Successful Management of Mixed Mycosis in HIV-Negative Patients With Different Immune Status: A Case Series Report

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**Objective:** The limited information available on mixed mycosis involving the lungs makes the understanding of mixed fungal diseases insufficient and affects prognosis. Our study aims to improve understanding by exploring experience in the successful management of mixed fungal infections.

**Methods:** Patients who had two types of mycosis involving the lung at the same disease course were retrospectively enrolled.

**Results:** Between September 2011 and December 2019, 17 patients with proven mixed mycosis were enrolled. Four patients were immunocompromised, with one case each of lung transplantation, corticosteroid treatment, STAT3 hyper-IgE syndrome, and anti-IFN-γ autoantibody-associated immunodeficiency syndrome. Among 13 patients who were not immunocompromised, 9 had type 2 diabetes mellitus. Eight cases were coinfection with Mucor and Aspergillus, 4 cases were Cryptococcus and Aspergillus, 2 cases were Talaromyces marneffei and Cryptococcus, 2 cases were Talaromyces marneffei and Aspergillus, and 1 case was Candida and Aspergillus. Seven patients were diagnosed with mixed pulmonary mycosis at almost the same time. Among the remaining 10 patients, the initial treatment was ineffective in four cases, and six patients showed a partial response to the initial antifungal treatment, but the original fungal lesions became re-enlarged. Three patients were admitted to the intensive care unit during hospitalization, and one patient died. Another Mucor coinfection patient died due to treatment refusal.

**Conclusion:** Mixed mycosis involving the lungs is not uncommon in patients without apparent immune deficiency diseases. During the management of mycosis, we recommend keeping mixed mycosis in mind for patients with a poor response to initial antifungal treatment, even in immunocompetent populations, and identifying the cause of illness through a rigorous procedure.

**Keywords:** immunocompetent, immunocompromised, Mucor, Aspergillus, Cryptococcus, Talaromyces marneffei
INTRODUCTION

Fungi can cause serious infections worldwide, with the number of serious cases reaching 150 million (Bongomin et al., 2017). Immunocompromised hosts, such as those with organ transplantation, hematopoietic stem cell transplantation, malignant tumors receiving radiotherapy and chemotherapy, and autoimmune disease receiving immunosuppressive therapy, are at risk of invasive pulmonary mycosis, which may lead to poor prognosis. On the basis of discharge diagnoses, the incidence of invasive fungal infections in France was 5.9/100,000 cases/year, with a mortality of 27.6%, both of which increased during the observation period (2001–2010) (von Lilienfeld-Toal et al., 2019). While, the proportion of pulmonary fungal infection increased annually from 26.5 per 1000 inpatients in 2013 to 42.6 in 2019 and from 3.07 per 1000 outpatients in 2013 to 8.48 in 2019 in Guangzhou, China (Li et al., 2021). The most common pulmonary mycoses are Aspergillus, Candida, Cryptococcus species, and Pneumocystis jirovecii (Bongomin et al., 2017; Azoulay et al., 2019; Zhan et al., 2021). Mixed fungal infections have also been reported (Wang et al., 2018; Awari et al., 2020); however, their characteristics have not yet been described in detail, and most are in immunocompromised hosts.

Existing data show that pulmonary fungal disease is not limited to immunosuppressed hosts. Reports of pulmonary fungal diseases in immunocompetent hosts are gradually increasing, such as cryptococcosis, Talaromyces marneffei infection, and aspergillosis, although the latter is one of the common infections in primary immunodeficiency diseases such as chronic granulomatous diseases (Mortaz et al., 2019; Abd Elaziz et al., 2020). In China, most patients with cryptococcosis do not have an immunocompromised status (Ye et al., 2012; Cheng et al., 2021). Talaromyces marneffei infection, an endemic fungal disease in Asian populations, also occurs in HIV-negative immunocompetent populations (Lee et al., 2019; Pan et al., 2020). Although some patients have been confirmed to have nonclassical immunodeficiency, such as hyper-IgE syndrome and anti-IFN-γ autoantibody-associated immunodeficiency syndrome, most patients do not have clear immunodeficiency or hypofunction (Lee et al., 2019; Pan et al., 2020). Due to the lack of specificity in the clinical manifestations of pulmonary mycosis in immunocompetent hosts, mycosis is often not the first consideration during clinical diagnosis and treatment. Incorrect diagnosis often delays treatment, which may affect the prognosis.

