**Infections and Chronic Inflammatory Bowel Disease**

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**Summary**  
In the more recent years since the introduction of anti-TNF therapy, the treatment strategy in chronic inflammatory bowel disease has developed more towards an early intensive, often double immunosuppression. While this leads to an improved therapeutic success, this intensified therapy also increases the risk for side effects and especially for infectious complications. The early detection of this complication in the immunocompromised patient is often more difficult due to the potential broad spectrum of infectious agents, the often atypical presentation in conjunction with the immunosuppression as well as often similar symptoms regarding intestinal infectious complications common for a flare of the underlying disease. In the first part, this overview will discuss the broad spectrum of potential infectious complications, using pulmonary infections as an example and presenting an algorithm for detection and therapy. In the second part, common intestinal infectious complications will be discussed from diagnosis to therapy.

**Introduction**  
While the current treatment strategy of immunosuppression in patients with active inflammatory bowel disease (IBD) leads to a remission or at least reduction of symptoms in most patients, this strategy also induces an increased risk for infections, intestinal as well as extraintestinal. In order to ensure a safe therapy, early detection of infectious complications and initiation of the treatment is crucial.
What Kind of Infections Can Potentially Be Expected under Immunosuppression in Patients with Inflammatory Bowel Disease?

Various case reports, cohort evaluations as well as analyses of multicenter trials regarding infectious complications have been published. Toruner et al. [1], for example, identified 100 consecutive IBD patients with opportunistic infections. The infectious spectrum in his cohort included viral infections, e.g. herpes simplex virus (HSV), herpes zoster, and cytomegalovirus (CMV), bacterial infections, e.g. *Escherichia coli*, *Mycobacterium marinum*, and *Streptococcus*, as well as fungal infections, e.g. *Histoplasma capsulatum* and *Cryptococcus neoformans* [1]. These examples just give an idea of the broad spectrum one can expect as potential infectious complications. In addition to more or less rare infections, the risk for classical infections such as urinary tract infections, skin infections, or respiratory infections is also increased in the immunocompromised host.

As the spectrum is broad, this article will focus on two major areas, i.e. infectious respiratory complications and infections mimicking an acute flare of the underlying IBD.

Infectious Respiratory Complications

Starting with respiratory complications, the most prominent, though not the most common candidate for respiratory infections with the introduction of anti-TNF therapy has been the reactivation of latent tuberculosis, leading to severe infections with a lethal course in some of these patients [2].

With the awareness of these potential complications and the introduction of mandatory screening for latent tuberculosis prior to anti-TNF induction, this specific complication could be dramatically reduced, demonstrating the effectiveness of prevention. In most countries, these preventive measures include taking the history regarding potential exposures to open tuberculosis in the past, chest radiography, and an interferon-gamma release assay [3]. However, several issues need to be taken into account. For one, testing under immunosuppression such as thiopurines might not yield valid results due to unresponsive lymphocytes, and negative pulmonary imaging does not rule out latent or active tuberculosis. Furthermore, one has to keep in mind that an exclusion of tuberculosis before the beginning of treatment does not rule out a new infection with tuberculosis later in life. Therefore, re-testing should always be considered in patients with signs of unknown infection, especially following potential exposure such as after travelling to countries with a high prevalence of tuberculosis [4]. One also has to keep in mind that when tuberculosis occurs in patients on anti-TNF therapy, for example, it is more commonly atypical (extrapulmonary in <50%, disseminated in 25% of cases), with intestinal tuberculosis mimicking active Crohn’s disease, therefore making the correct diagnosis even more difficult [3, 5].

A further strategy to reduce pneumonia is to vaccinate. Guidelines such as from the European Crohn’s and Colitis Organisation (ECCO) recommend vaccination against seasonal influenza on a yearly basis as well as pneumococcal vaccination according to national guidelines [3].

Immunocompromised patients presenting with respiratory symptoms should be carefully examined for pneumonia. In the case of classical signs for bacterial pneumonia, positive auscultation, and typical infiltrate in a chest radiograph, antibiotic treatment in these patients under immunosuppression should always cover *Streptococcus pneumoniae*.

Patients with inflammatory disease on immunomodulator therapy with pneumonia should also be tested for *Legionella pneumophila* [3]. Testing can be done by screening a urine sample for *L. pneumophila* antigen, for example. The most common route of transmission is airborne from warm water supplies containing water that was not heated above 70°C as well as air conditioning and humidifiers. As there is no available vaccination, the best preventive measurements are maintenance of water systems and adequate heating of the warm water supply. Curative treatment consists of macrolide or fluoroquinolone antibiotics [6, 7].

