Review Article

The ‘black fungus’ Co-Infection in COVID-19 Patients : A Review

Jessica Novia¹, Friska Wilda², Alius Cahyadi³*, Marcella Adisuhanto³

¹ Marianum Halilulik Hospital, Belu, East Nusa Tenggara
² Panti Wilasa Citarum Hospital, Semarang
³ Department of Internal Medicine, School of Medicine and Health Sciences, Atma Jaya Catholic University of Indonesia, Jakarta

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ABSTRACT

Mucormycosis is one type of fungal disease, associated with a poor prognosis if not promptly diagnosed and managed because its highly aggressive tendency. Although it is a rare disease, a rapid increase in cases of mucormycosis associated with COVID-19 is being reported. Mostly, risk factors for this disease are uncontrolled diabetes mellitus, other immunosuppressive conditions and corticosteroid therapy. Immune dysfunction, lung pathology and corticosteroid therapy in COVID-19 patients making it more susceptible to develop fungal infection including mucormycosis. The combination of steroid therapy and underlying diabetes mellitus in COVID-19 also can augment immunosuppression and hyperglycemia. Control of hyperglycemia, early treatment with liposomal amphotericin B, and surgery are three important factors in mucormycosis therapy that essential for successful management. However, in this COVID-19 pandemic situation, that management strategies are compromised. First, hyperglicemia can be aggravated by glucocorticoid, therapy that used widely for COVID-19 especially in severe case. Second, patients with ARDS and multiorgan dysfunction can prevent timely diagnostic for imaging and other testing, so appropriate therapy that should be given will be delayed. Last, the essential service in hospital such surgery in this pandemic era reduced significantly to prevent the spread of COVID-19. This review was created with the aim mucormycosis co-infection can be considered in patients with COVID-19, especially with known risk factor. Prompt and rapid diagnosis are important for effective therapy and decreasing case fatality rate. The use of steroid in mild cases, utilization of higher doses of steroid and drugs that targeting immune pathway should be avoided.

Keywords: Mucormycosis; Black Fungus; Coronavirus; COVID-19

ABSTRAK

Mucormycosis merupakan salah satu penyakit infeksi jamur dengan tingkat penularan yang tinggi. Jika tidak segera didiagnosis dan diterapi, maka berhubungan dengan prognosis yang buruk. Walaupun penyakit ini jarang ditemukan, tetapi data penelitian terbaru melaporkan peningkatan signifikan kejadian mucormycosis pada pasien COVID-19. Umumnya, penyakit diabetes melitus yang tidak terkontrol, kondisi imunosupresif lain dan terapi kortikosteroid merupakan faktor risiko terjadinya mucormycosis. Disfungsi sistem imun, kelainan patologis paru dan terapi kortikosteroid pada pasien COVID-19 membuat pasien lebih berisiko untuk mengalami infeksi sekunder termasuk mucormycosis. Kombinasi terapi steroid dan adanya komorbid diabetes melitus pada COVID-19 juga lebih meningkatkan kondisi imunosupresi dan hiperglikemia. Kontrol hiperglikemia, pengobatan awal dengan liposomal amfoterisin B, dan pembahana adalah tiga aspek penting dalam terapi mucormycosis yang merupakan faktor penentu keberhasilan penatalaksanaannya. Walaupun demikian, dalam situasi pandemi COVID-19 ini, strategi penatalaksanaan tersebut sulit tercapai. Pertama, kondisi hiperglikemia dapat diperburuk dengan glukokortikoid, yang merupakan terapi yang digunakan secara luas untuk COVID-19 terutama pada kasus berat. Kedua, pasien dengan ARDS dan disfungsi multiorgan dapat membuat uji diagnosis seperti pencitraan dan tes lainnya menjadi terlambat dilakukan sehingga diagnosis dan terapi pasien akan tertunda. Terakhir, di era pandemi
In this review, we would like to summarize recent data concerning mucormycosis co-infection in COVID-19 patients, epidemiology, pathogenesis and treatments. Mucormycosis disease progression is rapid and have angioinvasive nature, so a prompt diagnosis and treatment should be started as soon as possible to reduce the mortality.

MUCORMYCOSIS

Mucormycosis, formerly known as zygomycosis is a fungal disease caused by a group of molds called mucormycetes. These diseases are most often caused by a fungus that is found in soil and decaying vegetation, usually inhaled by humans from the air. There are various ways a person can contract mucormycosis such as by spores inhalation, food containing spores consumption, and spores-contaminated wound. This infection is mostly attacking immunocompromised individuals or taking medicines that weakened their immune system. The most common fungal species that result in mucormycosis are the Rhizopus species and Mucor species.