Diagnosis and treatment are even more difficult in mixed fungal disease because of unsatisfied positivity in culture and pathology. The need for timely inoculation, destruction of hyphae by tissue grinding, and empiric use of antifungal drugs is one of the reasons for negative culture. The morphological similarity of mold makes pathological diagnosis dependent on experienced technicians. Awari et al. and Wang et al. reported some cases of mixed mycoses of Aspergillus and Cryptococcus in both immunocompromised and immunocompetent patients (Wang et al., 2018; Awari et al., 2020). Coinfection with Talaromyces marneffei and Cryptococcus neoformans has also been reported in a non-HIV patient (He et al., 2020). However, the characteristics of fungal coinfection are still undefined. Here, we retrospectively analyzed a group of successfully managed mixed mycoses to explore the clinical characteristics and main points of diagnosis and treatment of mixed mycoses.

METHODS

This was a retrospective study conducted at the First Affiliated Hospital of Guangzhou Medical University. Between Jan 1, 2011 and Dec 31, 2019, patients who were hospitalized in the Department of Pulmonary and Critical Care Medicine with a diagnosis of mixed mycosis that involved pulmonary were searched from the electronic medical record database and enrolled as the study group. This study was approved by the ethics committee of the First Affiliated Hospital of Guangzhou Medical University (2018-119). Because this is a retrospective study, informed consent was not required.

Study Population
Inclusion criteria were the following: I. patients who had two types of fungal infections involving the lung at the same disease course; II. patients who had follow-up data in the outpatient medical records database; and III. patients who were HIV negative.

Mycosis was diagnosed based on the European Organization for Research and Treatment of Cancer and the Mycoses Study Group Education and Research Consortium (EORTC/MSGERC), which is a graded diagnostic standard, including the classifications of “proven,” “probable,” and “possible” invasive fungal disease (IFD) (Donnelly et al., 2019). The present study included only patients with proven or probable IFD. The diagnostic criteria for proven IFD included a specimen obtained by needle aspiration or biopsy showing the unique morphology of the fungus by histopathology, cytopathology or direct microscopic examination; a specimen obtained by a sterile procedure from a normally sterile and clinically or radiologically abnormal site consistent with an infectious disease process, recovering a hyaline or pigmented mold or yeast by culture; a positive blood culture for mold (Aspergillus was excluded) or yeast; or a positive test for cryptococcal antigen in cerebrospinal fluid or blood. The diagnosis of probable IFD should meet all three criteria, including host factors, clinical features and mycological evidence, with details in the referenced guideline (Donnelly et al., 2019).

Exclusion criteria included the following: I. two fungal infections did not occur at the same disease course, and the first mycosis was cured before the onset of the second mycosis; II. only one fungal infection was diagnosed; and III. there was incomplete information.

Data Collection
Clinical features of enrolled patients were collected, including medical history, underlying diseases, use of corticosteroids and/or immunosuppressants before the onset of mycosis, clinical symptoms and signs, imaging and laboratory examination data,
types of culture specimens, diagnosis procedure, treatment, and outcome. If the patient went to the outpatient clinic, the follow-up and results within 1 year were also collected from the database.

Statistics
SPSS 25.0 software was used for data processing of the results of this study. Because the treatment drugs and prognosis of mucormycosis are different from those of other fungal infections, comparisons were made between Mucor mixed infection and non-Mucor mixed infection cases. All P values are two-sided tests, and \( P < 0.05 \) means that the difference is statistically significant. Independent sample t-tests were used for continuous variables, and Fisher’s exact test and \( \chi^2 \) test were used for categorical variables.

RESULTS
General Characteristics
Between September 2011 and December 2019, 17 cases of mixed fungal infection were confirmed in our hospital. The average age was 50.2 ± 16.9 years, ranging from 21 to 76, with 12 (70.6%) males. Four patients were immunocompromised, with one patient receiving bilateral lung transplantation for destructive pneumocephalitis, one patient receiving long-term corticosteroid therapy for nephrotic syndrome and hypoalbuminemia, one case of STAT3 hyper-IgE syndrome and one case of anti-IFN-\( \gamma \) autoantibody-associated immunodeficiency syndrome (Table 1). Among the 13 patients who did not have apparent immune deficiency, 10 had underlying diseases, including hypertension, diabetes mellitus and hyperthyroidism (Table 1).

Among all 17 patients, 8 cases were coinfection with Mucor and Aspergillus, 4 cases with Cryptococcus and Aspergillus, 2 cases with Talaromyces marneffei and Cryptococcus, 2 cases with Talaromyces marneffei and Aspergillus, and 1 case with Candida and Aspergillus. The lesions of all patients with mucormycosis were confined to the lungs, but some patients with Talaromyces marneffei infection and cryptococcosis had involvement of extra-pulmonary organs, including bone, skin, lymph nodes, and/or central nervous system (CNS) (Table 1).