A further pulmonary infection to be considered in patients under immunosuppression is an infection with *Nocardia* species, an aerobic Gram-positive, weakly acid-fast actinomycete [8, 9], especially in patients receiving anti-TNF and particularly when also being treated with corticosteroids [3]. *Nocardia* is found worldwide in soil and can lead to pulmonary infections through inhalation, with hematogenous dissemination to the brain in up to 33% [8]. Screening for infection requires examination of sputum, pleural or bronchial lavage fluid by Gram stain, and a modified acid-fast stain. Antibiotic treatment consists of a continuous treatment with sulfamethoxazole/trimethoprim and/or ceftriaxone until the disappearance of all lesions, which can take month [8].

In addition to bacterial-caused pneumonia, patients under immunosuppression appear to be also at risk for parasitic as well as fungal infection, though the knowledge and recommendations regarding IBD patients are mainly based on case reports and smaller case series. As these pulmonary infections often present with atypical symptoms and cannot be detected purely by clinical examination and standard chest radiography in many cases, suspicion should always be raised in immunosuppressed IBD patients complaining about breathlessness, cough, and reduced strength leading to a low threshold for performing CT scans and bronchoscopy with bronchoalveolar lavages (BAL). The screening should then include *Mycobacteria*, *Strongyloides stercoralis*, *Nocardia*, *Histoplasma*, *Cryptococcus sp.*, *Aspergillus sp.*, and *Pneumocystis jiroveci*. The diagnosis of *P. jirovecii*, an atypical fungus, is based on identification in bronchopulmonary secretions or BAL fluid, and the gold standard for *Candida* infection is a positive culture from normally sterile body sites, meaning that cultures from other sites have to be interpreted in the clinical context as cultures.
Infections Mimicking a Flare

Clostridium difficile

Epidemiology

In the past few years, the incidence of *Clostridium difficile* infection (CDI) has increased rapidly, with an infection rate that has doubled from 1996 to 2003 [16]. The burden of CDI has increased dramatically, and it is now recognized that CDI is responsible for 20–30% of cases of antibiotic-associated diarrhea and up to 75% of cases of antibiotic-associated colitis [17, 18]. IBD has been found to be associated with *C. difficile*, and patients with IBD appear to be at an increased risk of developing CDI with a poorer outcome of CDI including higher rates of colectomy and death as well as higher rates of recurrence [19–22]. According to the recent studies, it appears that CDI is more common in patients with ulcerative colitis (UC) compared to Crohn’s disease patients where an increase of CDI cases was only found in patients with large bowel disease [23]. In one study, CDI rates in UC patients have doubled from 2.66 to 5.12% in 7 years [24]. In addition to the increased prevalence, the number of IBD hospitalizations complicated by CDI has increased from 1.4% in 1998 to 2.9% in 2007 [25]. It has to be noted that *C. difficile* in IBD patients may not only affect the large bowel but also the small bowel, with high mortality rates which had frequently been shown in this population during the last decades [26]. CDI is often seen in patients who underwent proctocolectomy for a severe IBD [27] and also in patients with *C. difficile* pouchitis with CDI involvement in up to 18% of cases [28].

Risk Factors

Risk factors associated with CDI in the general population are antibiotic use, older age, hospitalization, immunosuppression, residence in long-term care facilities, cancer, and gastrointestinal disorders. IBD has been found to be an independent risk factor for CDI with a threefold increase compared with non-IBD patients [29]. There are conflicting data whether other factors associated with IBD including immunosuppression also play a role. It was found that glucocorticoid initiation with or without use of other immunosuppressive agents cannot differentiate colonization from infection in these cases. However, the sensitivity for e.g. *Aspergillus* sp. in BAL is only 50% and a definitive diagnosis may require invasive procedures such as biopsies. Similarly, the sensitivity for detection of *Histoplasma capsulatum* is only 50% [3]. Though the risk for these infections appears to be increased under immunosuppression, the actual number of IBD patients affected is fortunately relatively low. This implicates, however, that the individual experience of the treating gastroenterologist/surgeon is usually relatively low. Therefore, in unclear cases one should always consider consulting an infectious disease specialist for expert opinion, where available. The same holds true for the treatment of more rare infectious complications. The first-line treatment of *P. jirovecii*, for example, comprises the application of trimethoprim plus sulfamethoxazole for about 3 weeks, which requires monitoring of the blood count and the renal functions as this treatment may cause bone marrow suppression or renal impairment requiring dose adjustment [3].

