

Inflammatory bowel disease management during the COVID-19 outbreak: The 10 do's and don'ts from the ECCO-COVID Taskforce

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Abstract

Our knowledge on COVID-19 is changing and evolving rapidly, with novel insights and recommendations, almost on a daily basis. It behooves the medical community to provide updated information on a regular basis, on best practice to facilitate optimal care of infected patients and appropriate advice for the general population. This is particularly important in the case of patients with chronic conditions, such as Inflammatory Bowel Disease (IBD).

In this paper, we compiled the existing evidence on the impact of COVID-19 in IBD patients and provide guidance and on the most appropriate care to adopt during the pandemic. Our review highlighted that IBD, *per se*, is not a risk factor for COVID-19. However, all IBD patients with symptoms should be tested for SARS-CoV-2 and the procedures for disease management shall be carefully adapted: i) in SARS-CoV-2 positive IBD patients, medical treatments should be re-evaluated (with particular focus on corticosteroids) always with the purpose of treating active disease and maintaining remission; ii) non-urgent surgeries and endoscopic procedures should be postponed for all patients; iii) online consultancy should be implemented; and iv) hospitalization and surgery should be limited to life threatening situations.

Introduction

At the end of 2019, a cluster of patients with pneumonia emerged in Wuhan, Hubei Province, China, with possible zoonotic origin.¹ This unidentified pneumonia was later found to be related to a novel coronavirus (CoV)² named as severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), by the International Committee on Taxonomy of Viruses (ICTV).³ To prevent the use of inaccurate or stigmatizing names, the World Health Organization (WHO) proposed a standard format for the disease nomenclature: COronaVirus Disease-2019 (COVID-19).⁴

SARS-CoV-2 is a single positive-stranded RNA virus enveloped in a lipid bilayer. The virus sneaks into the human body via the mucous membranes (mouth, nasal, ocular) and enters host cells through the binding of viral spike proteins (S-protein) to the human protein receptor angiotensin converting enzyme-2 (ACE2).⁵ This receptor is abundant in lung, heart, kidney, adipose tissue, esophagus, stomach, bladder, ileum, and colon.^{6,7}

Transmission of COVID-19 is efficient. Aerosol and fomite transmission of SARS-CoV-2 is plausible, since the virus can remain viable and infectious in aerosols for hours, and on surfaces up to days (depending on the inoculum shed).⁸ In fact, the concentration of viral RNA in airborne samples is minimal or close to zero⁹, but it can be detected on fomites including plastic.¹⁰ Transmission occurs from close contact and respiratory droplets, without evidence of airborne transfer¹⁰. Thus, people who are in close contact with patients, their family members or healthcare workers are part of the high-risk population.¹¹

The spread of SARS-CoV-2 identified two important modes of disease transmission: i) local transmission, at each local epicenter; and ii) transmission via international travelers which favored the global spread of the infection, fueling COVID-19 pandemic. Large outbreaks were reported in closed communities and hospitals, raising the possibility of 'superspreading' events, as reported before in previous CoV outbreaks.¹² By May 2020,

the pandemic had affected almost 2 million people, worldwide, accounting for over 125,000 deaths, with the numbers rising sharply (*Worldometer's COVID-19 data*).¹³

SARS-CoV-2 infection is associated with five different clinical courses: asymptomatic infection, mild to medium cases, severe cases, critical cases and death.¹¹ While highly transmissible, more than 80% of the patients have mild disease.¹⁴ In the face of these challenging circumstances, gastroenterologists need to adapt priorities, reset standards of quality of care, and guide patients by providing relevant information.

In this context, the European Crohn's and Colitis Organization (ECCO) gathered a group of gastroenterologists, with particular interest in opportunistic infections, and infectious disease experts, to deliver guidance to physicians in the setting of gastrointestinal (GI) diseases during COVID-19 pandemic. The aim of the current paper is to provide health care professionals with understanding and knowledge of the optimal care we can provide to our patients, including those taking immunomodulatory treatment.