Clinical Symptoms, Signs and Laboratory Tests
Cough, expectoration and fever were the most common symptoms. More than 40% of patients had hemoptysis or shortness of breath. Only 3 patients (17.6%) had an elevated white blood cell count. There were no decreased immunoglobulin G/M/A levels or CD4+ or CD8+ T lymphocyte cell counts in patients without apparent immune deficiency diseases (Table 2).

Mycological Results and Biopsy
The culture was positive for Aspergillus fumigatus in case 6 and case 7 and for Talaromyces marneffei in case 11. A galactomannan antigen test was positive in case 14 and case 16. Biopsy was positive in all patients but case 16 (Table 1).

Manifestations of CT Images
All 17 patients underwent chest CT examination. The most common features of CT images were mediastinal lymph node enlargement (12 cases, 70.6%), cavity (10 cases, 58.5%), effusion or consolidation (9 cases, 52.9%), multiple nodules (7 cases, 41.2%) and pleural effusion (7 cases, 41.2%, Table 3 and Figure 1). However, none of the 17 cases showed classical features of lung mycosis, such as halo signs and crescent signs. There was also no vascular truncation sign or anti-halo sign. Lesions were less commonly located in the lingual lobe than in other lobes (Table 3).

Clinical Course, Treatment, and Prognosis
Seven patients were diagnosed with mixed pulmonary mycosis at almost the same disease course. Among the remaining 10 patients, the initial treatment was ineffective in three cases (cases 8, 10, and 12; fluconazole in two cases and voriconazole in 1 case), and the patients were diagnosed with mixed mycoses after review of the original biopsy from the hospital to which they were first admitted (Figure 2). They showed good response to adjusted treatment. The original fluconazole, second-line amphotericin B and voriconazole were ineffective in case 2. She was diagnosed with mixed mycosis after surgical resection. In six cases (cases 3, 5, 11, 13, 14, and 15), mixed mycosis diagnosis was made after bronchoscopy biopsy of re-enlarged original fungal lesions, which showed a partial response to initial antifungal treatment (fluconazole in 1 case, voriconazole in 1 case, itraconazole in 2 cases and posaconazole in 2 cases) (Figure 1). Among the 10 cases of mixed mycoses that were not diagnosed at the same time, initial antifungal drugs in 6 cases had no activity on the second fungus.

Three patients were admitted to the intensive care unit during hospitalization. However, patient 7 died even after treatment in the intensive care unit, and patient 6 died due to treatment refusal. Patient 2 showed a poor response to antifungal treatment, and pneumonectomy was conducted. Patients 5 and 11 suffered from a recurrence of disseminated Talaromyces marneffei infection even under the course of antifungal treatment. The remaining patient showed improvement in the CT scan with no recurrence.

