Giant Intramural Hematoma of the Colon in Acquired Factor VIII Inhibitor

Hematoma Intramural Gigante do Colon em Doente com Inibidor do Factor VIII Adquirido

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INTRODUCTION

Acquired hemophilia is a rare bleeding disorder caused by the development of autoantibodies against plasma coagulation factors, most frequently factor VIII.¹ Normal hemostasis requires at least 25% of factor VIII activity, which is decreased in patients with acquired inhibitors against factor VIII.² Standard coagulation studies can be normal, including thrombin and prothrombin times (PT) as well as platelet count and function. The typical findings of acquired hemophilia are a prolonged activated partial thromboplastin time (aPTT) and a low factor VIII level, with the latter not being routinely requested.¹ Treatment priorities are to stop the bleeding and to eradicate factor VIII autoantibodies. Immunosuppressive treatment with corticosteroids and cytotoxic drugs alone or in combination is regarded as the appropriate treatment.³

Intramural colonic hematomas are rather uncommon⁴ and are mostly characterized by abdominal pain, rectal bleeding and bowel obstruction depending on the size and location of the hematoma. The most frequent causes of intramural bowel hematoma are abdominal trauma, anticoagulant therapy, blood dyscrasias or iatrogenesis.⁵ The aim of this paper is to highlight the occurrence of acute abdominal obstruction due to spontaneous intramural colonic hematoma resulting from the unusual cause of acquired hemophilia.

CASE REPORT

A 66-year-old man, diagnosed four years before with Crohn disease affecting the right colon, presented to the emergency department complaining in the last day of 14 episodes of diarrheal stools with hematochezia and abdominal pain in the left lower quadrant, without previous story of colorectal trauma, as well as signs of colonic infection. Blood analysis revealed a normochromic normocytic anemia [haemoglobin 8.9 g/dL (13.5 – 17.5)] normal platelet and white blood cell counts [176 × 10⁹/L, 10.1 × 10⁹/L] and a prolonged aPTT [49.1 s (25.1 – 36.5 s)] with normal PT and international normalized ratio (INR). Colonoscopy showed a clot 50 cm from the anal verge which occluded the lumen. The patient received transfusion of three units of packed red blood cells (PRBCs), one gram of fibrinogen and three units of plasma in the emergency room. After being admitted to the Gastroenterology unit, he presented clinical worsening,

RESUMO

O hematoma intramural do colon é uma entidade rara, principalmente quando associada ao desenvolvimento de anticorpos anti fator VIII. Apresentamos um homem, 66 anos, com dor abdominal, hematoquezias e presença de coágulos no cólon esquerdo, radiologicamente sem sinais de hemorragia ativa ou oclusão intestinal e sem alterações analíticas nas provas de coagulação, mas com presença de anticorpos contra o fator VIII. Por agravamento e instabilidade clínica foi submetido a laparotomia exploradora, tendo-se verificado a presença de hematoma intramural do colon esquerdo, pelo que se procedeu a hemicolecetria esquerda. Foi medicado com hemoderivados e corticosteroides com evolução clínica favorável. O diagnóstico do hematoma intramural espontâneo pode ser um desafio, principalmente na ausência de suspeita clínica. O seu reconhecimento precoce é essencial para uma evolução favorável. Este caso releva uma causa rara de hemorragia e oclusão intestinal, bem como a dificuldade e importância do diagnóstico clínico na ausência de exames complementares elucidativos.

Palavras-chave: Colon; Hematoma; Hemofilia A; Factor VIII

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ABSTRACT

Intramural hematoma of the colon is very rare, particularly when associated with the development of autoantibodies against factor VIII. We report a case of a 66-year-old man with abdominal pain, hematochezia and clots in the left colon, without any radiologic signs of active bleeding or bowel occlusion or analytical changes in routine coagulation screening, but with positive autoantibodies against factor VIII. The clinical instability prompted surgical exploration. An intramural hematoma of the left colon was found, and a left colectomy was performed. The patient was treated with hemoderivatives and corticosteroids with clinical improvement. The diagnosis of spontaneous intramural hematoma might be a challenge, particularly in the absence of clinical suspicion. An early recognition is essential for a positive outcome. This case highlights a rare cause of bleeding and intestinal obstruction, but also the difficulty and relevance of establishing a clinical diagnosis when diagnostic tests are not completely informative.