1-General Measures

Do's

- Do avoid contact with infected people
- Do avoid touching eyes, nose or mouth with unwashed hands
- Do clean hands using soap and water or an alcohol-based solution
- Do avoid crowded places
- Do use face masks according to local policies
- Do ensure flu vaccine for all non-vaccinated IBD patients
- Do ensure pneumococcal vaccination for all non-vaccinated IBD patients
- Do discourage travelling

According to WHO^{15,16} and to the European Centre for Disease Prevention and Control (ECDC)¹⁷, social distancing, hand hygiene and respiratory etiquette should be adopted by all the population to prevent the spread of the infection. Hand hygiene is considered to be

a rational precaution, involving limited costs without associated risks. Hands should be washed regularly, using soap and water for 20–40 seconds, taking care to ensure no areas are missed¹⁷ (ECDC). In community settings, alcohol-based hand sanitizers provide limited added benefit over soap and water and, if used, should contain 60–85% alcohol. A combination of personal protective and environmental measures increases effectiveness. Such measures include routine cleaning of frequently used surfaces and objects (phones, tablets, keyboards), and minimizing the sharing of objects¹⁷ (ECDC). Surgical masks may be used as an infection control measure or as a mitigation measure, in community settings, when used by individuals with respiratory symptoms but that did not seek medical attention yet. People at high risk of exposure include care providers with extensive face-to-face contact. A recent meta-analysis on influenza prevention suggested that N95 respirators should not be recommended to general public and low risk medical staff.¹⁸ Travelling should be limited, for everyone, as much as possible, with the aim of reducing: i) acquisition of infection by travelers to areas or countries where community transmission is ongoing; ii) importation of cases from affected countries; and iii) transmission among travelers.

Patients with IBD, taking immunomodulatory therapy, have increased risk of influenza and pneumococcal disease.^{19,20} To avoid pulmonary comorbidities with influenza and pneumococcal disease during this outbreak, and next winter, all IBD patients should be vaccinated. Influenza and pneumococcal vaccination are recommended for the majority of patients with immune mediated disorders, including IBD.^{21,22} Since influenza infection rates can be reduced by annual vaccination, flu vaccine is recommended in all non-vaccinated IBD patients, and especially during COVID-19 pandemic. Although not numerous, some cases of influenza coinfection with SARS-CoV-2 have been described.^{23,24} During the next seasonal outbreak, SARS-CoV-2 and influenza are likely to occur simultaneously. Patients

who have not been previously vaccinated for *Streptococcus pneumoniae*, should receive a dose of PCV13 followed by a dose of PPSV23, at least 8 weeks later.²⁵

2-Should we stop drugs in patients without symptoms suggestive of COVID-19 (not tested or tested negative)?

Do's

- Do continue immunomodulators
- Do continue biologics
- Do continue JAK inhibitors
- Do reduce corticosteroids whenever possible
- Do keep infusions in an infusion center whenever possible

Don'ts

- Don't reduce the dose of immunomodulators or biologics to prevent SARS-CoV-2 infection
- Don't switch infliximab to adalimumab in a stable patient, unless it is not possible to provide intravenous infusions
- Don't assume that IBD patients are at increased risk of being infected

Don't Know

- Patients with IBD, who are exposed to SARS-CoV-2, have a higher risk of developing symptomatic or severe COVID-19

So far, IBD patients have not shown increased susceptibility to COVID-19^{26–28}. This is clearly evidenced in very recent studies, in Spanish, French and Italian IBD cohorts studied, in which the risk of infection, associated mortality and incidence of COVID-19 were similar to those reported for the general population.^{26,27} A recent study from Wuhan, performed during the local outbreak of the disease, including 318 patients with IBD, did not report a single COVID-19 case.²⁸ However, all biological and immunomodulatory treatments were discontinued rendering interpretation difficult.²⁹ The therapeutic approach for these patients should instead consider the risk of viral infection weighed against the risk

