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**Endoscopic evaluation of immunotherapy-induced gastrointestinal toxicity**

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**Abstract**

Immunotherapy is any treatment aimed at boosting or enhancing the immune system. It includes a wide range of options, from vaccines to treatment for conditions such as allergy and cancer. In the case of cancer, unlike other available treatments, immunotherapy is not aimed at destroying the tumor cells but at stimulating the patient’s immune system so that it attacks the tumor. In cancer, immunotherapy provides a series of advantages. Nevertheless, immunotherapy administered for treatment of cancer is associated with immune-mediated enterocolitis. Colitis mediated by monoclonal anti-cytotoxic T lymphocyte–associated antigen 4 and to programmed cell death protein 1 and its ligand PDL1 shares characteristics with chronic inflammatory bowel disease (IBD), and similar findings have been reported for both the endoscopy images and the segment involved. The most frequent lesions on endoscopy are ulcer and erythema, and the most frequently affected site is the sigmoid colon. A segmental pattern has been reported to be slightly more frequent than a continuous pattern. In addition, upper gastrointestinal lesions have been reported in up to half of patients, with the most frequent findings being gastritis and erosive duodenitis. As is the case in IBD, systemic corticosteroids and immunosuppressive treatment (anti-TNF agents) are the approaches used in patients with a more unfavorable progression. Immunotherapy must be suspended completely in some cases.

**Key words:** Enterocolitis; Ipilimumab; Immunotherapy; Immune-related adverse event; Nivolumab; Toxicity; Endoscopy

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**Core tip:** Widespread use of immunotherapy in various types of cancer has led to reports of new associated adverse effects resulting from increased stimulation of the immune system, which can confuse the body’s own tissues and organs with foreign matter, thus leading it to attack the body’s healthy tissue. The most frequent immune-mediated adverse effects include asthenia, general malaise, fever, gastrointestinal toxicity (abdominal pain, diarrhea, and colitis), cutaneous toxicity, hypothyroidism and hepatitis. This review of the endoscopic evaluation of immunotherapy-induced toxicity presents the most typical endoscopic images, the differential diagnosis based on these images, and the initial management.

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**WHAT DO WE UNDERSTAND BY THE TERM “IMMUNOTHERAPY”?**

Immunotherapy is any treatment aimed at boosting or enhancing the immune system. It includes a wide range of options, from vaccines to treatment for conditions such as allergy and cancer. In the case of cancer, unlike other available treatments, immunotherapy is not aimed at destroying the tumor cells but at stimulating the patient’s immune system so that it attacks the tumor. In cancer, immunotherapy provides a series of advantages, such as targeted treatment, which only acts on tumor cells without damaging healthy cells, and immunological memory, which can subsequently be reactivated to recognize and attack the tumor once the immune system is stimulated. Immunotherapy is also subject to disadvantages, such as the time necessary for it to take effect-the immune response is not immediate but gradual-and the associated adverse effects. There are 2 main types of immunotherapy: Specific immunotherapy, which causes a response to a specific cell or antigen and includes vaccines and adaptive cell therapy; and nonspecific immunotherapy, which is aimed at stimulating the whole immune system and includes cytokines and regulatory proteins such as antibodies to cytotoxic T lymphocyte–associated antigen 4 (CTLA-4) or the programmed cell death protein 1 and its ligand PDL1 (PD1/PDL1) pathway. Also of interest are monoclonal antibodies, which are included in another group, known as passive immunotherapy, and comprise molecules designed to recognize tumor cells or substances that the tumor needs for growth. These are administered intravenously and can destroy tumor cells or deprive them of the essential components they require for growth. They are sometimes combined with other treatments in order to enhance their effect[1,2].

Ipilimumab is a monoclonal anti-CTLA-4 immunoglobulin 1 (IgG1) antibody that activates destruction of regulatory T cells by stimulating antibody-mediated cytotoxicity, thus halting the immunosuppressive effect. Ipilimumab is currently approved for the treatment of advanced melanoma (unresectable or metastatic) in monotherapy or in combination with nivolumab (anti-PD1)[3-8].

