Pulmonary Hypertension Caused by Fibrosing Mediastinitis

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ABSTRACT

Pulmonary hypertension (PH) is a progressive and severe disorder in pulmonary hemodynamics. PH can be fatal if not well managed. Fibrosing mediastinitis (FM) is a rare and benign fibroproliferative disease in the mediastinum, which may lead to pulmonary vessel compression and PH. PH caused by FM (PH-FM) is a pathologic condition belonging to group 5 in the World Health Organization PH classification. PH-FM has a poor prognosis because of a lack of effective therapeutic modalities and inappropriate diagnosis. With the development of percutaneous pulmonary vascular interventional therapy, the prognosis of PH-FM has been greatly improved in recent years. This article provides a comprehensive review on the epidemiology, pathophysiologic characteristics, clinical manifestations, diagnostic approaches, and treatment modalities of PH-FM based on data from published reports and our medical center with the goal of facilitating the diagnosis and treatment of this fatal disease. (JACC: Asia 2022;2:218–234) © 2022 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Pulmonary hypertension (PH) is a progressive, severe, and hemodynamic disorder that may cause high mortality if not well treated.1 PH caused by fibrosing mediastinitis (PH-FM), a rare type of the condition in group 5 PH according to the World Health Organization PH classification, has a poor prognosis because of a lack of effective therapeutic modalities. Misdiagnosis and underdiagnosis of PH-FM are common.1,2

Fibrosing mediastinitis (FM), also known as sclerosing mediastinitis or mediastinal fibrosis, is a rare and benign fibroproliferative disease in the mediastinum.3 Proliferative fibrous tissue gradually replaces normal fat tissue and wraps, infiltrates, and compresses the adjacent structures in the mediastinum, such as pulmonary vessels, superior vena cava (SVC), bronchus, esophagus, and pericardium. The aberrant behavior of the proliferative fibrous tissue may cause PH, SVC syndrome, atelectasis, and obstructive pneumonia, among which PH and the resulting right heart failure are the most prevalent sequels leading to death.3 In recent years, with the development of percutaneous pulmonary vascular interventional therapy, the prognosis of PH-FM has been substantially improved.3,4 This paper provides a comprehensive review of the etiology, epidemiology, pathophysiologic characteristics, clinical manifestations, imaging features, diagnostic criteria, and treatment modalities of PH-FM according to published reports (see the search method in Supplemental...
PH Caused by FM

**Etiology, Pathogenesis, and Epidemiology**

FM is currently deemed as an abnormal immune proliferative reaction in the mediastinum in response to the triggering factors, including pathogens (*Histoplasma capsulatum, Mycobacterium tuberculosis, Blastomyces* and *Aspergillus* species, and so on), sarcoidosis, autoimmune diseases (immunoglobulin G4-related disease, Behcet’s disease, and systemic sclerosing disease, among others), iatrogenic (radiotherapy and esophageal injury, and so forth), and idiopathic FM. FM is categorized into granulomatous FM (also known as the focal subtype) and nongranulomatous FM (also designated as the diffuse subtype or idiopathic FM). Granulomatous FM is more prevalent, accounting for 80% to 90% of FM, and is elicited by infectious pathogens and inflammatory diseases such as sarcoidosis, whereas nongranulomatous FM is induced by autoimmune diseases, which often involve extramedialstinal disorders, such as retroperitoneal fibrosis, sclerosing cholangitis, autoimmune pancreatitis, and Riedel thyroiditis (Table 1).

The pathogenesis of FM remains elusive. It has been hypothesized that FM may progress through 3 stages following infection with *Histoplasma capsulatum*. The first stage is an acute infection of *Histoplasma capsulatum*. Patients may exhibit fever, myalgia, cough, severe pneumonia, and respiratory failure; however, most patients present with covert infection and have no salient clinical manifestation. The second stage is mediastinal granuloma, a cheese-like mass formed by enlarged and fused lymph nodes. Patients in the first stage rarely progress to the second stage. The last stage is FM, which elicits clinical symptoms because of the compression of mediastinal structures by excessive proliferation of fibrous tissue. Although the pathogenesis of FM caused by *Mycobacterium tuberculosis* infection or other triggers has not been reported, it may be similar to that of FM following *Histoplasma capsulatum* infection. PH-FM is mainly ascribed to the pathogenic factors causing stenosis of pulmonary arteries (PAs) and pulmonary veins (PVs).