of disease recurrence. Quantifying the immediate and long-term risks of a new disease is challenging, especially within an immunosuppressed population. In theory, immunosuppression may reduce viral clearance, but may also reduce the cytokine storm implicated in Adult Respiratory Distress Syndrome (ARDS). However, current evidence does not suggest that patients on immunomodulatory therapy are faring worse than the general population, in the COVID-19 pandemic, either in risk of contracting SARS-CoV-2, or in disease severity. A study of 191 patients, in Wuhan, who had completed inpatient treatment for COVID-19, did not identify immunosuppressed status (due to transplantation, chemotherapy, or other conditions requiring immunomodulatory treatment) to be a risk factor for adverse outcomes.³⁰ A national Chinese dataset of 1099 hospitalised patients identified only 2 patients with immunodeficiency, neither of which developed severe disease.³¹ A large European center for pediatric liver transplantation, located in Lombardy, showed that among patients in follow-up for cirrhosis, transplantation, autoimmune liver disease, or chemotherapy for hepatoblastoma, none developed clinical pulmonary disease, despite several testing positive for SARS-CoV-2.³² To date, the SECURE-IBD database has identified 959 cases of COVID-19 in IBD patients. Within this cohort, immunomodulators, anti-TNF therapy, anti-integrins and anti-IL12/23 were not associated with increased mortality.³³ Therefore, in balance, we believe that immunomodulatory drugs should be maintained in IBD patients without COVID-19 infection.

There are biological reasons to support why COVID-19 may not pose increased risk to IBD patients on immunomodulatory therapy. **First**, it has been suggested that the soluble form of ACE2 acts as a competitive binding partner for SARS-CoV-2 sequestering the virus, hindering binding to the virus's cell surface receptor, the full-length ACE2 protein.^{34,35} Elevated levels of ACE2 were measured in the plasma from patients with IBD³⁶ with evidence of limited infection progression and low susceptibility to infection.³⁷ The role of

ACE2 was also evidenced through the use of a chemical inhibitor of ACE2 (GL1001), that was able to reduce dextran sulfate sodium (DSS) colitis severity, in the mouse model, suggesting that ACE2 plays a pathogenic role in colitis.^{35,38} **Second**, tumor necrosis factor (TNF) inhibitors might be effective in reducing organ damage.³⁹ This effect is achieved through the decreased shedding of the ACE2 ectodomain (mediated by TNF- α -converting enzyme) which is essential for the penetration of SARS-CoV-2 into the cell. **Third**, as IFN- γ and TNF production has been associated with severe SARS-CoV infection, inhibition of TNF has been proposed as a treatment for the cytokine release syndrome that can occur in some of these patients.⁴⁰ **Fourth**, mercaptopurine and 6-thioguanine have potential antiviral activity against MERS and SARS, at least *in vitro*.⁴¹ **Fifth**, tacrolimus is a potent *in vitro* antiviral for human coronaviruses.⁴³ **Sixth**, most of the drugs used in IBD have a long elimination half-life and keep some activity even after treatment cessation.⁴⁴ **Seventh**, viral infections did not increase among IBD patients receiving ustekinumab and vedolizumab therapy. In addition, in a phase I clinical trial, the viral load of HIV patients on vedolizumab did not change significantly. In conclusion, the available data suggest that immunosuppressed patients are not at increased risk for severe disease and complications, compared with the general population.

In circumstances where it is not possible to safely run an infusion service, it may be reasonable to consider switching to sub-cutaneous alternatives. However, this practice must be used judiciously since elective switching from infliximab to adalimumab is associated with a loss of tolerance and efficacy within 1 year.⁴⁵

3-Should we stop IBD drugs in patients who are SARS-CoV-2 positive, whether symptomatic or asymptomatic?**Do's**

- Do postpone biologics administration
- Do stop/reduce corticosteroids whenever possible
- Do stop azathioprine/mercaptopurine therapy
- Do stop azathioprine in patients in combination therapy with an anti-TNF
- Do stop JAK inhibitors

Don'ts

- Do not continue prednisone at doses above 20mg/day
- Do not restart the treatment until nasopharyngeal swabs PCR-SARS-CoV-2 tests (if available) with a negative result