The most advance therapy attempts to inhibit the PD-1/PD-L1 checkpoint pathway. The PD1 receptor is found in lymphocytes and acts as an inhibitory-type checkpoint by interacting with its ligands PDL1 and PDL2. Tumor cells are capable of expressing PDL1, thus inhibiting the action of the cytotoxic T cell on them and generating tolerance. In this way, anti-PD1 drugs (nivolumab, pembrolizumab, durvalumab) and anti-PDL1 drugs (avelumab, atezolizumab) overcome tumor immune tolerance and enable the action of cytotoxic T cells. Anti-PD1 and anti-PDL1 agents have several indications in various scenarios, both in monotherapy and in combination therapy in patients with melanoma, lung cancer, renal cell carcinoma, urothelial carcinoma, classic Hodgkin lymphoma, and head and neck tumors[9].

**PATHOPHYSIOLOGY OF GASTROINTETINAL TOXICITY**

Toxicity of the various types of immunotherapy, or immune-related adverse effects, is closely associated with the mechanism of action of the different treatments, for example, the toxicity of interleukin 2 (IL-2) results from release of nitric oxide, IL-1, tumor necrosis factor alpha, and interferon gamma. In adoptive cell therapy, the initial toxicity is caused by lymphodepleting chemotherapy. After T-cell infusion, immunotoxicity manifests as fever, tachycardia, vascular hyperpermeability followed by multiorgan failure. In the most severe cases, this is due to cytokine release syndrome[10]. Blockade of CTLA-4 suppresses the function of regulatory T cells, which contribute to local inflammation in the gastrointestinal mucosa. It has been suggested that colitis associated with CTLA4 could result, in part, from this immunosuppressive function[11]. PD-1 and CTLA-4 blockade can generate toxicity that mimics autoimmune diseases[12].

**HOW TOXIC IS IMMUNOTHERAPY FOR THE GASTROINTESTINAL TRACT?**

Adverse effects involving the digestive system are recorded in around one-third of all patients receiving immunotherapy, specifically monoclonal anti-CTLA-4 IgG1 and anti-PD1 IgG4 antibodies[13].

Immune-mediated enterocolitis is one of the most common adverse effects, especially with ipilimumab. Up to one-third of patients treated with ipilimumab experience diarrhea, and immune-mediated colitis is observed in 7%-22% of cases[14]. In contrast, immune-mediated enterocolitis associated with nivolumab is less common, affecting around 10% of all patients who receive it[15]. The combination of ipilimumab and nivolumab is even more toxic than when each agent is used separately[8].

**ENDOSCOPY**

***When should it be performed?***

Colonoscopy with biopsies is the standard diagnostic approach for patients with lower digestive symptoms (*e.g.*, diarrhea, hematochezia). It is recommended in patients who receive immunotherapy and have persistent diarrhea or associated poor prognostic factors (hospitalization due to oral intolerance or absence of response to corticosteroids). Upper gastrointestinal symptoms (*e.g.*, dysphagia, gastroesophageal reflux, epigastralgia) are not uncommon and necessitate gastroscopy[16].

***Which are the main endoscopy findings?***

**In the colon:** In their series of 39 patients receiving anti-CTLA-4, Marthey *et al*[17] reported the most common lesions in endoscopy to be ulcer (79%) (Figures 1 and 2), erosion (13%), and erythema (8%) (Figure 3). The rectum and/or sigmoid colon were involved in 97% of cases, with extensive colitis being observed in 66% of patients. The distribution of the lesions was patchy in 55% of cases. The ileum was affected in only 5 patients.

In their study of 40 patients receiving treatment with anti-CTLA-4 agents who developed diarrhea and underwent flexible sigmoidoscopy or colonoscopy, Beck *et al*[18] reported findings for 36 cases. Again, the most common findings were erythema and ulcer (in 63% of patients). Endoscopy revealed inflammation several months after onset of enterocolitis, thus suggesting that in some cases, enterocolitis induced by anti-CTLA-4 agents can progress to inflammatory bowel disease (IBD).

We can find similar results for the type of involvement in immune-mediated colitis caused by anti-PD1 agents, which is less frequent than that associated with anti-CTLA-4 agents. Collins *et al*[15] studied a series of 20 patients with diarrhea who were receiving treatment with anti-PD1 agents and in 12 of whom colonoscopy findings were abnormal. The most frequent location was the descending colon (83%), and a patchy pattern, rather than a continuous pattern, was the most common (found in approximately 73% of patients). The most common lesions are also the same as those described above, namely, erythema, erosion, and ulceration.