DISCUSSION

Epidemiology

Globally, mucormycosis prevalence around 10,000 cases in the world except India and after merging with India to become 910,000 cases globally. Mucormycosis found in tropical and subtropical climates, such as Indonesia. Indonesia is a tropical country, warm and
humid, with numerous environmental fungi. Unfortunately, the prevalence in some developing countries, including Indonesia, are still unclear because the cases remain undiagnosed due to difficulty in collecting tissue samples and limited facilities of mycology laboratories. The etiologic agents mostly are *Rhizopus* spp., *Mucor* spp., and *Lichtheimia* (formerly *Absidia* and *Mycocladus*) spp. Genera of other mucorales such as *Rhizomucor*, *Saksenaea*, *Cunninghamella*, and *Apophysomyces* are less common.

**Clinical Manifestation of Mucormycosis**

Mucormycosis have six major clinical form based on clinical manifestation and anatomic position of the invasion including rhinocerebral, pulmonary, cutaneous, gastrointestinal, disseminated and unusual form such as endocarditis, osteomyelitis, peritonitis and renal infection. The initial symptoms of mucormycosis are non-specific. The most common form is rhinocerebral mucormycosis. Presentation usually begins with pain and numbness in the eyes and face, followed by conjunctival suffusion and blurred vision. Fever does not occur in almost 50% of cases. Mostly, leukocytosis may occur. If it is not properly dealt, it could spread to the ethmoid sinus into the orbit caused damage to the function of extraocular muscle and proptosis with chemosis. In initial phase of the infected area appears normal and the concomitant progression of the disease becomes erythema with or without edema, then appears purplish and lastly formed eschar blackish necrotic tissue (Figure 1). Infection may also extend to the mouth and cause the formation of a necrotic ulcer on the palate. This finding indicates that the disease has spread.

Pulmonary mucormycosis patients usually present with high-grade fever (>38°C) and non-productive cough. Less common symptoms such as pleuritic chest pain and dyspnea. In rare circumstances, can present in endobronchial tree and causing airway obstruction. Cutaneous mucormycosis can classified as localized if affect skin and subcutaneous tissue, or deep extension if invades deeper to muscle, tendon or bone. Typical presentation is necrotic eschar with erythema and induration in surrounding skin. Gastrointestinal mucormycosis is less common type and hard to diagnose in living patients. Most affected organ is stomach followed by colon and ileum. The presentation usually nonspecific such as neutropenic fever and hematochezia. If severe, this disease can invade blood vessels in bowel and resulting in perforation, peritonitis, hemorrhage and sepsis. Disseminated mucormycosis occur when spreading hematogenously to other organs. Commonly, site of spread is brain, but also can found in liver, spleen, heart and other organ. The presentation may vary according to location and degree of tissue invasion in that affected organ.
Diagnosis

The diagnostic pathway was designed by the European Confederation of Medical Mycology and the Mycoses Study Group Education and Research Consortium (ECMM/MSGERC) consensus. The ability in diagnosing mucormycosis basically depends on the well-trained staffs, techniques and imaging types, and mycological and histological investigations. A prompt referral to the highest care level was recommended for patients with suspected mucormycosis.13

Diagnostic for mucormycosis such as radiologic features is nonspecific and have wide range of types. The most common features that can be identified for pulmonary mucormycosis are presence of nodules, consolidations, reverse halo sign, large perilesional halo (Figure 2) and cavitation. Reverse halo sign characterized by peripheral consolidation with central ground glass and large perilesional halo characterized by ground-glass halo around lesion that very extensive and bigger than the lesion itself.17,18

In rhinocerebral mucormycosis, sinus involvement usually occurs and must be identified in radiologic findings. The most common paranasal sinus involved are maxillary, ethmoid and sphenoid. Mucosal thickening and bone erosion in imaging also the common features. Signal characteristics and contrast enhancement can be seen in CT scan, Most common form is mild enhancement. Less common, non-enhancing and heterogenous pattern can also be found. If the imaging present with non-enhancing opacification of sinuses, presence of retro antral, facial and orbital fat stranding and hypodense soft tissue extension indicated aggressive infection. (Figure 3). Lastly, imaging must identify extra sinus extension such as orbit and face.19