Don't Know

- Patients taking oral budesonide and beclomethasone therapy must stop treatment if tested positive
- IBD-related drugs protect against severe forms of COVID-19 (related to cytokine storm)

In IBD patients, aged over 65 years, disease activity and comorbidities were significantly associated with COVID-19 pneumonia and COVID-19 related death, whereas concomitant IBD treatments were not.⁴⁶ In these patients, COVID-19 complications and lethality seem to be unrelated to the use of immunomodulatory therapy.⁴⁶

However, there is more evidence to suggest that corticosteroids may pose significant risk to the IBD patient population. Steroid use is associated with increased morbidity and mortality in other viral infections, including influenza, SARS-CoV and MERS-CoV, along with complications in survivors. A recent uncontrolled study, in the setting of COVID-19, suggested that patients taking high steroid dosages have significantly worse clinical outcomes (ARDS, shock, secondary infection) than patients without a history of steroid use.⁴⁷ Within the IBD population, at the time of writing, patients who entered into the IBD-SECURE registry, with steroid use at presentation, had strikingly high rates of ICU

admission (19%) and death (11%); although this data is currently unadjusted for age or comorbidity.³³ Concerning SARS-CoV-2 infection, steroids were not effective for the treatment of lung injury or shock.⁴⁸ However, short-term steroids (≤ 0.5 -1 mg/kg for 7 days) may be beneficial to control overwhelming inflammation and cytokine-related lung injury, particularly in severe forms of ARDS.⁴⁹⁻⁵¹ Data on low-dose and short-term steroids, budesonide and beclomethasone therapy are not currently available.

The first analysis of the SECURE-IBD data also identified 5-ASAs as a risk for severe COVID-19 infection (OR 3.1, 95% CI 1.3-7.7).³³ This may be justified by methodological reasons or by the action of 5-ASA on PPAR- γ , and subsequent perturbation of ACE2, the binding site of SARS-CoV-2. Thus, at the light of this data, and in the absence of more detailed information on the interaction between 5-ASA, PPAR γ , and ACE2, we recommend pausing 5-ASA therapy, in patients with confirmed infection.

Lymphopenia is associated with worse prognosis of COVID-19.⁵² The Saint-Antoine experience, with a total observation time of more than 15,000 person-years (4,800 for thiopurines, 3,800 for anti-TNF), reported 31 cases of serious viral infection (EBV, CMV, VZV, HSV), mostly in patients exposed to thiopurines.⁵³ This can be related to the ability of azathioprine/6-mercaptopurine and tofacitinib to reduce the number of activated T cells and affect T-cell activation and effector function.³⁵ Tofacitinib has also been associated with viral infections (VZV)⁵⁴ and demonstrated inhibition of interferon- α production *in vitro*.⁵⁵ The decision to pharmacologically immunosuppress a patient with COVID-19 remains difficult. The possible beneficial effects in reducing inflammation should be carefully weighed against the potential deleterious impairment of anti-microbial immunity.

4-Can we start drugs for an IBD flare?

Do's

- Do test all IBD patients for SARS-CoV-2
- Do treat active IBD according to the standard guidance, as before the pandemic
- Do consider subcutaneous drugs to minimize hospital visits, and home delivery service

Since COVID-19 symptoms can mimic IBD manifestations, all patients with a suspected IBD flare should be tested to exclude SARS-CoV-2 infection.⁵⁶ In a recent ECCO survey, most of the physicians (75.1%) considered SARS-CoV-2 testing unnecessary even in patients treated with immunomodulators or biological drugs (62.8%). On the other hand, in IBD patients with suspicious symptoms or active disease, SARS-CoV-2 testing was supported by 54.6% of the responders.⁵⁷ In early reports, 2%-10% of patients with COVID-19 had GI symptoms such as diarrhea and vomiting.⁴⁷ In the Lu et al cohort, 12% of the patients presented with GI symptoms at onset and 50% during hospitalization.⁵⁸