Comparisons of Patients With Mucor Coinfection and Without Mucor Coinfection
Because voriconazole is ineffective for Mucor, which has a poor prognosis, comparisons were made between patients with Mucor and without Mucor coinfection to discriminate the characteristics between these two groups. There were 8 cases with Mucor coinfection and 9 cases without Mucor coinfection. Patients with Mucor coinfection seemed to be older than patients without Mucor coinfection; however, the difference was not significant (55 ± 16 years vs. 45 ± 16 years, \( P = 0.22 \)). Diabetes mellitus was more common in patients with Mucor coinfection.
| Case No. | Sex | Age | Smoking history | Comorbidity | Diagnosis of mycosis | Classification of IFD by EORTC/MSG | Host factors | Clinical features | Mycological evidence | Biopsy | Clinical course of illness | Outcome |
|----------|-----|-----|-----------------|-------------|---------------------|-----------------------------------|--------------|------------------|---------------------|---------|--------------------------|---------|
| 1        | Female | 67  | No              | Congenital lung cyst, hyperthyroidism | Chronic pulmonary aspergillosis and pulmonary cryptococcosis | Proven | NA | Lesions | Negative | Aspergillus and Cryptococcus | Diagnosis of mixed mycosis was made after surgery and itraconazole was prescribed | Survival |
| 2        | Female | 50  | No              | Type 2 diabetes mellitus, hypertension | Disseminated cryptococcosis (lung and lymph nodes) and pulmonary aspergillosis | Proven | NA | Consolidation | Negative | Aspergillus and Cryptococcus | Disseminated cryptococcosis (lung and lymph nodes) was diagnosed at first, with poor response to 5 months of treatment with fluconazole. Surgery was performed after intolerance to amphotericin B and a poor response to voriconazole for 1 month. Biopsy showed coinfection of Aspergillus and Cryptococcus. | Survival |
| 3        | Female | 66  | No              | Type 2 diabetes mellitus | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven | NA | Lesions and cavity | Negative | Aspergillus and Mucor | Voriconazole was prescribed after consideration of invasive fungal disease in the lung and poor status of the patient. The lesion showed a good response, but was re-enlarged 2 months after treatment. Biopsy via bronchoscopy confirmed the mixed mycosis. There was a good response after posaconazole treatment. | Survival |
| 4        | Male   | 57  | Yes             | Type 2 diabetes mellitus | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven | NA | Consolidation and cavity | Negative | Aspergillus and Mucor | Mixed infection was diagnosed concurrently, with a good response to amphotericin B liposome treatment. | Survival |
| 5        | Male   | 24  | No              | Hyper-IgE syndrome and bronchiectasis | Disseminated Talaromyces marneffei infection (lung, fungemia) and invasive pulmonary aspergillosis | Proven | STAT3 deficiency | Cavity | Negative | Aspergillus and Talaromyces marneffei | Disseminated Talaromyces marneffei infection was diagnosed at first and was treated with itraconazole because of intolerance to amphotericin B due to vomiting. Four months later, surgical resection was performed for new onset of hemoptysis and gradual enlargement of original lesions in chest CT scans. Mixed infection was diagnosed concurrently. But the patient was severely ill with poor condition and refused treatment. | Recurrence of Talaromyces marneffei infection during antifungal treatment |
| 6        | Female | 61  | No              | Type 2 diabetes mellitus | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven | NA | Consolidation and cavity | Aspergillus fumigatus by culture | Aspergillus and Mucor | Mixed infection was diagnosed concurrently. But the patient was severely ill with poor condition and refused treatment. | Death |
| 7        | Male   | 66  | Yes             | Lung transplantation, hypertension | Invasive pulmonary aspergillosis and pulmonary candidiasis | Proven | Lung transplantation | Consolidation | Aspergillus fumigatus by culture | Aspergillus and Candida | Invasive pulmonary aspergillosis was diagnosed after a positive result in sputum culture. The patient showed a poor response to voriconazole. Mixed mycosis was diagnosed after bronchoscopy, and amphotericin B was | Death |