**Esophagus-stomach-duodenum:** In their study of 22 patients who underwent gastroscopy, Marthey *et al*[17] found lesions in 13 (60%), the most common being gastritis (Figures 4 and 5) and erosive duodenitis.

Similar results were reported by Collins *et al*[15], who found that 63% of patients also presented upper gastrointestinal lesions. Erythematous gastritis was reported in most cases.

**Biliary tract:** The scientific literature also contains references to biliary involvement, more specifically in cases of extrahepatic cholangitis[19] and toxic hepatitis[20]. In such cases, endoscopy can be based on retrograde cholangiopancreatography or cholangioscopy. To our knowledge, no endoscopic images of the biliary tract obtained by cholangioscopy have been reported.

**Small intestine:** To our knowledge, there are no published capsule endoscopy–based data on involvement of the small intestine as a result of toxicity induced by immunological therapy.

***Other aspects associated with endoscopy***

Other lesions seen in endoscopy include exudates, granularity, and loss of vascular pattern. These findings are similar to those of IBD (ulcerative colitis and Crohn disease). Endoscopy findings from the first 2 wk of treatment with the drug and before the onset of symptoms have not been shown to predict the development of immune-mediated colitis. Endoscopy-confirmed involvement and the need for infliximab are not more common in patients receiving high-dose anti-CTLA-4 agents, thus suggesting that the severity of enterocolitis is not dose-dependent[21,22].

**WHICH DIFFERENTIAL DIAGNOSIS SHOULD BE MADE?**

Suspicion of toxicity due to immunotherapy should be based on the presenting complaints, of which diarrhea is the most common. Therefore, immune-mediated colitis should be taken into consideration in any patient receiving treatment with anti-CTLA-4 and/or anti-PD1 agents and who presents compatible symptoms. Other possible presenting complaints are abdominal pain, vomiting, hematochezia, weight loss, and/or fever. Onset of symptoms may be at any time during treatment and even several months after the last dose. The main conditions in the differential diagnosis are tumor progression, the infectious causes of diarrhea, and the development of IBD (Table 1). Therefore, imaging tests should also form part of the extension study for the primary tumor. Feces should also be tested for parasites and *Clostridium difficile*, and other tests, such as diagnostic colonoscopy, should be performed[21].

The main laboratory abnormalities in patients receiving immunotherapy are anemia, increased C-reactive protein, and low levels of serum albumin, all of which are nonspecific and play no role in the differential diagnosis. Therefore, endoscopy is the key to diagnosis. However, the result of a macroscopically normal endoscopy does not rule out the diagnosis, and biopsy specimens should be taken throughout the colon and assessed according to the segment they came from. Furthermore, infection by cytomegalovirus should also be ruled out by immunohistochemical staining of the biopsy specimens.

Histopathology findings are compatible with acute colitis, which is characterized by a marked inflammatory cellular infiltrate in the lamina propria consisting of neutrophils, lymphocytes, plasma cells, and eosinophils. Occasional findings include foci of neutrophilic cryptitis, crypt abscess, gland destruction, and erosions of the mucosal surface[23].

The histological characteristics of immune-mediated colitis are often nonspecific and may mimic those of other types of colitis. However, a variety of histologic characteristics that can act as useful pointers have been reported. Active colitis, together with major apoptosis of the epithelial cells in the crypt, has been recognized as the most useful characteristic. Other, less common associated patterns include lymphocytic and collagenous colitis. The correlation with the clinical history and, in particular, exposure to the drug plays an essential role in enabling the pathologist to differentiate immune-mediated colitis from infectious colitis, IBD, and drug-related colitis[24].

**HOW SHOULD THE DISEASE BE TREATED?**

Management of the patient with suspected immune-mediated enterocolitis should be multidisciplinary, involving oncologists, gastroenterologists, endoscopists, and the intensive care unit.