In the context of SARS-CoV-1, the virus was cultured from stool samples during the SARS 2002-2003 outbreak.^{59,60} Now, SARS-CoV-2 has been cultured from stool samples during the COVID-19 outbreak.⁶¹ However, in a series of 20 COVID-19 patients the infectious virus could not be isolated from stool samples, despite the high virus RNA concentration.⁶² In SARS-CoV-2 infection, nasopharyngeal swabs are more sensitive than oropharyngeal ones and are best taken when the first symptoms emerge⁶³. Swabs from both sites are often combined to increase sensitivity. Due to its simplicity, easy methodology and to the range of extensively validated standard operating procedures, RT-PCR is now the preferred and most widely used method for detection of current infection.⁶⁴ According to recent evidence, the sensitivity of many of the available RT-PCR tests, for

detecting SARS-CoV-2, may be lower than optimal. Several factors warrant consideration. These include poor sample quality, variable presence of virus in lower *versus* the upper respiratory tract and the timing of the samples. A study including 1014 suspected COVID-19 cases, who underwent multiple RT-PCR testing and chest computed tomography (CT),⁶⁵ showed that 88% of patients had positive chest CT scans, whilst RT-PCR positivity was found only in 59%. Combining epidemiologic evidence with the analytical sensitivity of the currently used RT-PCR assays, it is not surprising that at least two gray zones could be identified: i) the initial phase of infection, when the patient is still asymptomatic or only mildly symptomatic; and ii) later stages in which virus shedding may persist, though below the analytical sensitivity of some RT-PCR assays.⁶⁶ Rapid antigen tests would theoretically provide fast results (15-30 minutes) and low-cost detection. These tests are in launch phase and might prove sufficiently sensitive and specific⁶⁷, making them potentially very useful in IBD.

Serological antibody tests should be a useful supplement to RNA or antigen detection. Many individuals will not have been tested having had asymptomatic or minimally symptomatic infections, or had a negative test result (due to sampling timing) or a false-negative result in nucleic acid amplification technology (NAAT) tests⁶⁸. Serological tests do not provide a direct detection of the virus, but rather a measurement of the immune response to the infection, which can provide insights into the kinetics of this response. In terms of public health, serological tests are vital to: i) recognise those who overcame the infection and have developed an immune response; ii) identify those who can return to work; and iii) select donors of convalescent plasma, a potential treatment for patients with severe COVID-19.⁶⁹ However, these tests do not provide a direct detection of the virus, but rather a measurement of the immune response to the infection, which can provide insights into the kinetics of this response. ELISA detected IgM in more cases than NAAT, on day

5.5 of illness, and the combination of both tests detected 98.6% of cases versus 51.9% with a single NAAT.⁷⁰

In the case of negative test, IBD patients should be treated according to the standard guidelines, and the physician should promote and advise home drug delivery and remote patients' programs.

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5-How can we continue IV infusion clinics?**Do's**

- Do implement screening procedures for suspected COVID-19 in all patients
- Do implement measures to minimize crowding
- Do implement at least 2m distance between chairs
- Do impose the use of surgical face masks in all patients
- Do continue biologics at regular intervals and doses

Don'ts

- Don't switch infliximab to adalimumab in stable patients unless it is not possible to provide intravenous infusions
- Don't allow accompanying persons inside the hospital

6-Do patients need specific protective equipment? Do physicians need specific equipment when seeing IBD patients?**Do's**

- Do avoid close contact with people and wash hands frequently
- Do use face masks (patient and physician)

Don'ts

- Don't see patients with accompanying persons unless strictly necessary

7-How to manage outpatient IBD clinics?**Do's**

- Do implement telemedicine
- Do monitor at distance
- Do web report outcomes
- Do promote local labs with e-mail reports
- Do implement point-of-care biomarkers
- Do implement calprotectin measurement at home
- Do implement measure of drug levels (TDM) with rapid tests
- Do cross sectional imaging, including intestinal ultrasound instead of invasive procedures