Keywords: Colon; Hematoma; Hemophilia A; Factor VIII

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with abdominal distension, generalized pain and persistent bleeding. Abdominal computerized tomography (CT) did not show signs of active haemorrhage or wall thickening. Reordered coagulation analysis indicated prolonged PT [14 seconds (9.4 – 12.5 s)], and aPTT [48.1 seconds (25.1 – 36.5 s)], as well as decreased activity of factor VIII [30.70% (50% – 150%)], increased factor VIII inhibitor titer [2.01 U Bethesda (negative < 0.5)], and increased functional Von Willebrand Factor [284.6% (48.8% – 163.4%)]. Since gastrointestinal bleeding persisted, the haemoglobin level continued to fall and the patient became unstable, and was then transfused with one PRBCs and three plasma units and admitted to the intensive care unit.

Colonoscopy was repeated, revealing at the same level of the previous clot a violaceous mucosa with necrosis and segments of destruction over around 20 cm. Surgical exploration was decided. After administration of 3000 U of factor VIII and four PRBCs, exploratory laparotomy was performed and found the left colon filled with an intramural hematoma from the sigmoid to the splenic flexure, causing distension of the proximal colon (Figs. 1–3). A left colectomy was done with mechanical closure of the distal stump and terminal colostomy. Histopathology confirmed an extended submucosal hematoma with necrosis of mucosa and without signs of Crohn disease (Fig. 4). On the first postoperative day, the patient received 3000U of factor VIII and started prednisolone (120 mg/day). He also received three more PRBCs on the third postoperative day and 238 000 U of anti-inhibitor coagulant complex during the 17 days following the postoperative day. The eradication of factor VIII inhibitors three weeks after surgery, allowed the start of corticosteroid tapering. A deep incisional surgical site infection was detected and treated with piperacillin-tazobactam and dressings. The patient was discharged on the 26th day after surgery and referred to the immunohemotherapy outpatient clinic. Corticosteroid tapering was completed four months after surgery, with normal activity of factor VIII and negative factor VIII inhibitor titre. Five months later, the patient underwent endoscopic evaluation, with intestinal mucosa without changes, and intestinal transit was reestablished.
with mechanical colonic anastomosis. The postoperative period was uneventful, and the patient was discharged on the fifth day.

**DISCUSSION**

Intramural hematoma of the colon is a rare event that can be secondary or ‘spontaneous’. The hemorrhage is usually located in the submucosal layer of the bowel due to the presence of more vascular structures. Haemorrhagic ascites can be present and is usually related to leakage of blood from an engorged, thickened and inflamed bowel wall with submucosal bleeding extending into all layers. It is believed that the progression of the symptoms is due to the establishment of an intramural osmotic gradient and the presence of the hematoma, leading to an expansion of the intestinal wall.

The diagnosis of intramural hematoma of the colon is a clinical challenge, because signs and symptoms are not specific and clinical suspicion is crucial. The clinical presentation depends on the location of the hematoma, and symptoms of either high or low bowel obstruction may predominate. Abdominal pain is present in almost all cases, being either diffuse or predominantly located in the site of the hematoma. Signs of peritoneal irritation are suggestive of complications such as necrosis, perforation or hemoperitoneum. Digestive bleeding occurs in about 40% of the cases. Abdominal CT may be important to the diagnosis, with suggestive images of circumferential wall thickening, intramural hyperdensity, luminal narrowing, intestinal obstruction, and hyperdense ascites. However there are no pathognomonic CT changes and plain abdominal x-ray films reveal only typical patterns of colic obstruction if present. Colonoscopy may be useful to show hematoma signs like “blue and roundish formations” in the submucosal layer.

In this case, the analytical results were key for the diagnosis of acquired haemophilia and the diagnosis of intramural hematoma was neither evident in the CT scan nor in the colonoscopy. Although rare, massive gastrointestinal tract bleeding can occur as a complication of Crohn’s disease and can be particularly difficult to control. It is also known that bleeding is more frequent among patients with colonic involvement, especially of the descending colon and rectum. Since the patient had a previous diagnosis of Crohn’s disease, we strongly considered it as a cause for the haemorrhage, even though the latest colonoscopy of the patient, one year before, showed just three superficial ulcers in the right colon. Nevertheless, diagnostic procedures that are used in patients with lower gastrointestinal bleeding are similar in patients with or without inflammatory bowel disease.

The decision for surgical intervention was determined by the persistent haemorrhage, signs of bowel obstruction and the clinical instability of the patient. The appropriate pharmacological treatment of patients with acquired haemophilia depends essentially on the natural history of any concomitant pathology and the clinical presentation of coagulopathy. A conservative treatment with corticosteroids is usually recommended for acquired inhibitor haemophilia in the first instance.