While there are no effective immunizations available for the infectious agents discussed above and screening for these is not recommended in asymptomatic patients prior to initiation of immunosuppression, chemoprophylaxis for *P. jirovecii* is a potential option, at least for some patients. The ECCO guidelines as well as the German Colitis guideline recommend a standard prophylaxis with co-trimoxazole, if tolerated, for those patients on triple immunomodulators with one being a calcineurin inhibitor or anti-TNF therapy [3, 10]. However, this recommendation is mainly based on expert opinion as risk-benefit studies for IBD patients are lacking, and therefore recommendations have to be extrapolated from patients with HIV infection and hematological diseases.

As discussed below in more detail for CMV, the potential spectrum for pulmonary infections also includes viruses such as from the family of herpes viruses, HSV, varicella zoster virus, Epstein-Barr virus, and CMV. Primary infection with e.g. HSV in immunocompetent individuals usually causes an asymptomatic or mild, self-limited oral-labial (HSV type 1) or genital (HSV type 2) infection [11]. In immunocompromised patients, especially in those treated with azathioprine, HSV infection has a greater potential for dissemination possibly causing severe systemic infections with significant morbidity and mortality including encephalitis, meningitis, pneumonia, esophagitis, colitis, and/or hepatitis [12–15]. While increasing titers of anti-HSV IgM (immunoglobulin M) can help making the diagnosis, the diagnostic gold standard is the PCR (polymerase chain reaction) from affected tissue [11]. Systemic therapy with the nucleoside analogue aciclovir is then required in symptomatic immunocompromised IBD patients [3, 11].

An algorithm for immunocompromised IBD patients with pulmonary symptoms is shown in figure 1.
current recommendations on CDI diagnosis implement a molecular assay or a two-step algorithm encompassing screening with enzyme immunoassay for glutamate dehydrogenase (GDH) followed by an enzyme immunoassay for toxins [36]. There is no evidence that testing for CDI should be done differently in IBD patients. On behalf of the physician, a high level of suspicion for CDI in IBD patients is required, and stool tests should be performed in every patient with IBD relapse. It should be noted that testing for CDI in non-IBD patients requires 1–2 probes; however, repeated testing up to 3 probes in IBD increases sensitivity.

**Treatment**

Even though local host defenses could be compromised in IBD due to the altered gut microbiota or the use of immunomodulators, treatment of CDI in IBD patients is not fundamentally different compared to non-IBD patients. Currently, there are no clear guidelines regarding the treatment of CDI in IBD patients. Most authors suggest that the initial episode of CDI in IBD patients with mild-to-moderate disease scores should be treated with metronidazole (table 1). Patients with severe or complicated disease should be treated with vancomycin during the initial episode. *C. difficile* resistant to metronidazole has generally been uncommon in IBD patients even though failure rates of up to 50% have been reported in this population [37, 38]. There is also no supporting evidence how to proceed with the ongoing immunosuppression in IBD patients with CDI. Most authors maintain immunosuppression in patients with CDI although an escalation of immunosuppression is avoided [39]. Severe CDI disease is usually treated in combination with vancomycin orally plus metronidazole i.v.

In addition to metronidazole and vancomycin, other antibiotics like rifaximin or fidaxomicin may be used in severe or recurrent CDI. Beside antibiotics, fecal microbiota transplantation may be another option to treat concurrent or particularly recurrent CDI in IBD patients. The rationale behind fecal microbiota transplantation comes from the notion that an introduction of feces from the healthy host would recolonize the bowel with microbiota needed to reestablish colonization resistance against *C. difficile*. Even though there is no general protocol for the administration of fecal microbiota transplantation, several attempts have been made to treat CDI in IBD patients with more or less reasonable results [40]. There is currently no evident role of probiotics in the prevention or treatment of CDI in IBD patients. The treatment of recurrent CDI is even more complicated, and conflicting data exist on how to treat in this situation. Options include the use of the same treatment regimen as for the initial episode up to the use of alternative antibiotics such as rifaximin, fidaxomicin, or fecal microbiota transplantation in this situation [41].

In summary, CDI in IBD patients is a major challenge as the clinical importance is still underestimated. Because of the high mortality and severity of CDI in patients with IBD, early recognition in these patients is of particular importance.

