in 55 years.

CLINICAL CASE

The patient was a white, 70 year old housewife with no family or epidemiological history of interest. She reported that at during her 50s she had had bouts of diarrhea accompanied by abdominal pain, mucus and pus with a frequency of 6 to 8 bowel movements daily and a duration of 5 to 7 days. Her condition had improved with diet. She was seen
by doctors several times, but they were unable to make a diagnosis. This condition alternated with periods of clinical improvement for approximately 5 years. For approximately one year she had been suffering from weakness of her shoulders which made it difficult to raise her arms, comb her hair and perform daily household chores. In addition, she had noticed difficulty in getting up from a sitting position and could not climb stairs or swallow solid foods. All of this was associated with skin lesions around the eyes, and on her chest and both hands.

**CLINICAL EXAMINATION**

Wet and colored mucous membranes and heliotrope rashes were found in both periorbital areas (Figure 1).

![Figure 1. Heliotrope erythema, violet lesions distributed bilaterally and symmetrically around the eyes and eyelids, accompanied by a certain degree of edema.](image)

Erythema affected several areas of the body but was most accentuated on the upper chest, arms and thighs (Figure 2).

In the physical examination, the patient presented the following characteristics:

- **Cardiorespiratory system:** normal vesicular murmur, rhythmic heart tones, blood pressure: 120/80 mm Hg, heart rate: 80 beats per minute (bpm).
- **Abdomen:** pain on deep palpation of left hemiabdomen but no visceromegaly.
- **Nervous system:** Patient was well oriented in time, space and person. No focal signs
- **Musculoskeletal system:** Patient suffered generalized joint pain especially in both hands and had Gottron’s sign of red-purple keratotic, atrophic erythema on the extensor surface of the backs of her hands and metacarpophalangeal and interphalangeal areas (Figure 3). This was accompanied by progressive myalgia which was most severe in the proximal regions of both arms. Muscular hypotrophy of several muscles was striking, osteotendinous reflexes and muscle strength were diminished in all four limbs.
  - Defecation was painful and produced stools containing mucous and blood.
  - Her vagina, the fundus of her uterus, and her breasts were without alterations.

In the analytical study the patient presented the characteristics described below:

- Hemoglobin: 11 g/L, leukocytes: 10.3 x 10⁹/L with normal differential, erythrocyte sedimentation rate: 70 mm/first hour. Blood sugar, creatinine, lipids, uric acid, Addis count and venereal disease research laboratory test (VDRL) are normals. Total proteins: albumin: 64.5%, α₁: 1.3%, α₂: 2.5%; β: 4.4%; G protein: 12.6%. LE cells, IgM rheumatoid factor, antinuclear antibodies, cryoglobulins, complement and antiphospholipid antibodies, negative.
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for bacteria and parasites. Human immunodeficiency virus, hepatitis B antigens, antibodies against hepatitis C, slow serology in search of brucella and blood cultures, negative.


- Skin and muscle biopsies (Figures 4 and 5): Perivascular infiltrate of lymphocytes in the deep plexus, predominantly lymphocytic inflammation of perimysial vessels, perifascicular atrophy.

A colonoscopy was done because of the possibility that colon cancer was the cause of DM. It showed erythematous, friable mucosa on a granular surface with a normal vascular pattern. A sample was taken for histopathological study which found dispersed diffuse intraepithelial lymphocytes, apoptotic bodies in superficial epithelial cells, Paneth cell

Figure 3. Gottron’s sign.

Figure 4. Panoramic histological skin section. Note the perivascular inflammatory infiltrate of the papillary dermis and deep dermis (H/E 20x).

Figure 5. Histological field of the deltoid muscle showing predominantly lymphocytic inflammation of the endomysium and perimysium with necrosis and fiber atrophy (H/E 40x).
metaplasia, and large numbers of mononuclear cells in the lamina propria. The immunohistochemical study found that more than 25% of the cells were positive for CD8. All these findings are compatible with UC (Figure 6A and B).