Don'ts

- Don't hospitalize patients unless strictly necessary
- Don't schedule unnecessary appointments, limit to what is strictly decision-making

The maximal viral shedding of SARS-CoV-2 occurs in the early stages of infection, meaning that spreading from asymptomatic patients is a reality. In hospitals and clinics caring for a large number of COVID-19 patients, the rate of infection is high among health

workers.⁷¹ In IV infusion clinics, the first measure is to identify the patients who are possibly infected, since the approach and safety requirements will be different. Thus, symptomatic (fever, cough, fatigue, myalgia, expectoration, diarrhea, and loss of sense of smell or taste) or suspected (contact with a positive person) patients should be promptly tested and submitted to protective measures. The use of masks, in patients attending hospitals, is highly recommended, even in those without symptoms.^{72,73} In the above mentioned ECCO survey, most respondents (physicians) confirmed the use of protections during consultations (72.2%) and considered it important to advise IBD patients to wear protection equipment in their daily lives (53%).⁵⁷

When considering patients who are in regularly monitoring, virtual clinics or online consultancy is advisable. Patients can be asked to have had laboratory tests in advance, and complete a simple questionnaire on symptoms, concomitant medications, and relevant questions. To replace visits, the IBD team can schedule phone calls with the patients, at the same date and at the same time as the scheduled visit. If social isolation prevents the patient from going to get a stool tests, a home faecal test for calprotectin is a valid alternative, if available. This would be for all patients, including those in remission receiving subcutaneous biologics or small molecules therapies.⁷⁴ Telemonitoring, using specific questionnaires and calprotectin home test measurement, was studied before the COVID-19 era. It proved to be as safe as conventional follow-up, both in paediatric⁷⁵ and adult⁷⁶ populations. In fact, faecal calprotectin levels showed a significant correlation with endoscopic extent, mucosal healing, and histological activity.^{77,78} Moreover, non-invasive imaging, such as ultrasonography, is essential for diagnosis and monitoring because it is a low risk procedure with the advantage of providing a rapid assessment of disease activity (location, extension, inflammation, and presence of complications like fistula, strictures or intrabdominal abscesses).⁷⁹

In the case of patients enrolled in clinical trials before the outbreak, the sponsors can be asked to: i) postpone non-necessary follow-up visits or to replace them by virtual clinics; ii) identify local labs that can guarantee the regular lab tests required by the protocol; iii) manage home delivery of study drugs, mainly those of oral and subcutaneous administration. In this scenario, patients would go to the hospital only for key visits (end of induction, re-randomization, end of study) and to receive intravenous drugs, when these cannot be suspended. The administration of intravenous drugs can also be adapted, mainly regarding dose intervals. For instance, infliximab administration can be postponed to every 10 weeks.⁸⁰ Regarding vedolizumab, the GEMINI trial showed that patients randomized to placebo can maintain remission up to week 24. Therefore, postponing vedolizumab 4-8 weeks may be reasonable. However, maintaining the original schedule remains the best strategy in most cases.

**8-Can IBD patients on immunomodulator/biological treatment continue working?
What about Health Care professionals with IBD?**

Do's

- Do avoid contact with infected patients
- Do use masks when working
- Do redirect towards the lowest risk zones
- Favor teleworking when possible

IBD patients on immunomodulators do not seem to have a higher risk of infection than the rest of the population. However, due to disease specificities and possible side effects of the treatment, avoidance of SARS-CoV-2 requires an additional set of precautions. Thus, patients should take extra protective measures, over and above those of the general population. Some societies use the concept of “stringent social distancing” and “shielding”, with measures ranging from avoidance of crowded places to advice to not leave the house

altogether. The specifics vary between nations depending upon SARS-CoV-2 prevalence and disease control strategies. Within some European countries people to shield include patients on anti-TNF induction regimes and high doses of steroids and immunomodulators.^{81,82}

It is not rare to find IBD patients working in health care systems. Considering that this population experiences higher rates of infections than general population, it may be necessary to introduce extra precautions in health care workers with IBD. If possible, these professionals should be redirected from high risk areas (like emergency departments, infectious disease units, and intensive care units) to low risk areas and, when possible, working from home should be implemented. Protective Personal Equipment (PPE) should be used according to international guidelines, when homeworking is not viable.