(Continued)
| Case No. | Sex | Age | Smoking history | Comorbidity | Diagnosis of mycosis | Classification of IFD by EORTC/MSG | Host factors | Clinical features | Mycological evidence | Biopsy | Clinical course of illness | Outcome |
|---------|-----|-----|-----------------|-------------|---------------------|-----------------------------------|--------------|------------------|---------------------|--------|--------------------------|---------|
| 8       | Male | 46  | Yes             | Type 2 diabetes mellitus | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven | NA | Lesions, consolidation and cavity | Negative | Aspergillus and Mucor | given subsequently. However, there was a poor response. Pulmonary mycosis was considered after bronchoscopy biopsy at another hospital. There was no change in the lesion after 2 months of voriconazole. Mixed mycosis was diagnosed after review of the biopsy and showed good response to amphotericin B. | Survival |
| 9       | Male | 25  | Yes             | Type 2 diabetes mellitus | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven | NA | Lesions, consolidation and cavity | Negative | Aspergillus and Mucor | Mixed infection was diagnosed concurrently, with a good response to amphotericin B liposome treatment. | Survival |
| 10      | Male | 40  | No              | None | Pulmonary cryptococcosis and chronic pulmonary aspergillosis | Proven | NA | Consolidation and cavity | Negative | Cryptococcus and Aspergillus | Pulmonary cryptococcosis was confirmed at first, with a poor response to 8 months of treatment with fluconazole. Mixed mycosis was diagnosed after review of the biopsy and showed good response to voriconazole with therapeutic drug monitoring. | Survival |
| 11      | Male | 21  | No              | Anti-IFN-γ autoantibody-associated immunodeficiency syndrome | Disseminated Talaromyces marneffei infection (lung, lymph nodes, bone) and invasive pulmonary aspergillosis | Proven | IFN-γ antibody deficiency | Consolidation and cavity | Talaromyces marneffei | Serum galactomannan antigen > 1.0 | Disseminated Talaromyces marneffei was diagnosed at first and showed a good response to amphotericin B and itraconazole successively. There was new occurrence of lesions and re-enlargement of original lesions 9 months later, and mixed mycosis was diagnosed, with a good response to successive amphotericin B and voriconazole. | Recurrence of Talaromyces marneffei infection during antifungal treatment |
| 12      | Male | 41  | No              | Nephrotic syndrome and hypertension | Disseminated cryptococcosis (lung and CNS) and pulmonary Talaromyces marneffei infection | Proven | Corticosteroid treatment for nephrotic syndrome | Lesions and cavity | Talaromyces marneffei and Cryptococcus | Pulmonary cryptococcosis was confirmed at first, with a poor response to 1 month of treatment with fluconazole. Mixed infection was diagnosed after review of the biopsy, with a good response to successive amphotericin B liposome plus fluconazole and voriconazole. | Survival |
| 13      | Female | 76 | No              | Type 2 diabetes mellitus | Pulmonary mucormycosis and invasive pulmonary aspergillosis | Proven | NA | Lesions, consolidation and cavity | Negative | Aspergillus and Mucor | Pulmonary mucormycosis was diagnosed at first and showed a good response to posaconazole because of intolerance to amphotericin B due to heart arrest. There was new occurrence of hemoptysis and re-enlargement of original lesions 5 months later, and invasive pulmonary aspergillosis was diagnosed, with a good response to voriconazole. | Survival |
| Case No. | Sex | Age | Smoking history | Comorbidity | Diagnosis of mycosis | Classification of IFD by EORTC/MSG | Host factors | Clinical features | Mycological evidence | Biopsy | Clinical course of illness | Outcome |
|---------|-----|-----|-----------------|-------------|---------------------|----------------------------------|--------------|-----------------|---------------------|---------|--------------------------|---------|
| 14      | Male | 37  | No              | None        | Pulmonary cryptococcosis and pulmonary aspergillosis | Proven          | NA            | Lesions         | Cryptococcal antigen in blood and Aspergillus and Cryptococcus | Pulmonary cryptococcosis was diagnosed at first and showed a good response to fluconazole. There was re-enlargement of original lesions 14 months later, and mixed mycosis was diagnosed, with a good response to voriconazole. | Survival |
| 15      | Male | 48  | No              | Type 2 diabetes mellitus and diabetic nephropathy | Pulmonary mucormycosis and pulmonary aspergillosis | Proven          | NA            | Lesions and consolidation | Negative | Aspergillus and Mucor | Pulmonary mucormycosis was diagnosed at first and showed a good response to successive amphotericin B plus posaconazole and posaconazole. There was enlargement of original lesions 8 months later, and mixed mycosis was diagnosed, with a good response to voriconazole. | Survival |
| 16      | Male | 62  | No              | Chronic renal failure | Disseminated Talaromyces marneffei infection (lung and skin) and pulmonary cryptococcosis | Proven          | NA            | Lesions         | Cryptococcal antigen in blood and Talaromyces marneffei by culture | Mixed infection was diagnosed concurrently, with a good response to successive amphotericin B and voriconazole. | Survival |
| 17      | Male | 66  | Yes             | Type 2 diabetes mellitus and hypertension | Invasive pulmonary aspergillosis and pulmonary mucormycosis | Proven          | NA            | Lesions and cavity | Negative | Aspergillus and Mucor | Mixed infection was diagnosed concurrently, with a good response to successive amphotericin B plus posaconazole. | Survival |
than in patients without *Mucor* coinfection (Figure 3). Patients with mixed *Mucor* infection had lower hemoglobin levels than patients without mixed *Mucor* infection (97 ± 15 g/L vs. 118 ± 23 g/L, *P* = 0.037). Cavity, multiple nodule effusion/consolidation and enlarged mediastinal lymph nodes were more common in *Mucor* mixed infection than in non-*Mucor* mixed infection (Figure 3). Mortality was similar in the two groups (12.5% vs. 11.1%, *P* > 0.05).