The differential diagnosis of a haemorrhagic disorder with normal baseline coagulation studies is limited to clinical suspicion, and a failure to recognize an uncommon acquired coagulation disorder can have serious consequences. For the treatment of colonic haemorrhage, surgical treatment is suggested for those patients with a doubtful diagnosis or those who exhibit a deteriorating condition, an irreversible intestinal obstruction, signs of bowel necrosis or peritonitis.

The surgical treatment with resection of the intestinal segment with hemostasis is the gold standard therapy. Very few cases have been treated by endoscopic evacuation with a good outcome, and only when mucosal perforation was absent.

A prompt and early recognition of nontraumatic, spontaneous, large-bowel intramural hematoma, and early involvement of surgical care is crucial for a positive outcome. Rare coagulation disorders, like acquired inhibitor haemophilia presented in this case, are important to consider in the differential diagnosis for a colonic haemorrhage or obstruction of unknown cause.

**CONCLUSION**

The present case reports an atypical cause of low gastrointestinal haemorrhage, associated with the presence of coagulopathy due to an acquired inhibitor of factor VIII. The diagnosis of intramural hematoma implies a high level of clinical suspicion due to its non-specific clinical presentation. The delay in diagnosis leads to a delay in effective treatment, with poor outcome and increased associated costs.

**CONFLICT OF INTEREST**

The authors certify that they have NO affiliations with or involvement in any organization or entity with any financial interest (such as honoraria; educational grants; participation in speakers’ bureaus; membership, employment, consultancies, stock ownership, or other equity interest; and expert testimony or patent-licensing arrangements), or non-financial interest (such as personal or professional relationships, affiliations, knowledge or beliefs) in the subject matter or materials discussed in this manuscript.

**PROTECTION OF HUMANS AND ANIMALS**

The authors declare that the procedures were followed according to the regulations established by the Clinical Research and Ethics Committee and to the Helsinki Declaration of the World Medical Association.

**DATA CONFIDENTIALITY**

The authors declare having followed the protocols in use at their working center regarding patients’ data publication.
Um Caso de Histiocitose de Células de Langerhans Orbitário num Adulto

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ABSTRACT

Langerhans cell histiocytosis remains an enigmatic disease with a very heterogeneous presentation. We describe a rare case of orbital Langerhans cell histiocytosis in a 39-year-old female patient who presented right orbital pain and edema of the upper right eyelid. Surgery showed a friable lesion and underlying bone irregularity. Morphological aspects and immunohistochemical profile favored the diagnosis of Langerhans cell histiocytosis, which was confirmed with evidence of Langerin expression. The staging tests did not reveal any organ involvement, so we decided to follow the algorithm proposed by Euro Histio Net: in case of unifocal disease and in a single organ, clinical surveillance was preferred. This case aims to raise awareness of a manifestation of Langerhans cell histiocytosis, which should always be considered as a differential diagnosis in adults with osteolytic orbital lesions.

Keywords: Adult; Histiocytosis, Langerhans-Cell; Orbital Diseases

RESUMO

A histiocitose de células de Langerhans permanece uma doença enigmática com apresentação muito heterogênea. Descrevemos um caso raro de histiocitose de células de Langerhans orbitária numa doente do sexo feminino, 39 anos, com dor orbitária e edema da pálebra superior direita. A tomografia computorizada das órbitas revelou uma lesão lítica próxima da glândula lacrimal. Na cirurgia observou-se uma lesão frátil e irregularidade óssea subjacente. Os aspectos morfológicos e perfil imunohistoquímico favoreciam o diagnóstico de histiocitose de células de Langerhans, confirmando-se com a evidência da expressão da Langerina. Uma vez que os exames de estadiamento não revelaram envolvimento de outro órgão, decidimos seguir o algoritmo proposto pelo Euro Histio Net: tratando-se de doença unifocal e uni-órgão, optamos pela vigilância. Este relato de caso visa alertar para uma manifestação rara da histiocitose de células de Langerhans, a qual deve ser sempre considerada como um diagnóstico diferencial em adultos com lesões orbitárias osteolíticas.

Palavras-chave: Adulto; Doenças da Órbitas; Histiocitose de Células de Langerhans

INTRODUÇÃO

Langerhans cell histiocytosis (LCH) is characterized by a proliferative lesion of pathological cells similar to Langerhans cells. Multiple clinical series have shown that the peak incidence of LCH occurs between the ages of 1 and 3 years, with most cases of multiple-organ system disease beginning before the age of 2 years old.1 Although this uncommon disease typically affects childhood, LCH does occur in adults, with a considerable paucity of clinical data.2 Positivity of cluster of differentiation (CD) 1a and Langerin in the lesion is the current gold standard for the diagnosis of LCH, but only in the correct clinical setting.3 The clinical