9-Can we perform non-urgent endoscopy? If not, what is non-urgent endoscopy?

Do's

- Do perform endoscopy as priority 3 months after the infection rates decrease
 - ◆ IBD patients with mild-moderate flare-up confirmed by biomarkers, to exclude CMV infection
 - ◆ Longstanding IBD in surveillance for CRC, if prior dysplasia
 - ◆ Endoscopic resection in IBD patients known to have LGD/HGD colonic lesions already detected
- Do perform endoscopy within 3-6 months after the infection rates decrease
 - ◆ Pouchitis
 - ◆ IBD patients with flare-up not confirmed by biomarkers
 - ◆ Longstanding IBD in surveillance for CRC

Dont's

- Don't perform non-urgent endoscopy
- Don't use surgical masks in the endoscopy room

Endoscopy is extremely important in the management of IBD. In the context of COVID-19, viral SARS-CoV-2 RNA was found in the stool of a significant percentage of infected patients and, in some cases, the test was still positive after nasal swabs became negative. Even though the presence of the virus in gut mucosa does not seem to have a direct impact in gastrointestinal symptoms, it is not known if viral shedding is associated with fecal-oral transmission. The SARS-CoV-2 infection rate among endoscopy personnel is significantly lower (4.3%) than the average infection rate reported for health care workers (about 10%).⁸³ However, it is higher than the values of the pre-COVID-19 era, when it ranged from 1.1 (in colonoscopies) to 3.0 (in esophagogastroduodenoscopy), for every 1000 procedures.⁸⁴ Thus, risk/benefit scales should be carefully weighted, ensuring that endoscopic procedures are only performed in patients who urgently require endoscopy.⁷⁹ A first approach is to stratify patients according to the risk of SARS-CoV-2 infection: i) patients with fever, fatigue or respiratory symptoms, myalgia, diarrhea, loss of sense of smell or taste, close contact with COVID-19 patient, or travelling to outbreak areas are considered to have high risk¹²; ii) the remaining patients are considered as intermediate risk patients.

Endoscopy should be performed in a negative-pressure room, if possible, in accordance with the guidelines for infection control in endoscopy.^{85,86} In addition, the staff of the endoscopy department should follow standardized precautions.¹² SARS-CoV-2 is easily inactivated by many common disinfectants and no additional approach should be implemented to clean and disinfect the endoscope.⁸⁷ Even so, due to the risk of aerosolization, the room should be kept empty for at least for 1 hour before the next procedure in the absence of negative pressure, and for 30 minutes in the case of a negative-pressure room.⁸⁸ The basic protection requirements of the medical staff, in the endoscopy centre, should reach Biosafety level 2 (wearing of disposable gowns, N95

masks, goggles, caps, and shoe covers during endoscopy) in all GI endoscopic procedures. Biosafety level 3 protection (PPE, respirators, plus negative-pressure rooms due to the aerosolization risk of SARS-CoV-2 virus during endoscopy) is required for all endoscopic procedures in SARS-CoV-2 infected or suspected patients, and in those with very high risk of potential exposure.