**TABLE 2** | Clinical characteristics of 17 patients with mixed fungi.

| Clinical features | N(%)
|-------------------|------
| Fever             | 11 (64.7%) |
| Cough             | 15 (88.2%) |
| Expectoration     | 15 (88.2%) |
| Hemoptysis        | 8 (47.1%) |
| Shortness of breath | 7 (41.2%) |
| Chest pain        | 4 (23.5%) |
| Weight loss       | 2 (11.8%) |
| Wet rales         | 4 (23.5%) |
| White blood cell count>10×10^9/L | 3 (17.6%) |
| White blood cell count<10×10^9/L | 14 (82.4%) |
| Hemoglobin (g/L, mean ± SD) | 108.5 ± 21.9 |
| Lactic dehydrogenase (U/L, mean ± SD) | 193.6 ± 62.2 |
| Aspartate aminotransferase (U/L, mean ± SD) | 29.3 ± 32.1 |
| Alanine aminotransferase (U/L, mean ± SD) | 20.9 ± 14.9 |
| Albumin (g/L, mean ± SD) | 37.4 ± 7.2 |
| C-reactive protein (g/L, mean ± SD) | 6.7 ± 7.5 |
| PCT (ng/L, mean ± SD) | 0.1 ± 0.2 |
| CD4+ T lymphocytes (cells/µl) | 780 ± 389 |
| CD8+ T lymphocytes (cells/µl) | 822 ± 475 |

Data are shown as numbers (%), unless otherwise specified.

**TABLE 3** | Manifestations of lung CT images and distribution of lesions in patients with mixed fungi.

| Imaging feature or location of lesions | N (%) |
|---------------------------------------|-------|
| Lobulation                            | 2 (11.8%) |
| Cavity                                | 10 (58.8%) |
| Necrosis                              | 1 (5.9%) |
| Single nodule/mass                    | 1 (5.9%) |
| Multiple nodules/masses               | 7 (41.2%) |
| Effusion or consolidation             | 9 (52.9%) |
| Lymph node enlargement                | 12 (70.6%) |
| Thickening of the pleura              | 1 (5.9%) |
| Pleural effusion                      | 7 (41.2%) |
| Left upper lobe                       | 8 (47.1%) |
| Left lingual lobe                     | 5 (29.4%) |
| Left lower lobe                       | 9 (52.9%) |
| Right upper lobe                      | 11 (64.7%) |
| Right middle lobe                     | 7 (41.2%) |
| Right lower lobe                      | 10 (58.8%) |

**DISCUSSION**

In the past, it was believed that pulmonary mycosis occurred only in immunosuppressed hosts, with *Candida*, *Aspergillus* and *Cryptococcus* as the most common opportunistic fungi (Pfaller and Diekema, 2004), but increasing evidence shows that pulmonary mycosis in patients without apparent immune deficiency diseases is not uncommon, especially pulmonary cryptococcosis, which mainly occurs in immunocompetent hosts in mainland China. In our study, patients without apparent immunocompromising disease or immunosuppressive treatment accounted for an extremely large proportion.

**FIGURE 1** | Serial morphologic changes on chest CT of patients with mixed mycosis. (A–D) CT images in case 3 showed that compared to the lesion at the time of pulmonary aspergillus diagnosis (A), the lesion in the right lower lobe was smaller at 3 months after voriconazole treatment (B), re-enlarged 5 months after voriconazole treatment (C), and almost disappeared at 9 months after the diagnosis of mixed mycosis and posaconazole treatment (D). (E–H) CT images in case 14 showed that compared to the lesion at the time of pulmonary cryptococcosis diagnosis (E), the lesion in the right lower lobe was smaller 8 weeks after fluconazole treatment (F), re-enlarged 10 months after fluconazole treatment (G), and almost disappeared at 1 year after the diagnosis of mixed mycosis and voriconazole treatment (H).
FIGURE 2 | Pathology of patients with mixed mycosis. (A) Multiple spores can be seen in the cytoplasm of the multinucleated giant cells in the granulomatous nodules of lymphoid tissue in case 5. GMS staining (400x) showed that the fungi grew in clusters, with separation within the fungus and a small black spot in the center, which was considered Talaromyces marneffei (long red arrow). (B) Focal necrosis containing round or semilunar spores can be seen under the bronchial mucosal tissue with GMS staining (400x), which was considered to be Cryptococcus (long red arrow) in case 10. (C, D) Two forms of hyphae and spores can be seen in the necrotic foci of lung tissue by GMS staining (400x) in case 15. Fungi with the same thickness of hyphae and acute angle of branches were considered Aspergillus (long red arrow) (C), fungi with different sizes of hyphae and strange shapes of branches were considered Mucor (long red arrow) (D). (E) Two forms of hyphae and spores can be seen in the necrotic foci of lung tissue by GMS staining (400x) in case 17. Fungi with different sizes of hyphae, strange shapes of branches and thickened capsules were considered Mucor (long red arrow); fungi with the same thickness of hyphae and acute angle of branches were considered Aspergillus (short blue arrow).
It should be noted that only 2 cases in this study were in classical immunocompromised hosts, and one of them developed a mixed infection with *Talaromyces marneffei* in the present study. Another two cases of *Talaromyces marneffei* mixed infection were confirmed to have atypical types of immunodeficiency after the occurrence of *Talaromyces marneffei* infection, one with STAT3 hyper-IgE syndrome and the other with anti-IFN-γ autoantibody-associated immunodeficiency syndrome. The former was originally characterized by recurrent cold staphylococcal abscesses, pneumonia, eczema, hyperextensibility, and extreme elevation of IgE levels (Holland et al., 2007). The latter is associated with severe disseminated nontuberculous mycobacterial infections and other disseminated opportunistic infections (*Salmonella*, *Histoplasma*, and *Cryptococcus*) in previously healthy adults (Hong et al., 2020). However, recent studies have shown that these populations are also susceptible to infection with *Talaromyces marneffei*, which was previously a significant infectious complication in HIV/AIDS patients but has now increased in incidence in patients with other immune defects (Cao et al., 2019; Lee et al., 2019). Therefore, it is necessary to identify the potential cause of immunodeficiency in cases of *Talaromyces marneffei* infection, and people with unapparent immunodeficiency need to pay attention to the possibility of mycoses, especially those caused by *Talaromyces marneffei*.