Treatment is mainly medical, and endoscopy is used only for diagnosis. Treatment of mild diarrhea (fewer than 3 watery stools per day) is based initially on oral antidiarrheal drugs together with fluid-electrolyte replacement. In moderate cases or absence of response, treatment should be started with oral corticosteroids (prednisone or equivalent at 0.5-1 mg/kg per day). In cases of severe diarrhea (more than 6 watery stools per day), treatment with anti-CTLA-4 and/or anti-PD1 agents should be suspended permanently, and intravenous corticosteroids should be started (methylprednisolone or equivalent 1-2 mg/kg per day). Patients who do not have a clinical response to intravenous corticosteroids after 3 d of treatment should start biologics (infliximab in a single dose of 5 mg/kg). The response to infliximab is generally fast, although some patients may require a second dose after 2 wk[21,25]. Marthey *et al*[17] reported that 37% of patients were treated successfully with corticosteroids. Biologic therapy was necessary owing to resistance to corticosteroids in 30% of cases (12 of 39 patients); infliximab was successful in 83% of cases (10 of 12 patients). Given the favorable response to infliximab, this therapy should be intensified rapidly in patients who do not respond to corticosteroids and whose clinical course is indolent. Treatment with corticosteroids during the first 5 d after onset of symptoms can enable more rapid resolution of symptoms than later initiation of treatment[26].

Perforation of the colon, while potentially fatal, is uncommon (< 1%). However, when surgery is necessary, colectomy should be subtotal and not segmental, since in most cases, enterocolitis induced by anti-CTLA-4 agents affects the whole colon[27-29].

**CONCLUSION**

Immune-mediated colitis is an emerging condition, given that the indications for immunotherapy, specifically anti-CTLA-4 and anti-PD1 agents, are expected to increase over time and for different types of tumor. Therefore, it is important to know the symptoms and determine the degree of involvement of immune-mediated colitis using endoscopy in order to initiate appropriate treatment early. The differential diagnosis should be based on infection, tumor progression, and IBD, with which the disease shares symptoms, endoscopy-confirmed lesions, and treatment.

The most common endoscopy-confirmed lesions of immune-mediated colitis are ulcers and erosions on edematous and erythematous mucosa. The most common location is the sigmoid colon, and a segmental pattern is slightly more common than a continuous pattern. Although histopathology is not specific in immune-mediated colitis, a biopsy must be taken to rule out other diseases and make a definitive diagnosis. Treatment includes systemic corticosteroids, although biologic therapy with infliximab may be necessary in some cases. Lastly, we believe that it is of the utmost importance to perform new studies that provide a detailed description of the adverse effects of regulatory proteins and of new, forthcoming agents in order to improve recognition and treatment of immune-mediate colitis in daily clinical practice.

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Grade D (Fair): 0

Grade E (Poor): 0

**Table 1** **Differential diagnosis**

|  |  |  |
| --- | --- | --- |
| **Disease** | **Endoscopy findings** | **Clinical characteristics** |
| IBD | UC: Continuous and circumferential mucosal inflammation starting in the rectumCD: Deep fissures, cobblestoning, segmental distribution, relative rectal sparing, and terminal ileal involvement | Rectal bleeding, abdominal pain, diarrhea, chronic anemia |
| Radiation colitis | Similar to IBD | Rectal bleeding, chronic anemia |
| Infectious colitis | Diffuse effects on the colon | Dysentery-like diarrhea, different agents, *Clostridium difficile* and CMV to be ruled out |
| Colitis associated with diverticulosis | Segmental distribution, peridiverticular, sigmoid colon affected, rectum and proximal colon are normal | Rectal bleeding, abdominal pain, diarrhea |
| NSAID-induced colitis | Any part of the intestine, isolated lesions | Recurrent abdominal pain, obstruction, perforation, hemorrhage, chronic anemia |
| Microscopic colitis | Normal endoscopy findings | Watery diarrhea |
| Ischemic colitis | Segmentary colitis (sigmoid /left colitis) | Acute onset of abdominal pain and rectal bleeding |

IBD: Inflammatory bowel disease; UC: Ulcerative colitis; CD: Crohn’s disease; CMV: Cytomegalovirus; NSAID: Nonsteroidal anti-inflammatory drug.

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**Figure 1 Mucosa at the rectosigmoid junction with mild erythematous spots and no erosions or ulcers.**

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**Figure 2 Mucosa at the rectosigmoid junction with erythema and fibrin-covered superficial erosions.**

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**Figure 3 Mucosa in the descending colon with extensive erythema and deep fibrin-covered ulcers.**

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**Figure 4 Erosion on the mucosa of the gastric antrum with generalized erythema.**

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**Figure 5 Petechiae on the mucosa of the gastric fold.**