During the pandemic outbreak, non-urgent endoscopic procedures should be postponed. These would include all the procedures that are not urgently required, and whose benefits do not outweigh the risk of SARS-CoV-2 infection. In this category, laucci et al⁷⁹ included new diagnosis of IBD with moderate/severe activity, severe acute flares of ulcerative colitis, refractory medical obstruction manageable through endoscopy or jaundice in patients with IBD, and primary sclerosing cholangitis with a dominant stricture. All the patients with scheduled endoscopy should be contacted, by phone, the week before, and again 1-2 days before the procedure, to identify patients with suspected COVID-19 or at risk of being infected. At the hospital, one patient checkpoint is mandatory, at every public entrance or endoscopy unit, and the endoscopist and/or the nurse must be double-checked. Urgent scenarios could be prioritize: i) in priority cases, endoscopy must be performed within 3 months: mild-to-moderate flare-ups, patients with longstanding IBD in surveillance for colorectal cancer with dysplasia, endoscopic resection in patients with low-grade or high-grade dysplastic colonic lesions, symptomatic patients with moderate pouchitis and altered blood test); and ii) in all the other cases, endoscopy can be delayed until 6 months after the infection rates decrease (mild pouchitis, IBD patients with flare-up not confirmed by biomarkers, longstanding IBD in surveillance for CRC).⁷⁹ Some conditions may require more prompt evaluation than others, depending on results of blood tests, non-invasive inflammatory markers, and previous history of dysplasia or cancer.

10-Can we still perform IBD surgery?

Do's

- Do emergency surgery in life-threatening situations
 - ◆ free bowel perforation
 - ◆ closed loop obstruction
 - ◆ acute severe colitis refractory to medical treatment
 - ◆ active bleeding not amenable to interventional therapy
- Do non-elective surgery as soon as possible
 - ◆ invasive colorectal cancer, high-grade dysplasia
 - ◆ intractable stenosis failing medical management
 - ◆ penetrating luminal disease resistant to medical therapy
 - ◆ perianal abscesses
- Do postpone surgery in uncomplicated IBD
- Do test all IBD patients with SARS-CoV-2-PCR before any surgery
- Do use N95 respirators as a minimal requirement for surgery

Don'ts

- Don't perform non-urgent IBD surgery

Emergency surgery is required in life-threatening situations, such as bowel perforation, closed loop, or medically refractory acute severe colitis. Conversely, non-urgent surgery should be postponed to protect patients and healthcare workers, and facilitate the management of healthcare resources, in times of scarcity of staff and material.^{89–91}

Non-elective surgery is indicated for invasive colorectal cancer/high-grade dysplasia, intractable stenosis, and abdominal/perianal abscesses not amenable to medical/interventional management.⁹² Perioperative complexity should be minimized, including liberal use of terminal ostomy instead of a high-risk anastomosis, and drainage procedures, with or without antibiotics, instead of complex surgery.⁹² Keeping track of deferred surgeries is the key to provide responsible surgical care after the pandemic.

Owing to the paucity of symptoms in the majority of COVID-19-positive patients and to the surgical morbidity reported during viral incubation (mortality 20.5%, ICU requirement

44.1%), it is mandatory to test all patients for SARS-CoV-2 before surgery.^{89,93,94} When rapid PCR tests are unavailable, chest CT can be the diagnostic. If neither testing nor imaging is available, all patients must be considered positive.⁹²

At surgery, the risk of aerosolization is high and contamination with body fluids is a concern.⁹³ To date, infectious virus was found in airway, blood, and feces.⁶¹ Aerosol formation is potentiated by tissue dissection with energy devices, while concentration of abdominal aerosol is inherent to laparoscopy.^{95–97} Consequent use of PPE is mandatory, including face shield/goggles, N95 mask, and proper donning and doffing.⁹⁵ The number of staff should be minimized and negative pressure operating rooms should be preferred. Scientific societies have issued conflicting statements regarding the use of laparoscopy, with no firm contraindication in the COVID-19 era. Avoidance of laparoscopy is thus debatable, as open surgery will translate in additional morbidity and resource utilization. Low energy setting and pneumoperitoneum pressure, avoidance of 2-way insufflators, closed circulation with efficient filtration of pneumoperitoneum, and gentle desufflation are advised.⁹⁸

Conclusions

In conclusion, the existing evidence shows that IBD is not a risk factor for COVID-19. However, medical treatments should be re-evaluated in SARS-CoV-2 positive IBD patients and corticosteroid therapy should be re-evaluated regardless of symptoms. A goal should be to treat active disease and maintain remission, while adopting the same protective measures as the general population. In addition, non-urgent surgeries and endoscopic procedures should be postponed.

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