The most prevalent trigger of FM is *Histoplasma capsulatum* infection (H-FM) in the United States and *Mycobacterium tuberculosis* infection (TB-FM) in China. The median age of patients with FM is 42 years in the United States, whereas the average age is 69.5 years in China from a small-sample study. Although histoplasma infection is common in the United States, it is often asymptomatic and rarely progresses to FM. It has been reported that 3 in 100,000 patients with histoplasma infection develop FM. Dines et al. showed that 11 of 31 patients with mediastinal granuloma eventually developed FM. The incidence of FM caused by *Mycobacterium tuberculosis* infection has not yet been reported. Although there is no official report on the prevalence of PH in FM, it is of note that most of the FM cases recruited in previous small-sized cohort studies are accompanied by PH upon admission.

In our center, we registered 161 patients with FM, and 139 of them (67 men) were evaluated by echocardiography. Among the evaluated FM patients, 91 (47 men; age range: 50-90 years) and the data from our medical center with the goal of facilitating the standardization of the diagnosis and therapeutics of this fatal disease.

**TABLE 1 The Etiologies of Fibrosing Mediastinitis**

| Classification | Etiologies                                                                 |
|----------------|---------------------------------------------------------------------------|
| Idiopathic FM  | Infection associated7-10, Histoplasmosis capsulatum, Tuberculosis, Aspergillosis, Mucormycosis, Blastomycosis, Actinomycosis, Nocardiosis, Coccidioidomycosis, Cryptococcosis, Non-infection associated5-14, Sarcoïdosis, Immunoglobulin G4-related disease, Behcet disease, Systemic sclerosing disease, Rheumatic fever, Hodgkin disease, Silicosis, Trauma, Iatrogenic5-18, Radiotherapy, Chest surgery, Esophageal fistula, Methysergide maleate, Microinvasive procedures in the mediastinum (endoscopic ultrasonic-guided fine-needle aspiration, drainage tube placement) |

**ABBREVIATIONS AND ACRONYMS**

- CT = computed tomography
- CTPH = chronic thromboembolic pulmonary hypertension
- CTPA = computed tomography pulmonary angiography
- CTPV = computed tomography pulmonary venography
- ECG = electrocardiogram
- FM = fibrosing mediastinitis
- H-FM = histoplasma capsulatum infection-related fibrosing mediastinitis
- Ig = immunoglobulin
- MRI = magnetic resonance imaging
- PA = pulmonary artery
- PAG = pulmonary artery angiography
- PET-CT = positron emission tomography/computed tomography
- PH = pulmonary hypertension
- PH-FM = pulmonary hypertension caused by fibrosing mediastinitis
- PV = pulmonary vein
- PVG = pulmonary vein angiography
- RHC = right heart catheterization
- SPECT/CT = single-photon emission computed tomography/computed tomography
- SVC = superior vena cava
- TB = tuberculosis
- TB-FM = Mycobacterium tuberculosis infection-related fibrosing mediastinitis
- V/Q = lung ventilation/perfusion
### TABLE 2 The Characteristics of Fibrosing Mediastinitis Caused by Different Triggers

