Ulcerative Colitis and Acute Thrombocytopenia in a Pediatric Patient: A Case Report and Review of the Literature

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Abstract

We report the case of a 14-year-old boy affected by ulcerative colitis (UC) and acute thrombocytopenic purpura (ITP) with simultaneous onset. UC diagnosis was based on symptoms, endoscopy and histology findings. ITP diagnosis was based on the normal bone marrow megakaryocyte count, the presence of platelet associated IgG and the absence of splenomegaly. Medical treatments including high doses of steroids, intravenous immunoglobulins were ineffective on ITP course, while UC course was mild for several months after the onset. When colonic inflammation became untractable not responsive at steroids and immunosuppressive agents, colectomy resolved both pathologies.

Keywords

Ulcerative Colitis, Thrombocytopenia, Children

1. Introduction

Ulcerative colitis (UC) is a chronic inflammatory bowel disease (IBD) frequently associated with extra-intestinal manifestations and various autoimmune disorders [1] [2]. Autoimmune antibody-mediated hematologic diseases such as autoimmune hemolytic anemia have been reported in patients with IBD, both in UC and in Crohn’s disease (CD) [3]-[6]. Immune thrombocytopenic purpura (ITP), defined as low platelet count in the absence of

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other clinically apparent causes of thrombocytopenia and caused by antibody-mediated platelet destruction, has been sporadically reported in adult patients affected by IBD [7]. Association between ITP and IBD in children is rarely described.

Onset clinical manifestation of both diseases could be presented by gastrointestinal bleeding.

UC treatment with steroids and immunosuppressive drugs is usually effective to treat ITP. Medical therapy of ITP includes besides corticosteroids and immunosuppressive agents such as cyclosporine and azathioprine, intravenous immunoglobulins. Recently infliximab has also been reported as a successful treatment for immune mediated extraintestinal manifestations of UC refractory to other interventions [8] [9].

We report the case of a boy with UC and refractory ITP, in whom colectomy resolved both diseases.

2. Case Report

A 14-year-old boy, without evidence of problems in the past medical history, was admitted to “Bambino Gesù” Children Hospital of Rome with a 15 days history of bloody diarrhea and abdominal pain. No bleeding from other organs was present. There was no evidence of purpura and mucosal petechiae. Family history was unremarkable.

Initial laboratory evaluation showed: hemoglobin 12 g/dl (normal values 13 - 16 g/dl), hematocrit 33.6% (36% - 49%), white blood cell count 4350/ml (4000 - 13,500/ml), platelet count 38 × 10³/mm³ (150 - 450 × 10³/mm³), moderate elevation of CRP and ESR levels. The assessment of the coagulation resulted into the normal ranges. ASCA-IgA and IgG, ANCA, antinuclear and anti-dsDNA antibodies resulted absent. Platelet-associated antibodies were present (antibodies anti GPIIb/IIIa, antibodies anti GPIa/IIa and antibodies anti GPIb/IX). Heterozygosis of MTHFR (C677T) was present. Stool evaluation was negative for infectious pathogens, virus, Clostridium Difficile and its toxin. Fecal calprotectin levels raised up to 3550 microg/gr (NV < 250 microg/gr).

Bone marrow biopsy revealed a normocellular marrow with a normal number of megakaryocytes. Hepatotropic viruses (B and C hepatitis) and cytomegalovirus, Ebstein-Barr virus, HHV-6 and parvovirus infections in blood and bone marrow were ruled out. No splenomegaly was found on abdominal ultrasonography. Upper endoscopy showed hemorrhagic gastritis and absence of Helicobacter Pylori. Colonoscopy showed macroscopic inflammation extending from cecum to rectum (pancolitis). Terminal ileum mucosa resulted normal. Histopathology confirmed the diagnosis of UC (see Figure 1). Treatment with 5-aminosalicylic acid and intravenous prednisolone at the dose of 1.5 mg/Kg/day was begun. There was a rapid improvement of gastrointestinal symptoms and an increase in platelet count, up to 63 × 10³/mm³.

During the follow-up the platelet count remained stable around 40 × 10³/mm³, but decreased to 20 × 10³/mm when reduction of steroids was attempted. No resolution of his thrombocytopenia was obtained with a course of intravenous immunoglobulins and steroid treatment was continued at the dose of 1 mg/Kg/day.

This resulted in a good control of the child’s UC with a stable Pediatric Ulcerative Colitis Activity Index (PUCAI) below 10 [10]. Endoscopy performed three months after the diagnosis showed a normal mucosa, and histopathology confirmed the remission of the inflammatory process.

Eight months later during a further attempt to reduce steroids the patient presented bloody diarrhea with more than 15 bowel movements/day and severe abdominal pain. Colonoscopy showed severe relapse of the UC. Total parenteral nutrition, antibiotics and intravenous cyclosporine as a rescue therapy was begun. Symptoms persisted, the patient needed blood transfusion, and the PUCAI score after 5 days of therapy was 80, platelets count was 29 × 10³/mm³ Surgical treatment was considered necessary as the disease was life-threatening and total colectomy was performed. Histology of the resected colon was in keeping with a diagnosis of severe, acute UC (see Figure 2 and Figure 3).