Histopathological examination for mucormycosis is important but not always reliable to differentiate with Aspergillus. Mucorales have primitive coenocytic hyphae which are fragile because of lack of regular hyphae-septations. They make aggressive tissue grinding can render fragile fungal elements become non-viable. The important differentiation between Mucorales and Aspergillus is on their hypha type. Mucorales hypha have wide diameter and non-septate while Aspergillus hypha is narrower and have many sepatation.20,21

Imaging

Fig. 2 (a) Frontal radiograph of the right lung shows a faint area of ground-glass opacity (dotted circle); (b) Coronal CT image obtained an area of nodular ground-glass opacity; (c) Coronal CT image shows enlargement of the lesion with development of the reverse halo sign.22
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**Fig. 3** (a) Coronal view of CT showing involvement of left maxillary sinus, nasal conchae, and ethmoidal sinus extending up to frontal sinus; (b) Axial view of CT showing destruction of posterior, medial, and anterior walls of left maxillary sinus.\(^{16}\)

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**Histopathology in Mucormycosis**

**Fig. 4.** Structure of *Mucor*. (a) Mucorales are irregular hyphae with wide width (6 to 25-micron diameter) are non-septate or sparsely septate, ribbon-like; (b) High-power photomicrograph shows a spherical structure called the sporangium. (Lactophenol cotton blue stain).\(^ {22,23}\)

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**Culture and Microscopy**

Culture is highly recommended for identification of fungal genus and species.\(^ {7}\) It should be noted that culture does not always work for several reasons, including improper sampling and incorrect sample treatment before the examination. In fact, only 15-25% of cases are positive.\(^ {24}\)

**Treatment**

A multimodal approach is needed in the management of mucormycosis, such as discontinuation of risk factors, early administration of antifungal therapy with optimal doses, and surgical intervention. Treatment should be started immediately if the diagnosis is suspected because the disease tends to spread rapidly throughout the
body, although the exact diagnosis has not been confirmed.13

**Prophylaxis**

Posaconazole delayed-release tablets are recommended for neutropenic patients or those with graft versus host disease.7

**First-line antifungal monotherapy**

Daily doses of liposomal amphotericin B ranged 5-10 mg/kg for any patients and ≤5 mg/kg if renal toxicity develops. In Central Nervous System (CNS) involvement, use of amphotericin B lipid complex 5 mg/kg per day. Treatment duration given usually weeks to months, depending on each patients condition. If the immune defect is resolved, such as well-controlled diabetes and resolved neutropenia, immunosuppressant can be tapered or stopped, therapy can be continued until resolution of signs and symptoms of infection, and substantial radiographical improvement.7 At the fourth week, the overall response rate was 36%, while in the twelfth week, the overall response was 45%,13

**First-line antifungal combination monotherapy**

There are no definitive data to guide the use of antifungal combination therapy.7

**Antifungal salvage treatment**

Daily Isavuconazole 200mg (after six doses of 200 mg q8h) and Posaconazole delayed-release tablets at a dose of 200 mg q6h or infusions are strongly supported as salvage treatment.7,13

**Surgery**

Aggressive surgery is often required not only on necrotic tissue but also on surrounding tissue that appears healthy because the Mucorales grow so rapidly.7

**THE LINK BETWEEN MUCORMYCOSIS CO-INFECTION COVID-19**

As previously stated that mucormycosis is mainly attacking immunocompromised patients, although possibly found in immunocompetent individuals.25,26 Generally, mucormycosis does not pose a serious threat to healthy individuals because immune system mainly polymorphonuclear cells can destroy the spores and hyphae.26–28 When patients are exposed to SARS-CoV-2, the virus will target the immune system. The relationship between COVID-19 and mucormycosis is the state of weakened of patients’ immune responses, with reduced numbers of T lymphocytes, CD4+, and CD8+ T cells and medical treatment with a steroid to reduce inflammation.29

One of the major risk factors that increasing morbidities and mortalities in COVID-19 associated with mucormycosis, is diabetes mellitus.30–32 In patients with diabetes, Rhizopus is the most commonly found fungus. The reason which allows them to survive in high acid and glucose are an enzyme properties and ketone reductase.33 Treatment pathway for patient with COVID-19 with mucormycosis co-infection including both diseases therapy. Therapy requires surgical debridement, antifungal treatment and stabilization of risk factor.8

Diabetic ketoacidosis (DKA) often occurs in severe infections, such as in COVID-19. Therefore, it is not surprising that patients with COVID-19 are more likely to develop mucormycosis because acidic conditions make mucorales species easier to grow.34 Research suggests SARS-CoV-2 induces damage of pancreatic islets resulting in acute diabetes and DKA.35 Another explanation for why the diabetogenic state occurs in patients with severe COVID-19 is due to cytokine storms that increase insulin resistance and high expression of angiotensin-converting enzyme 2 receptors in pancreatic islets. Increased serum ferritin levels in severe COVID-19 also one of the possible roles of blood acidosis for mucormycosis susceptibility.34,36–38