| Features                                      | H-FM<sup>a</sup> | TB-FM<sup>b</sup> | Idiopathic FM<sup>c</sup> |
|----------------------------------------------|------------------|--------------------|--------------------------|
| **Pathophysiology**                          |                  |                    |                          |
| Granulomatous                                 |                  |                    |                          |
| Focal, invasive, calcified lesions in the mediastinum caused by abnormal immune reaction to pathogen |                  |                    |                          |
| Granulomatous                                 |                  |                    |                          |
| Focal, invasive, calcified lesions in the mediastinum caused by abnormal immune reaction to pathogen |                  |                    |                          |
| Nongranulomatous                              |                  |                    |                          |
| Extensive, diffuse, noncalcified proliferation of fibrous tissue in the mediastinum caused by autoimmune diseases, radiotherapy, IgG4-related disease, and drugs |                  |                    |                          |
| **Prevalence**                                |                  |                    |                          |
| 3/100,000 persons                             | Unknown          | Unknown            |                          |
| **Demographics**                              |                  |                    |                          |
| Median age: 42 y (range: 21-75 y)              | Unknown          | More common in China |                          |
| More sex differences                         | Unknown          | Unknown            |                          |
| More common in the United States             | Unknown          | Unknown            |                          |
| **Symptoms (depending on affected structures)**| Common symptoms are dyspnea, cough, chest pain, SVC syndrome (less in TB-FM), hemoptysis, pleural effusion (less in H-FM) | Common symptoms are dyspnea, cough, chest pain, SVC syndrome (less in TB-FM), hemoptysis, pleural effusion (less in H-FM) | Common symptoms are dyspnea, cough, chest pain, SVC syndrome (less in TB-FM), hemoptysis, pleural effusion (less in H-FM) |
| **Location in mediastinum**                  |                  |                    |                          |
| Focal > diffuse                               | Diffuse or focal | Bilateral > unilateral |                          |
| Unilateral > bilateral                       | Diffuse or focal | Bilateral or unilateral |                          |
| **Mainly affected mediastinal structures**   |                  |                    |                          |
| Pulmonary arteries                            | Pulmonary arteries | Pulmonary arteries | Pulmonary arteries |
| Bronchi                                       | Bronchi          | Bronchi            | Bronchi |
| SVC                                          | Pulmonary veins (less) | Pulmonary veins | Pulmonary veins |
| Pulmonary veins (less)                        | SVC (less)       | SVC                | SVC |
| **Extrathoracic involvement**                | None             | None               | Retroperitoneal fibrosis |
|                                              |                  |                    | Autoimmune pancreatitis |
|                                              |                  |                    | Sclerosing cholangitis |
| **Imaging**                                  |                  |                    |                          |
| Mediastinal widening                         | Mediastinal widening | Mediastinal widening |                           |
| Nodal calcification                          | Nodal calcification | Nodal calcification |                           |
| Abnormal soft tissue (focal)                  | Abnormal soft tissue (focal) | Abnormal soft tissue (focal) |                           |
| Compression of mediastinum structures including vessels, airway, and SVC | Compression of mediastinum structures including vessels, airway, and SVC | Compression of mediastinum structures including vessels, airway, and SVC |                           |
| **Treatment**<sup>5,6,8,25,29,36,74,75,83</sup> |                  |                    |                          |
| Medication                                   |                  |                    |                          |
| Antifungal and anti-inflammatory drugs are ineffective<sup>c</sup> | Anti-TB and anti-inflammatory drugs are ineffective<sup>c</sup> | Anti-TB and anti-inflammatory drugs are ineffective<sup>c</sup> | Corticosteroids are of varied benefit<sup>d</sup> |
| Rituximab may be effective<sup>c</sup>        | Rituximab may be effective<sup>c</sup> | Rituximab may be effective<sup>c</sup> | Rituximab may be effective<sup>c</sup> |
| Vasodilator drugs for pulmonary hypertension may be ineffective<sup>c</sup> | Vasodilator drugs for pulmonary hypertension may be ineffective<sup>c</sup> | Vasodilator drugs for pulmonary hypertension may be ineffective<sup>c</sup> | Vasodilator drugs for pulmonary hypertension may be ineffective<sup>c</sup> |
| **Surgery**                                  |                  |                    |                          |
| Uncertain efficacy and high operation-related mortality rate (20%) |                  |                    |                          |
| **Endovascular therapy**                     | Preferred and effective for relieving pulmonary artery, pulmonary vein, and SVC stenosis, but the rate of intrastent restenosis is high in pulmonary veins and SVC | Preferred and effective for relieving pulmonary artery, pulmonary vein, and SVC stenosis, but the rate of intrastent restenosis is high in pulmonary veins and SVC | Preferred and effective for relieving pulmonary artery, pulmonary vein, and SVC stenosis, but the rate of intrastent restenosis is high in pulmonary veins and SVC |

<sup>a</sup>There may have some bias because of limited evidence, and there are some overlaps with H-FM, TB-FM, and idiopathic FM in pathophysiology, location in mediastinum, distribution of affected structures in mediastinum, symptoms, and imaging findings in some patients. It may depend on whether there is active fungal or TB infection and if the fibrous lesion is active. It was reported that rituximab was effective for patients with progressive FM in a small-sample case series study. Evidence comes from a small-sample retrospective study.<sup>7</sup><sup>2</sup> Evidence comes from a small-sample retrospective study.<sup>7</sup><sup>3</sup>

<sup>b</sup>H-FM = Histoplasma capsulatum infection-related fibrosing mediastinitis; IgG4 = immunoglobulin G4; SVC = superior vena cava; TB-FM = Mycobacterium tuberculosis infection-related fibrosing mediastinitis; TB = tuberculosis.