Because it was difficult to reach the left colon to take a biopsy sample, an opaque enema was performed (Figure 7). The findings coincided with those reported by the colonoscopy, ruling out the possibility of a neoplasm.

The patient was treated with 1 mg/kg/day of prednisone together with 500 mg of azulfidine every 6 hours for the first weeks followed by tablets every 12 hours after discharge. This treatment was effective, and the patient’s clinical condition improved significantly.

**DISCUSSION**

The diagnosis of an inflammatory myopathy should be considered when patients present weakness of the upper and lower extremities without sensory symptoms and when rashes characteristic of DM are present. (1-4) Heliotrope, violet, periorbital rashes with macular edema rashes are pathognomonic. On other occasions, Gottron’s sign appears. This sign consists of violet papules on the backs of the joints of both hands, as observed in this patient. Similarly, the shawl sign may occur. It consists of periungual telangiectasias, capillary thrombosis, poikiloderma, and areas of photosensitivity in exposure zones.

Other symptoms and signs that may occur include interstitial pneumonitis, cardiac involvement and dysphagia, as observed in this patient. Enzymatic alterations characteristic of DM pose differential diagnoses with malignant diseases, hepatopathies and myocardial infarction. DM in association with malignant processes constitutes a paraneoplastic syndrome. Some authors indicate that its incidence is between 13% and 15% while others place it between 20% and 30%. (5, 6) The data offered by skin and muscle biopsies show inflammation of the endomysium and perimysium. Perifascicular atrophy can occur, indicating the diagnosis.

UC occurs in most cases between the third and seventh decade of life. (7) This patient was 70 years old. It should be remembered that the cause and the mechanism by which tissue damage occurs is unknown. (8) This patient was diagnosed using the Mendeloff criteria of a history of diarrhea or rectal bleeding for 6 weeks or more, sigmoidoscopy revealing friability of the mucosa with contact bleeding or petechial hemorrhaging with inflammation of the mucosa, barium enema showing evidence of ulceration, narrowing of the colon, and characteristic macroscopic or microscopic alterations. (9) All of these alterations were present in this case.

The appearance of extracolonic complications is well-known with an incidence between 25% and 36%. (10) The most frequently affected organs are the joints, skin, eyes and hepatobiliary system. (7, 10, 11) As can be seen, involvement of DM with UC occurs very infrequently. No cases were found in our search of the medical literature. Musculoskeletal symptoms are the most frequent symptoms of this disease. (10) They include arthritis similar to

![Figure 6. A and B. Histological micrograph of the colon biopsy showing severe chronic inflammation of the entire mucosa in which cryptitis typical of the disease is seen (H/E 40x).](image-url)
CONCLUSIONS

This is a case report of a patient with a previous history of gastrointestinal symptoms who came to our hospital with joint, muscular and dermatological manifestations. Physical examination and complementary examinations confirmed a diagnosis of DM associated with UC although the possibility of colon neoplasia had been suspected initially. This diagnosis must always be considered for timely treatment.

REFERENCES


rheumatoid arthritis, and ankylosing spondylitis and acute toxic arthritis have both been described. (11) Only generalized arthralgia was reported by this patient.

It is estimated that the risk of colorectal cancer in UC is 4 to 20 times higher than in the general population. (7) Taking this possibility into account, all studies necessary to rule out this disease were performed, and UC was diagnosed by endoscopic, imaging and histological studies.

Steroids are the treatment of choice for DM. (12) Prednisone at a dosage of 1 mg/kg/day after an initial dose of 60 mg is considered to be the most advisable. After 6 to 8 weeks of treatment the dose can be slowly reduced. Treatment should last from 12 to 18 months. For the treatment of UC, first generation aminosalicylates were used. The patient received 500 mg of azepidine in maintenance doses every 6 hours. Following discharge, she continued to take two tablets daily. With this treatment, a notable improvement was achieved both in clinical terms (serum enzyme levels) and in terms of systemic manifestations.

Figure 7. Barium enema of colon. Note the loss of haustra and stiffening of the wall of the descending and sigmoid colon.