In the present study, all but one patient infected by *Mucor* had type 2 diabetes mellitus. The incidence of diabetes mellitus was more common in the mucormycosis group than in the non-mucormycosis group. This result was consistent with the literature indicating that diabetes mellitus, with or without ketoacidosis, is one of the risk factors for *Mucor* infection (Cornely et al., 2019). Although the mechanism of immunodeficiency in diabetic patients is still unclear and needs in-depth research, if diabetic patients with pulmonary mycosis do not have a good response to the existing antifungal treatment that does not cover *Mucor*, attention should be given to the possibility of *Mucor* coinfection. Additionally, there were 3 non-immunocompromised hosts with mixed fungal diseases and no underlying diseases. It is valuable information for clinicians because not only single pulmonary fungal disease but also mixed fungal diseases can occur in patients without apparent immune deficiency diseases. According to our experience, during the follow-up, if the patients with proven pulmonary fungal disease showed a poor response to antifungal treatment, the existence of another fungal disease needs to be considered even in patients without apparent immune deficiency diseases.

Regardless of the immune status, among the single IFDs, *Aspergillus* and *Cryptococcus* are the most common pathogens, while *Mucor* and *Talaromyces marneffei* are relatively rare (Letourneau et al., 2014; Yan et al., 2016; Cao et al., 2019). *Pneumocystis jiroveci* is also one of the most common pathogens in immunosuppressed hosts (Azoulay et al., 2019). However, the mixed mycoses reported in the literature are mainly mixed infections of *Aspergillus* and *Cryptococcus* (Lin et al., 2006; Enoki et al., 2012; Wang et al., 2018; Awari et al., 2020). There have also been reports of *Aspergillus* and *Mucor* coinfection in the lung or outside the lungs (Bergantim et al., 2013; Mahadevaiah et al., 2013; Webb et al., 2013). Most *Cryptococcus* and *Talaromyces marneffei* coinfections have been reported in HIV patients (Le et al., 2010; Groll et al., 2021). However, in the present study, mixed fungal infections with *Aspergillus* and *Mucor* were the most common types, followed by

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**FIGURE 3** | Clinical features of mixed mycosis patients with and without *Mucor* infection. *Comparisons between *Mucor* mixed mycosis and non-*Mucor* mixed mycosis showed significance (P < 0.05).
Aspergillus and Cryptococcus coinfection, and coinfection with Talaromyces marneffei and other fungi was not uncommon in HIV-negative patients. This information is valuable for clinicians. When the initial antifungal treatment is not effective and mixed fungal diseases are considered, the possibility of Mucor should be considered, especially in patients who have diabetes mellitus. Therefore, fungus coinfection is worthy of consideration.

The symptoms, signs and imaging findings of mixed fungal lung infection lack specificity. There is no halo sign or crescent sign, which is not consistent with the imaging features of previous invasive pulmonary aspergillosis with a "halo sign at the early stage and crescent sign at the later stage" (Park et al., 2010). Classical chest CT images of Mucor infection, such as reversed halo signs and vascular occlusion, were also lacking in the present Mucor patients. Although Mucor patients more commonly showed multiple nodules and/or masses, effusion or consolidation and enlarged mediastinal lymph nodes in chest CT scans than non-Mucor coinfection patients, it was still difficult to diagnose Mucor coinfection, for which the prescription of targeted treatment amphotericin B liposome is recommended (Cornely et al., 2019). Hence, the diagnosis of proven invasive pulmonary mycosis mostly depends on histopathologic, cytopathologic, or direct microscopic examination of a specimen obtained by needle aspiration or biopsy.