CLINICAL MANIFESTATIONS

The clinical symptoms of PH-FM depend on the involved structures in the mediastinum.<sup>2,6-8,25,28-40</sup>

Years; median age: 66 years) had confirmed or suggested tuberculosis (TB) infection according to the medical history, results of TB tests, and findings of imaging examinations. Furthermore, 83 of these 91 patients exhibited systolic pulmonary artery (PA) pressure of no <40 mm Hg. The prevalence of the involvement of the PAs, bronchi, PVs, and SVC were 100%, 98.9%, 94.5%, and 13.2%, respectively (Y. Cao, unpublished data, 2021) (Table 2). Disparate preferences and extents of involvement in the mediastinal structures were also found in patients with FM. For instance, Peikert et al<sup>25</sup> reported that the focal subtype accounts for 95% and the diffuse subtype for 5% in the FM cases studied; 72% of the cases involved the right side, 5% the left side, and 23% both sides. Moreover, 42% of the cases involved PAs, 13% PVs, and 42% the SVC. By contrast, another report demonstrated that of the 28 patients with TB-FM, 85% presented with bilateral lesions, 3% with SVC compression, 100% with PA compression, and 66% with PV involvement.<sup>28</sup> Additionally, another small-sized study showed that all the patients had bilaterally involved mediastinum and none had involved SVC.<sup>24</sup> The differences in the scope and location of the involved structures may be due to the distinct triggering factors of this disease. Collectively, H-FM predominantly affects the structures in the right upper mediastinum, including the PAs, bronchus, and SVC, but exerts fewer effects on PVs. TB-FM mainly involves the structures at bilateral hilar regions, such as PAs, bronchi, and PVs but rarely involves the SVC (Table 2).
In general, shortness of breath is the most frequent symptom of FM. The involved mediastinal structures include PAs, PVs, the SVC, and bronchi, with the corresponding symptoms as dyspnea, hemoptysis, SVC syndrome, and coughing. Specifically, compression of the PAs may incur PH and severe hemoptysis as a result of systemic collateral hyperplasia; compression of the PVs can lead to PH and pulmonary venous congestion, which, in turn, cause mild to moderate hemoptysis and refractory pleural effusion, and eventually, pulmonary vascular compression may lead to severe PH, right heart failure, and even death. Compression of the SVC can cause SVC syndrome. If the bronchus is involved, the patient may experience wheezing, cough, atelectasis, and obstructive pneumonia. Involvement in the recurrent laryngeal nerve and phrenic nerve may cause hoarseness or shortness of breath; involvement with the esophagus and the thoracic duct can elicit dysphagia and chylothorax, respectively; and compression of the pericardium may cause pericardial tamponade and constrictive pericarditis. The SVC syndrome is more prevalent in H-FM than in TB-FM, whereas refractory pleural effusion exhibits the opposite trend.

**IMAGING FINDINGS**

**ELECTROCARDIOGRAM AND ECHOCARDIOGRAPHY.**

The typical electrocardiogram (ECG) manifestations of PH-FM are similar to those of pulmonary arterial hypertension and PH caused by other types of PA stenosis. The characteristic ECG includes right deviation axis, uncertain axis, R/S <1 in lead I, R/Q >1 in lead aVR, and right bundle branch block and T-wave inversion/flatness in limb and/or precordial leads, indicating enlargement of and endocardial ischemia in the right ventricle caused by pressure overload. These typical ECG characteristics are termed PAS syndrome. The abnormal ECGs can be partially or totally recovered following the amelioration of pulmonary vascular stenosis (Figure 1).

Echocardiogram has no finding specific for PH-FM. All PH signs can be present in patients with PH-FM, including elevated systolic right ventricle pressure, enlarged right heart, and compressed left ventricle. Proximal PV lesions can be revealed by echocardiography in some patients with PH-FM.

