CASE REPORT

Systemic lupus erythematosus complicated by intestinal vasculitis and pneumatosis intestinalis
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ABSTRACT
Gastrointestinal manifestations in systemic lupus erythematosus (SLE) are not uncommon. Non specific symptoms are often observed, such as abdominal pain, nausea, vomiting and diarrhea. On the other hand, pneumatosis cystoides intestinalis, which is characterized by multiple gas-filled cysts located throughout the intestinal wall, is a rare condition in SLE. We describe a case of a 20-year-old man who was admitted with fever, weight loss, headache and arthralgia and had a diagnosis of systemic lupus erythematosus. During his hospital stay, he developed abdominal symptoms that suggested intestinal vasculitis. The computed tomography of the abdomen showed the double halo sign, or target sign and pneumatosis cystoides intestinalis. The patient presented complete recovery after conservative treatment, with intestinal rest and total parenteral nutrition.

Keywords: systemic lupus erythematosus, vasculitis, pneumatosis cystoides intestinalis, gastrointestinal tract.

INTRODUCTION
Pneumatosis intestinalis (PI) is an unusual condition, characterized by multiple cysts filled with gas in the intestinal mucosa, submucosa or subserosa. The first description dates from 1730, but Meyer was the first to use the term.¹ There are several associated conditions and among the conjunctive tissue diseases, the most common is systemic sclerosis.¹ It rarely occurs in systemic lupus erythematosus (SLE), with only 14 cases described to date in the literature. We describe herein a rare case of PI in a 20-year-old man with a recent diagnosis of SLE.

CASE REPORT
A previously healthy twenty-year-old male patient, presented fever for 10 days, accompanied by weight loss, severe headache and history of two painful oral ulcers.

At admission, he presented regular general status, dehydration and looked pale. He was conscious, oriented and febrile. Cervical lymphadenomegaly was observed, without oral ulcers. Chest examination was normal. The abdomen was flat, flaccid and hydro-aerial noises were present. The patient presented pain in the right hypochondriac region; the liver was palpated 4 cm below the right costal border and the spleen could not be palpated. The patient presented knee and elbow arthritis. The neurological assessment was normal.

Laboratory assessment showed: anemia (Hemoglobin: 11.0 g/dL); lymphopenia (331/mm³); thrombocytopenia (63,000/mm³); Coombs reagent; reticulocytes: 0.3%; normal liver function tests; ESR 44 mm/h; creatinine 1.4 mg/dL; urea 64 mg/dL; partial urinalysis with protein +++, 10,000 leukocytes, 40,000 erythrocytes, > 10 hyaline cylinders; negative blood cultures.

Due to the presence of fever and headache, a cerebrospinal fluid sample was collected, which showed the presence of aseptic meningitis – 11.6 leukocytes; 61% monomorphonuclear and 29% polymorphonuclear; protein 72.9 mg/dL; glucose 43 mg/dL; VDRL, gram, culture,
Ziehl, serological test for herpes and direct mycological examination were negative.

The serological tests for hepatitis B and C, HIV and toxoplasmosis were negative; for CMV and EBV, IgG was positive and IgM was negative, respectively. The Antinuclear Antibody (ANA) test was positive, with a titer higher than 1:640, homogenous nuclear pattern; C3 and C4 were decreased (52 and 8 mg/dL, respectively); ENA profile was non-reagent and Anti-DNA was > 1:640.

The echocardiogram showed an ejection fraction of 74% (> 58%), small pericardial effusion, with no vegetation. Systemic lupus erythematosus (SLE) was diagnosed (arthritis, serositis, hematological alterations, nephritis, anti-DNA and ANA). The patient started to present mental confusion, generalized tonic-clonic seizures, abdominal pain and nausea. A skull computed tomography (CT) with contrast showed diffuse hypodensities in the brain hemispheres, which could not rule out vasculitis. Pulse therapy with 1g/day of methylprednisolone IV was started due to the severe Central Nervous System involvement, followed by 1g of cyclophosphamide IV on the fourth day, which was immediately followed by complete neurological recovery and worsening of the abdominal picture, with gas and stool retention. The abdomen was slightly distended and hypertympanic; hydro-aerial noise was absent and the patient presented diffuse pain on palpation, with rebound tenderness (Blumberg sign).

The acute abdomen X-ray showed hydro-aerial levels and small bowel dilatation, with “stack-of-coins” appearance (Figure 1).