However, the reality is that the diagnosis of pulmonary mycosis based solely on pathology may lead to misdiagnosis and missed diagnosis. Especially when distinguishing various molds, such as Aspergillus and Mucor, confusion occurs easily because of the similarity in the morphology of these two molds. Discrimination highly depends on skill and experience. In this study, the pathological judgment of some cases also exhibited this problem. It is worth noting that for 3 patients, mixed mycosis was missed in the biopsy in the hospital to which they were first admitted. The diagnosis of mucormycosis was made after reviewing the original pathological slides from the hospitals to which they were first admitted. As stated in the EORTC/MSGERC consensus definitions of IFDs, amplification of fungal DNA by PCR combined with DNA sequencing when molds are seen in formalin-fixed paraffin-embedded tissue can be used as one of the criteria for the diagnosis of infection by molds in these situations (Donnelly et al., 2019).

Additionally, lesions in 7 patients re-enlarged after the original effective treatment and were diagnosed after a second biopsy and pathological examination. Hence, based on the comprehensive patient situation, clinicians must also have rigorous procedures in the management of patients and analyze the reasons for ineffectiveness of original treatment or re-enlargement of lesions, including the sufficient triazole concentration in blood (Di Paolo et al., 2021), drug resistance (Yousefian et al., 2021), and the possibility of mixed infection with multiple pathogens such as bacteria, tuberculosis, other fungi, etc. As shown in this study, mixed mycosis is also one of the factors that needs to be considered, even in immunocompetent patients. Hence, we recommend that the following procedures can be considered in patients with pulmonary mycosis when poor response to initial antifungal treatment happens or the initial lesions grow again, including whether treatment drug monitor shows adequate concentrations of azoles (Di Paolo et al., 2021), whether there is new occurrence of symptoms or characteristic imaging of other types fungal infections other than tuberculosis or non-tuberculosis mycobacteria, whether current in use antifungal agents could cover suspected pathogens such as fluconazole is invalid for aspergillus and voriconazole is invalid for mucor. At this time, targeted fungal etiology and pathological examination are required to confirm the diagnosis.

Except for early complete surgical treatment being strongly supported for mucormycosis whenever possible, systemic antifungal treatment is recommended as the first choice for other fungi by guidelines (Cornely et al., 2019; Garcia-Vidal et al., 2019; Groll et al., 2021). In the present study, except for one patient who died because of refusing further treatment, all mucormycosis patients recovered after receiving systemic amphotericin B treatment. One mucormycosis patient also received surgery after systemic antifungal treatment. Three cases with non-Mucor mixed infection received surgery after poor response to systemic antifungal treatment or for diagnosis purposes. Hence, surgery is an alternative to systemic antifungal therapy in some mixed pulmonary mycoses with a poor response. In contrast to the high mortality in immunocompromised hosts in the literature, most mixed mycosis cases in the present study had a good response.

The major limitations of our study were that it was a retrospective, single-center study. There may have a bias in patient enrollment. Another limitation was that not all patients were screened for gene mutations, such as the STAT 3 gene and the level of anti-interferon γ antibody. In the present study, only three cases with Talaromyces marneffei infection were tested for the STAT 3 gene and anti-interferon γ antibody, with one of them positive for each.

**CONCLUSION**

Mixed pulmonary mycosis is not uncommon in immunocompetent hosts. During the management of mycosis, we recommend keeping mixed mycosis in mind for patients with a poor response to initial antifungal treatment, even in immunocompetent populations. A rigorous procedure may help to differentiate mixed mycosis from single mycosis.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

This study was approved by the ethics committee of the First Affiliated Hospital of Guangzhou Medical University (2018-119).
Written informed consent for participation was not required for this study in accordance with the national legislation and the institutional requirements.

**AUTHOR CONTRIBUTIONS**

Conception and design: YZ, CL, and FY. Collection and assembly of data: YZ, CL, SL, JZ, ZL, and YG. Data analysis and interpretation: YZ. Manuscript writing: YZ, CL, and FY. Final approval of manuscript: YZ, CL, SL, JZ, ZL, YG, and FY. All authors contributed to the article and approved the submitted version.

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