It has been proven that by administration of systemic corticosteroids could cut down death rates in COVID-19 patients on invasive mechanical ventilation.39,40 According to the European Organization for Research and Treatment of Cancer and the Mycoses Study Group Education and Research Consortium
The resultant tissue damage leads to the release of free iron into the circulation, which further exacerbates the mucormycosis process.\textsuperscript{51}

“Endothelialitis” in patients with severe COVID-19 is also one of the associations between COVID-19 and mucormycosis.\textsuperscript{52,53} Important initial steps of mucormycosis are endothelial adhesion and penetration.\textsuperscript{7} In addition, acidemic states, and hyperglycemia induce the endothelial receptor glucose-regulated protein (GRP 78) and the mucorales adhesin spore coat protein homologs (CoTH), creating a “perfect storm” for increased adhesion and penetration of mucorales to the endothelium.\textsuperscript{54}

Based on the available literature regarding mucormycosis co-infection COVID-19, there were six studies reporting 28 patients that have reported rhino-orbito-cerebral mucormycosis. It is important to remember that mucormycosis can occur at any time after a COVID-19 infection, either during the hospital stay, or a few days to weeks after discharge. Therefore, all physicians

\textbf{Fig. 5 Pathogenesis of Mucormycosis}\textsuperscript{44,45}
could be more aware of these side effects of the kinds of treatment patients are given and how could patients be more aware of what they could face because of the medicine that they are taking, especially if having underlying conditions. They should knowledgeable about the red flag symptoms of invasive mucormycosis.11,29,55–58

Alekseyev, et al presented a 41-year-old man with a history of type 1 diabetes mellitus (T1DM), COVID-19 pneumonia and rhinocerebral mucormycosis. He was treated with steroids and hydroxychloroquine before, as the recommended regional COVID-19 practice guideline at the time. For his diabetic ketoacidosis (DKA) treated with intravenous fluids and an intravenous insulin, cefepime and amphotericin B IV, along with three surgical debridements for the rhinocerebral mucormycosis. The patient successfully discharged and continued the treatment at home. 59

Another study by Kanwar et al, they presented a 56-year-old man with COVID-19 and underlying end-stage renal disease. This patients also developed mucormycosis as a complication of COVID-19. He received a five-day therapy of methylprednisone, one dose of tocilizumab, and one unit of convalescent plasma. At first hospital admission, blood cultures collected were negative for bacterial and fungal organisms. He was discharged home seven days later but five days later he was readmitted because shortness of breath. Polymerase chain reaction (PCR) examination for COVID-19 was positive again and chest radiograph showed increasing density and pleural effusion. He was started on empiric intravenous (IV) vancomycin and piperacillin-tazobactam. Sputum sample was collected and showed filamentous fungal elements on fungal stain that was suspected from Mucorales group because non-septate hyphae. Empiric amphotericin B was started and antibacterial medications were discontinued, unfortunately the patient developed cardiac arrest and died the following day. 60

Maini et al, reported a 38-year-old man with COVID-19 confirmed, no history of diabetes or other condition. He was monitored in ICU and started on remdesivir IV, methylprednisolone IV and dexamethasone. After 12 days of treatment, the glycated hemoglobin (HbA1C) level was 12.3%. Eighteen days later, the patient complaint of swelling and pain in his left eye, then underwent MRI scan and histopathologic examination from sinus sample. Patient was then diagnosed as sino-orbital mucormycosis. Medical treatment was changed into amphotericin B and patient was going into surgical debridement. After a total of 38 days of hospitalization, he was discharged and continued treatment at home. 61

CONCLUSION

In COVID-19, due to immune system dysregulation, diabetogenic state, endothelialitis, and the widespread use of steroids as therapy against COVID-19 may lead to the development/exacerbation of pre-existing fungal diseases. Physicians should be aware of the development/exacerbation of pre-existing fungal infection among COVID-19 patients, especially if rhino-orbital-cerebral presentations are noted. A multidisciplinary approach should include the recognition of host factors, assessment of clinical manifestations, use of appropriate imaging modalities, histology and microbiology with any appropriate surgical consultation and treatment. The use of steroids should be monitored to achieve a therapeutic effect at the lowest dose and shortest durations to lower the risk of development/exacerbation of pre-existing fungal infection.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.
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