### Table 1. Treatment of CDI infection in IBD (according to IDSA guidelines [36])

| Disease category                        | Treatment                                                                 |
|----------------------------------------|--------------------------------------------------------------------------|
| Mild-to-moderate disease – initial episode | metronidazole 500 mg three times/day for 10–14 days                      |
| Severe disease, uncomplicated – initial episode | vancomycin 125 mg four times/day for 10–14 days                          |
| Severe disease, complicated – initial episode | vancomycin 500 mg four times/day plus metronidazole 500 mg every 8 h i.v. |
| Recurrence                              | same as for initial episode; alternative options: rifaximin, fidaxomicin, fecal stool transplantation |
Cytomegalovirus Infection

The impact of CMV in IBD remains controversial [42]. The pathogenicity of CMV reactivation in IBD patients has been a debate in the literature during the last decade. However, despite arguments on the non-pathogenicity of CMV reactivation, there is overall agreement that IBD patients are at an increased risk of CMV reactivation and that failure to ensure the appropriate diagnosis and treatment may lead to significant morbidity and mortality. Subclinical reactivation of latent CMV infection may occur frequently and is often associated with immunomodulator therapy [42, 43]. Usually, the reactivation of latent CMV is asymptomatic and does not induce serious tissue damage. It is therefore required to distinguish between CMV infection which may be detected by CMV DNA and CMV disease which may cause organ damage in the colon, liver, lung, or other organs [44]. Mild CMV reactivation without tissue damage is nearly always self-limited even under continuation of immunomodulator or biological treatment [45–47]. Therefore, a stop of immunomodulator treatment in patients with CMV reactivation is usually not required.

Clinical Presentation/Diagnosis

Screening for CMV infection in all IBD patients is not indicated unless the patients are steroid-resistant. In patients with acute steroid-resistant colitis, CMV should be excluded [44]. Different techniques are used to diagnose CMV infection including endoscopy, histology, serology, as well as CMV antigen and CMV DNA testing. The use of the different techniques is controversial, and there is no international gold standard for the detection of CMV infection. The use of CMV serology is of limited value for diagnosing an acute infection because of its failure to detect CMV reactivation. However, CMV serology may identify patients who are CMV-IgG-positive and thus at risk for CMV reactivation [48]. CMV antigen and DNA testing may act as reliable markers for disseminated infection. Even though the antigenemia assays are only semiquantitative, they may act as indirect markers and are sufficient to monitor infection as well as antiviral treatment in the immunocompromised host. More sensitive is the detection of DNA viral load by PCR [49]. The advantages of qualitative and quantitative testing as well as of rapid and high sensitive results favor the detection of CMV DNA load by means of PCR in the blood or in tissue biopsies [49, 50]. In steroid-refractory colitis, CMV has been detected by PCR in up to 36% of patients [51]. Detection of CMV DNA viral load of more than 250 copies/mg may be used as a predictor of steroid-resistant disease [52]. An alternative for the detection of CMV antigen or CMV DNA viral load may be the use of histopathology combined with immunohistochemistry. Monoclonal antibodies against CMV early antigen are highly specific and sensitive for verifying CMV infection in tissue biopsies [53]. Therefore, CMV is most commonly excluded by tissue PCR or immunohistochemistry [44].

Treatment of CMV Colitis

In the case of severe steroid-resistant colitis with CMV detected in the mucosa during immunomodulator therapy, antiviral therapy should be initiated and immunomodulators considered to be discontinued until colitis symptoms improve [44]. The therapy of choice for CMV infections is ganciclovir for a period of 2–3 weeks [54]. After 3–5 days a switch to oral valganciclovir for the rest of the period of 2–3 weeks may be considered [49, 55, 56]. In patients where ganciclovir is associated with side effects such as myelotoxicity or in those with ganciclovir resistance, foscarnet for a period of 2–3 weeks may be an alternative [42, 57]. Subclinical CMV reactivation or mild CMV infection does not require antiviral treatment or interruption of immunomodulator therapy as it usually completely recovers [54]. In the case of CMV disease with CMV reactivation causing organ disease such as hepatitis, pneumonia, esophagitis, colitis, or meningoencephalitis, immediate antiviral treatment with ganciclovir and discontinuation of immunosuppression is usually required due to the poor outcome in this situation [55, 58, 59].

All in all, screening for CMV infection is usually not required in IBD patients before immunomodulator therapy has been started. In patients with acute steroid-resistant colitis or in patients with acute flares under steroid treatment, CMV infection should be excluded either by PCR or immunohistochemistry. According to the recent ECCO guidelines, patients with severe steroid-resistant colitis with CMV infection during immunomodulator therapy should be treated with antiviral drugs until colitis symptoms improve. In the case of CMV disease with systemic organ manifestation, immunomodulator therapy should be discontinued.

In summary, there is a broad spectrum of potential infectious complications in immunocompromised patients. Some infections mimic a flare of the underlying chronic bowel disease, while others present in an atypical fashion due to the immunosuppression. This demonstrates the necessity for carefully taking the patients’ history especially regarding new symptoms. Furthermore, patients should clearly be advised to report new symptoms to the treating physician. Carrying a document listing the taken immunosuppression, which can be presented to the emergency room team in the case of urgent hospital admissions, might contribute to infectious signs not being underestimated in the often young IBD patients.

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