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Intestinal pseudopolyps in a patient with Crohn's disease and renal transplant – An unexpected diagnosis



Pseudopólipos intestinales en un paciente trasplantado renal y con enfermedad de Crohn – un diagnóstico inesperado

reaction analysis of the biopsy specimens was performed and confirmed the presence of Leishmania infantum. HIV otes structures of Leishmania sppl (Fig. 2). Polymerase chain stool cultures were negative. The ileocolonoscopy showed infiltrate and macrophages with spherical and small amastigof the mucosa, with an inflammatory lymphoplasmocytic exam revealed preservation of the glandular architecture right colon and in the terminal ileum, and the pathological the right colon (Fig. 1). Biopsies were performed in the scarring areas and pseudopolyps of the terminal ileum and hepatosplenomegaly. Lab workup showed pancytopenia and months after transplant, with clinical improvement. years and he was immunosuppressed with tacrolimus and thy. He referred several trips to the Middle East in recent Crohn's disease (CD), stricturing phenotype, diagnosed in A 41-year-old male, journalist, with a history of ileal bowel movements. Physical examination was relevant for years later, the patient reported a slight increase of his immunosuppressive drugs, prednisolone. Due to CD activity and malabsorption of and renal transplant in 2014 due to IgA nephropainfliximab was initiated a . Wo few

screening test was negative. The patient started therapy with liposomal amphotericin B.

Leishmaniasis is a chronic protozoan disease of the mononuclear phagocytic system.¹ Leishmania spp is endemic in several regions of the world, including the Mediterranean área.¹ The incubation period is usually long, and under conditions of immunosuppression, there is evidence of activation of latent infection several years after exposure to the parasite.¹ TNF- α has a major role in mediating host protection against visceral leishmaniasis (VL), so the use of anti-TNF agents may potentially cause worsening or reactivation of latent infection.¹,²

Cutaneous leishmaniasis is the most common leishmanial syndrome worldwide.³ VL, which reflects dissemination of Leishmania parasites throughout the reticuloendothelial system, is potentially life threatening without treatment.² VL is a systemic disease characterized by hepatosplenomegaly, fever, cachexia, hypergammaglobulinaemia, and pancytopenia.¹ Nevertheless, asymptomatic leishmanial infection has been reported previously.¹

The diagnosis of enteric VL is histological, requiring visualization of amastigotes inside macrophages of the intestinal lamina própria.^{4,5} A characteristic endoscopic image of this invasion has not been described so far, and the diagnosis is established by taking biopsies.^{4,5} A correct diagnosis of VL is challenging and easy to miss, especially in cases that are not clinically suspected, as the case reported. Liposomal amphotericin B is the preferred treatment choice.⁵

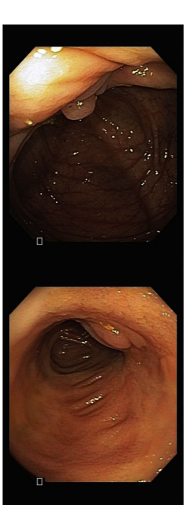


Figure 1 lleocolonoscopy imaging showing scars and pseudopolyps of the terminal ileum and the ileocecal valve

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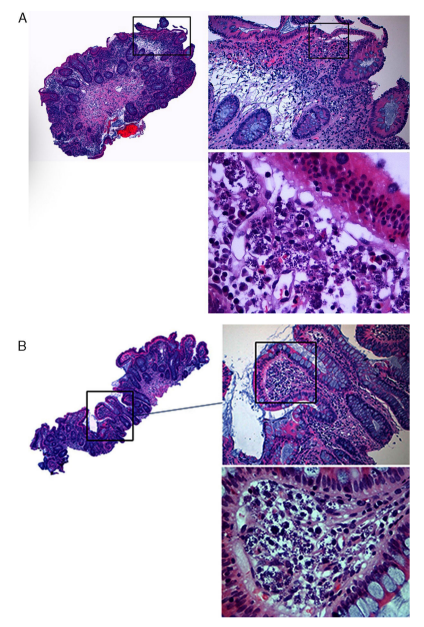


Figure 2 Histological showing small amastigotes structures of *Leishmania spp*, in the right colon mucosa (A) and in the terminal ileum (B).

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