Mucormycosis in a patient with acute myeloblastic leukemia following liver transplantation for Wilson’s disease

Aleksandra Anna Łanocha1,A,C-D,E, Renata Guzicka-Kazimierczak1,B,C,E, Barbara Zdziarska1,B,C,E, Marta Wawrzynowicz-Syczewska1,B,C,E

1 Pomeranian Medical University, Szczecin, Poland

A – Research concept and design, B – Collection and/or assembly of data, C – Data analysis and interpretation, D – Writing the article, E – Critical revision of the article, F – Final approval of article


Abstract

A case is presented of mucormycosis in a patient with acute myeloblastic leukemia following liver transplantation for Wilson’s disease. A 58-year-old female was admitted to the Department of Haematology with deterioration of her general condition, loss of appetite, tiredness and difficulty with mental contact for a few days. Blood and urine cultures for bacteria and fungus, galactomannan antigen were negative. Whole body computed tomography demonstrated bilateral hilar lymphadenopathy with necrotic lesions: splenomegaly with a hypodensive lesion 13 × 20 × 19 mm and lower pulmonary infiltrates suggested fungal etiology. Magnetic resonance imaging of the brain showed thickened meninges. Finally, mucormycosis was diagnosed. Treatment with amphotericin B lipid complex was started, resulting in an partial improvement of the general condition and decreased level of inflammatory markers. However, the patient’s condition continued to deteriorate, with sepsis etiology Escherichia coli, and despite the intensive managements she eventually died.

Key words

Mucormycosis, acute myeloblastic leukemia, Willson’s disease, liver transplantation

INTRODUCTION

Mucormycosis (zygomycosis) is a rare but still fatal opportunistic invasive fungal infection caused by filamentous fungi from the orders of Morales and Entomophthorales [1]. The incidence of invasive fungal infections, (IFI) has increased in recent years as a result of severe and long-lasting immunosuppression and neutropenia [2]. After aspergillosis and candidiasis, mucormycosis is the third most common cause of IFI in patients with haematologic malignancies after chemotherapy and solid organ transplantation [3]. Mechanisms of disease are related to conspicuous deficient action of polymorphonuclear and mononuclear phagocytes [4]. Data shows an increasing incidence of mucormycosis from 0.7/million in 1997 to 1.2/million in 2006, and in patients with haematologic malignancies from 0.02/million to 0.2/million [5]. Sarvestani et al. [6] observed that the prevalence rate of mucormycosis infection was generally 4.27 per 100 patients with haematologic malignancies. The clinical and radiological manifestations of mucormycosis are unspecific. Diagnosis and routine investigation includes also microbiological, histological and cytological findings [7].

Wilson’s disease (WD) is an inherited disorder of copper metabolism with a wide spectrum of clinical symptoms, mainly hepatic or neurological. Liver transplantation may correct the underlying hepatic metabolic defect and is indicated for the <5% of WD patients with acute failure as the first presentation of disease [8]. A case is presented of mucormycosis in a patient with acute myeloblastic leukemia (AML) following liver transplantation for WD.

In 2001, a 58-year-old female was diagnosed with WD (hepatic neurological type) and in 2012 underwent liver transplantation. In April 2015, after reconstruction of the arterial anastomosis of the transplanted liver due to hepatic artery stenosis and cholecystectomy, the patient was diagnosed with acute myeloblastic leukemia FLT3(-), NPM1(-), CEBPA(-). She was treated with chemotherapy according to the Polish Adults Leukemia Group (PALG): induction of DAC (daunorubicine, cytarabine and cladribine) <60-year-old protocol, followed by reinduction chemotherapy CLAG-M (cladribine, cytarabine, mitoxantron and GCS-F). Antifungal prophylaxis with posaconazole was used. On day 30, after the 2nd cycle of chemotherapy, pancytopenia was present and aplastic anaemia was diagnosed in bone marrow. Oral administration of a low dose of methylprednisolone was therefore started. The patient was admitted to the Department of Haematology with deterioration of her general condition, loss of appetite, tiredness and difficulty with mental contact for a few days. On physical examination, the patient was unconscious, dehydrated, with petechiae, haematoma on the dorsum of the nose and tachycardia. A blood test revealed pancytopenia with a white blood cell count of 0.44 G/l, platelet count – 1.0 G/l and agranulocytosis – 0.06 G/l, an increased level of inflammatory markers (CRP-115.7mg/l; prokalcytonine2,74ng/ml), hyperglycaemia 483mg/dl, ketoacidosis, increased aminotransferase (ALT: 187 U/l, AST: 305 U/l), increased GGTP: 98 U/l and lactic dehydrogenase- 960 U/l, normal total bilirubin 0.40 mg/dl, creatinine 1.00 mg/dl.

