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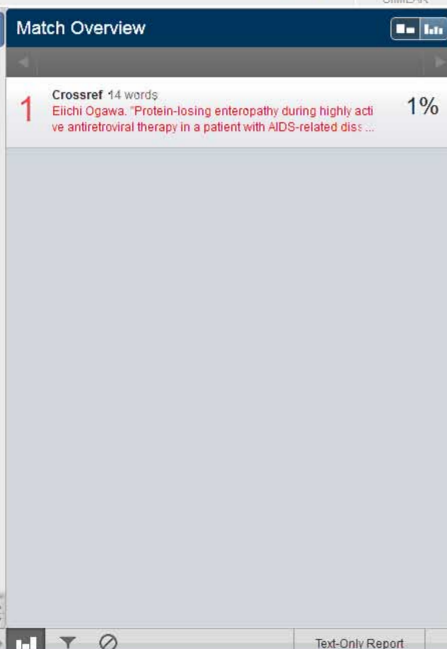
Manuscript Type: CASE REPORT

Protein-losing pseudomembranous colitis with cap polyposis-like features

Wolfgang Kreisel, Guenther Ruf, Richard Salm, Adhara Lazaro, Bertram Bengsch, Anna-Maria Globig, Paul Fisch, Silke Lassmann, Annette Schmitt-Graeff

Abstract

Protein-losing enteropathy (PLE) is characterized by loss of serum proteins into the gastrointestinal tract. It may lead to hypoproteinemia and clinically present as protein deficiency edema, ascites, pleural or pericardial effusion and/or malnutrition. In most cases the site of protein loss is the small intestine. Here we present an unusual case of severe PLE in a 55-year old female with a one-year history



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A subsequent colonoscopy revealed polyps similar to those seen during the patient's ... with foci of cryptitis, which was suggestive of inactive ulcerative colitis. ... reports of lower limb edema as a result of protein-losing enteropathy from cap polyposis. ... Similar histologic features are present in conditions such as prolapsing ...

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Inflammatory cap polyposis in a 42-year-old male - NCBI - NIH

https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3604675/

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Jean-Claude Givel, Neil Mortensen, Bruno Roche - 2009 - Medical

There are striking clinical features of alopecia, white nails and protein-losing enteropathy ... Inflammatory cap polyposis is a rare manifestation of mucosal prolapse [33]. ... It is a close mimic of pseudomembranous colitis and of other polyposis. ... Occult, diffuse "linitis plastica" like adenocarcinoma is another not infrequent 95 ...





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缺少字词: pseudomembranous

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