Steroids were tapered in few days and patient was discharged. Two months after colectomy, platelet count improved and reached 85 × 10³/mm³. Six months afterwards was 150 × 10³/mm³ and at 10 months platelet count was normal (240 × 10³/mm³).

The informed consent was obtained from the patient and his family to report this case.

3. Discussion

The association of IBD with ITP is rare and it has been recently recognized and described. Since the first description by Edwards and Truelove in 3 adult patients among 624 adults affected by UC, few occasional cases have been published [8] [9] [11]-[24]. Only 8 papers report of pediatric patients [12] [14] [22] [23]. We de-
scribed one more pediatric patient in whom UC and ITP had a simultaneous onset. The diagnosis of UC in our patient was based on symptoms, endoscopy and histology findings. The diagnosis of ITP was based on bone marrow examination that revealed a normal megakaryocyte number, the presence of platelet-associated IgG, and absence of splenomegaly. There was no evidence of any hematological disease, infection or immune deficiency. Disseminated intravascular coagulation (DIC) was ruled out by the presence of normal prothrombin time and normal levels of fibrinogen and fibrin degeneration products.

ITP in our patient had a partial response to high doses of steroids, with platelets number rising up to $60 \times 10^3$/mm$^3$ during full steroid treatment, while every attempt to reduce the dose was followed by a fall in the platelet count. It has been postulated that thrombocytopenia with platelet associated antibodies occurs during a severe relapse of UC. On the other hand increased colonic permeability to luminal antigens may produce antibodies which may result in cross-reactivity with platelet antigens, resulting in platelet destruction. This would be
true during an acute exacerbation because of enhanced exposure to luminal antigens and that is what happens in
the majority of cases of ITP associated with IBD [12]. On the contrary of what is reported in the literature, in our
case low number of platelets were not associated, at least during the first months of the disease, with exacerba-
tion of gastrointestinal symptoms and colonoscopy performed during his minimal platelet count (20 × 10^3/mm^3)
showed an improvement of mucosal injury. It is any way possible that, once activated the autoimmune mech-
nism, a subclinical injury of the mucosa may perpetuate the ITP.

Several treatments have been used in the management of both diseases. Courses with steroids in association
with mesalazine have been frequently able to induce remission of both diseases. Sometimes high doses of ster-
oids, immunosuppressive agents and intravenous immunoglobulin or splenectomy have been required to control
ITP [12] [13] [25]-[28].

For at least eight months our patient obtained a good control of his colitis with mesalazine and steroids while
ITP was partially controlled even during clinical and histological remission of colitis. The relapse of his colitis
was severe and a rescue medical therapy became necessary. Intravenous Cyclosporine A (CyA) was started at
the dose of 3 mg/kg/day. CyA has been effectively used for the treatment of severe steroid-refractory UC [10]
[29] [30] and it has also been reported to be useful in patients with corticosteroid and splenectomy-refractory
ITP [26]. In our patient rescue therapy with IV CyA was unsuccessful and surgery was unavoidable. Colectomy
was resolutive for both pathologies. Platelets number gradually rose after surgery—when all medical therapies
were stopped—and reached sub-optimal levels within 6 months after colectomy. The platelet count was com-
pletely normalized 10 months after colectomy and continued to be normal during the follow up of almost 18
months. In the past six adult patients affected by ITP and UC have been described to undergo colectomy, in five
of them the surgical intervention was resolutive for both ITP and UC [15]-[18] [21]. Higoushi et al. reported
four pediatric cases of UC and ITP one of which underwent colectectomy because of fulminant colitis unrespon-
sive to medical treatment. Medical therapy was stopped after surgery but she continued to have episodes of
thrombocytopenia treated with IV Ig and steroid [12]. In another pediatric patient ITP developed after colectomy
[20]. Hisada et al reported a case of acute ITP associated with preexisting ulcerative colitis in a 36-year-old man
in whom colectomy was performed at the patient’s request because of life-threatening colonic hemorrhage without
attempt a rescue therapy with intravenous cyclosporine or infliximab and resulted in cure of both diseases [19].

A recent Italian survey described the phenotype and disease course of pediatric onset UC diagnosed at a terti-
ary referral Center for Pediatric Gastroenterology. In this retrospective evaluation, 14% out of the 110 involved
underwent. The presence of extensive disease, severe colitis and need of corticosteroids at the diagnosis were
associated in this study with an increased risk of colectomy [31]. Our patient had all the risk factors that can lead
to approach the surgical option.
In conclusion we have reported for the first time the case of a pediatric patient with UC and thrombocytopenia who did not respond to the medical treatment in which colectomy has been resolutive for both diseases. We therefore consider that colectomy must be considered in children when both diseases are resistant to intensive medical management.

References


List of the Abbreviations

Ulcerative Colitis: UC
Idiopathic Thrombocytopenic Purpura: ITP
Inflammatory Bowel Diseases: IBD
Crohn’s Disease: CD
Disseminated Intravascular Coagulation: DIC
Cyclosporine A: CyA
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