In the body computed tomography of the head, thickening of the mucosa of the maxillary sinus (greater on the right), ethmoid cells, frontal and sphenoid sinus were described.
Blood and urine cultures for bacteria and fungus, and galactomannan antigen were negative. The patient's diabetic state was controlled with insulin and wide-spectrum antibacterial therapy (ceftriaxone, amikacin) was initiated. On the days 2 and 3 of the hospital stay, the patient developed a fever; meropenem, vancomycin, caspofungine were started. Aspergillosis was suspected and a whole body computed tomography was therefore performed and treatment with voriconazole administered. The whole body CT demonstrated bilateral hilar lymphadenopathy with necrotic lesions: left side – 29 × 23 × 25 mm and right side – 21 × 14 × 13 mm, multiple nodules, parenchymal density and pleural effusions, splenomegaly with a hypodense lesion measuring 13 × 20 × 19 mm, and lower pulmonary infiltrates, suggested fungal etiology (Fig. 1 and 2). Magnetic resonance imaging of the brain showed thickened meninges (Fig. 3). Cultures from cerebrospinal fluid were negative; cerebrospinal fluid analysis excluded myeloblasts.

During hospitalization, necrotic eschar was present on the dorsum of the nose. Scrapings for microbiological tests were therefore taken which showed Rhizomucor. Control blood and urine cultures were positive for _Candida glabrata_ in the urine. In the treatment, voriconasole was changed to amphotericin B lipid complex (on day 10 of hospitalization), resulting in a partial improvement in the patient’s general condition, and a decreased level of inflammatory markers. However, her condition continued to deteriorate, with sepsis etiology _Escherichia coli_, and despite the intensive managements she eventually died.

DISCUSSION

Mucormycosis is an uncommon and life-threatening, fungal infection caused by _Mucorales_ spp. [3]. Risk factors are diabetes, acidosis, steroid use, haematologic malignancy, transplantation, severe trauma, burns, neutropenia and renal failure [9, 10]. Mucormycosis can be classified as one of six forms: rhinocerebral, pulmonary, cutaneous, gastrointestinal, disseminated and uncommon presentations [11]. Patients with leukaemia have traditionally been considered to represent the majority of cases with pulmonary mucormycosis [12]. In the pulmonary or sinus form, exposure occurs by inhaling aerosolized mucor spores from the environment. Gastrointestinal mucormycosis, resulting from ingestion of spore-contaminated food, and cutaneous mucormycosis (very rare cases) due to direct skin implantation of fungal spores [2]. The patient in the presented case had the mentioned key predisposing factors: diabetes, neutropenia, acute myeloblastic leukaemia, chemotherapy, immunosupression after liver transplantation, treated with steroids. The clinical...
signs and symptoms of mucormycosis are nonspecific. There are no biomarkers to identify disease. β-D-glucan test and Aspergillus galactomannan tests do not detect antigen of the Mucorales cell wall [13]. Identification of mucormycosis is very difficult, most often comes from microscopic, culture and/or histopathological examination of relevant samples. To diagnose mucormycosis and assess disease staging, extensive clinical examination for signs dissemination, CT of brain, sinuses and chest, bronchoscopy with bronchoalveolar lavage and biopsy suspicious lesions of hard palate, skin, sinuses necessary [13]. New diagnostic approaches contain molecular testing and various other clinical samples including formalin-fixed tissues (MALDI-TOF) and Mucorales-specific T-cell detection [14, 15]. Unfortunately, molecular procedures are not widely available and diagnosis is based on fungal isolation and/or direct examination in clinical samples. Management includes association between antifungal, surgery and control of diabetes, metabolic abnormalities, such as haemochromatosis, haematopoietic growth factor of neutropenia, discontinuation/tapering of steroids and reduction of immunosuppressive therapy. Small focal lesions of mucormycosis should be surgically resected. Amphotericin B, posaconazole, isavuconazole are the most potent agents. Echinocandins in association with L-AmB; L-AmB associated with posaconazole can be used in salvage therapy [13, 14]. Amphotericin B is considered the drug of choice for primary treatment of mucormycosis. The standard daily dose of liposomal amphotericin B and lipid complex amphotericin B is 5 mg/kg/day. Although dosages beyond 5mg/kg/day may be considered, for example when there is central nervous system or oseoarticular involvement [10]. Posaconazole may be considered as salvage treatment of mucormycosis or as a first-line treatment only in cases when amphotericin B treatment is absolutely contraindicated. Isavuconazole is a new broad-spectrum triazole, used for the treatment of mucormycosis when amphotericin B is not feasible. In the presented case, aspergillosis was initially suspected, therefore voriconazole was administered. On day 10 of hospitalization, voriconazole was changed to amphotericin B lipid complex, resulting in an partial improvement of the general condition and decreased level of inflammatory markers.

Mucormycosis still remains a fatal prognosis, with high mortality rates, ranking from 20–50% if localised, up to 70–90% in cases of disseminated disease [10, 16]. To the best of the authors’ knowledge, this is the first described case of mucormycosis in a patient with WD after liver transplantation and acute myeloblastic leukaemia.

The presented case illustrates the dramatic course of mucormycosis and confirms the fatal prognosis associated with high mortality. The lack of availability of fast and reliable diagnostic tests means that it still remains a problem. The management of mucormycosis, including antifungal therapy, surgery and control of underlying conditions, should be started as soon as possible.

Acknowledgements
The authors would like to thank Dr Ewa Gabrysz-Trybek for assistance with preparation of the Figures.

Conflicts of Interest
The authors declare no conflict of interest.

Ethical Approval
No approval was required.

Funding Source
The research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

REFERENCES

Generation of the DOI (Digital Object Identifier) – task financed under the agreement No. 618/P-DUN/2019 by the Minister of Science and Higher Education.