The ultrasonography (US) showed enlarged kidneys and loss of cortico-medullary differentiation; moderate amount of free fluid in the peritoneal cavity, edema and thickening of the small bowel loop, with increased echogenicity of the mesenteric fat around it. The abdominal CT showed, in addition to the US findings, liquid distension and thickening of the small bowel loop wall, associated to PI (Figure 2).

The probable diagnosis of vasculitis of the small vessels of the intestinal wall was attained through the CT, which justified the diffuse enteric involvement, loop edema and PI.

The surgical approach was chosen together with the conservative treatment – antibiotic therapy by IV route, bowel rest and total parenteral nutrition (TPN) – while waiting for the response to the pulse therapy. As there was free fluid in the cavity, a paracentesis was performed and bacterial peritonitis was ruled out.

There was improvement in abdominal pain. After 8 days, the patient started to have bowel movements. Four days later,
The clinical picture is variable. It can manifest worsening of the pre-existing leukopenia with intestinal ischemia) and escape of contrast medium in intestinal microthrombosis, as unspecific abdominal discomfort, diarrhea, abdominal distension, to acute abdomen with massive bleeding. The absence of hydro-aerial noise is also sensitive to detect collections of intramural gas, such as PI.\textsuperscript{4,8,9,10,11,12,13}

Arteriography is not useful, as the disease usually affect small vessels. Endoscopy and colonoscopy are useful to locate ulcers and perform biopsies.\textsuperscript{5}

The differential diagnosis is a challenge due to the multiple causes of abdominal pain in the patient with SLE.\textsuperscript{14}

When there is no evidence of intestinal perforation, ischemia is potentially reversible and a conservative treatment is indicated. That includes high-dose corticoids (prednisone 1-2 mg/kg/day) or pulse therapy with methylprednisolone 1 g/day, for 3 days. Intestinal rest must be associated (TPN). Prokinetic agents can improve the peristaltism and reduce intraluminal pressure. Intravenous antibiotic therapy is used as adjuvant in the presence of IP; it aims at reducing bacterial overgrowth and gas production by anaerobic bacteria.\textsuperscript{3,4,5,15}

Cyclophosphamide pulse therapy, 0.75-1 g/m\textsuperscript{2}, is indicated in refractory cases.\textsuperscript{2,4,13}

There have been reports on the use of inhaled oxygen or hyperbaric chamber in an attempt to remove the gas from the cysts and of octreotide to improve motility and reduce bacterial overgrowth.\textsuperscript{10}

Prognosis is poorer when there is surgical indication, which can be improved when the procedure is performed early.\textsuperscript{5,5,7} Early clinical suspicion and treatment are essential for a good evolution. Factors associated with recurrence are thickening of the intestinal wall > 9 cm at the tomography and lower cumulative dose of immunosuppressive agents.\textsuperscript{15}

Although gastrointestinal tract vasculitis and IP are rare manifestations of SLE, the present case report describes a possible association between the two conditions.

**DISCUSSION**

The incidence of vasculitis of the gastrointestinal tract varies between 0.2% and 53% in lupus, with a close correlation with disease activity. There have been 14 cases of PI associated with SLE described in the literature. Underlying vasculitis was demonstrated in 50% of them.\textsuperscript{2,3,4,5,6}

The vasculitis usually affects small vessels of the intestinal wall. The histological assessment demonstrates atrophy and degeneration of the media, fibrinoid necrosis, thrombosis and lymphocytic infiltrate in lamina propria; the immunohistochemical analysis shows deposit of immune complexes, complement and fibrinogen.\textsuperscript{4,7} PI is supposedly caused by an increase in the intraluminal pressure, mucosal injury and gas production by bacteria in the mucosa.\textsuperscript{1,6,8} The clinical picture is variable. It can manifest as unspecific abdominal discomfort, diarrhea, abdominal distension, to acute abdomen with massive bleeding. The use of high-dose immunosuppressive drugs and corticoids can mask peritonitis. The absence of hydro-aerial noise is essential for a good evolution. Factors associated with recurrence are thickening of the intestinal wall > 9 cm at the tomography and lower cumulative dose of immunosuppressive agents.\textsuperscript{15}

An oral diet was re-introduced and was well tolerated. A new abdominal CT performed 10 days later demonstrated significant improvement in the intestinal loop distension and wall edema, with disappearance of the PI.

The patient was discharged with oral prednisone (1 mg/kg/d), in addition to planned monthly pulses of cyclophosphamide IV. During the follow-up, he did not present new exacerbations of the intestinal picture.

**REFERENCE**

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