**CHEST X-RAY.** Patients with PH-FM usually show signs of FM, including widening of the mediastinum, atelectasis, mediastinal masses and hilar enlargement, and pulmonary congestion or interstitial pulmonary edema in the upper lobes, as well as signs of PH, including a prominent main PA and enlarged right heart on chest radiographs. The terminology “FM dyad” and “FM triad” have been used to describe the chest x-ray manifestations of PH-FM. The FM dyad includes one of the signs of PH such as prominent main PA and atelectasis, whereas FM triad refers to the dyad plus refractory pleural effusion (Figure 2A). Additionally, pulmonary congestion or interstitial pulmonary edema in the upper lobes, as one of the signs of upper PVs stenosis, could have the same implication as refractory pleural effusion for PH-FM with PV stenosis. Accordingly, the FM triad could be modified to an FM dyad plus refractory pleural effusion or pulmonary congestion or interstitial pulmonary edema in the upper lobes. These 4 signs could be termed as “FM tetralogy” (Figure 2A). Our preliminary data indicated that the sensitivity of atelectasis in combination with pleural effusion for diagnosing PH-FM was 50% and that the specificity was >90% (Y. Cao, unpublished data, 2021). Therefore, the FM dyad and FM triad may provide important clues for diagnosis of PH-FM.

**VENTILATION/PERFUSION SCINTIGRAPHY.** Ventilation/perfusion scintigraphy (V/Q) scan is a preferred screening method for chronic stenotic pulmonary vascular disease. V/Q scan imaging in PH-FM

| TABLE 3 | The Involved Mediastinal Structures and the Associated Symptoms and Signs in Patients With Pulmonary Hypertension Caused by Fibrosing Mediastinitis |
|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| Involved Structures | Associated Symptoms/Signs | Reference(s) |
| Pulmonary artery | Dyspnea, Chest pain, Hemoptysis, Pulmonary hypertension, Enlarged right heart, Prominent main pulmonary artery, Mosaic perfusion | 8,20,26,30,35,46 |
| Pulmonary vein | Dyspnea, Hemoptysis, Refractory pleural effusion, Pulmonary edema, Enlarged right heart, Prominent main pulmonary artery, Mosaic perfusion, Thickening of interlobular septum | 20,26,31,32,35,48 |
| Bronchus | Cough, Dyspnea, Lithiopathy, Postobstructive pneumonia, Atelectasis, Thickening of interlobular septum | 20,26,34 |
| Superior vena cava | Superior vena cava syndrome | 33 |
| Esophagus | Dysphagia, Odynophagia | 30 |
| Pericardium | Constrictive pericarditis | 35 |
| Thoracic duct | Chylothorax | 37 |
| Recurrent laryngeal nerve | Hoarseness | 38 |
| Coronary artery | Angina, Acute coronary syndrome, Sudden death | 39,40 |
patients usually shows multiple adjacent segmental perfusion defects with mismatched ventilation impairment (Figure 3). These imaging characteristics, known as mismatched perfusion defects (Figure 3), are similar to those exhibited by pulmonary embolism. Therefore, misdiagnosis between PH-FM and pulmonary embolism should be avoided. In addition, the PH-FM patients with severe bronchus
stenosis and atelectasis may have impaired ventilation, which could be matched with the perfusion defects induced by the stenosis of corresponding PAs. Single-photon emission computed tomography/computed tomography (SPECT/CT) integrates functional imaging with anatomic imaging and has been proven as effective as V/Q scan in diagnosing chronic thromboembolic PH (CTEPH). In view of the definite imaging characteristics such as FM dyad and triad, SPECT/CT might have advantages over V/Q scan in the differential diagnosis of PH-FM from other stenotic pulmonary vascular diseases.
CONTRAST-ENHANCED CHEST COMPUTED TOMOGRAPHY. PH-FM usually manifests as focal or diffuse hyperplasia of fibrous tissues compressing adjacent structures with or without calcification on chest computed tomography (CT) (Figures 2B, 2C, and 4A to 4C). Other findings on CT include atelectasis, mosaic perfusion, ground glass density shadow, thickening of the interlobular septum, and hyperplasia of the systemic collateral vessels. The FM dyad and triad can be observed in typical chest CT (Figures 4A to 4C). Contrast-enhanced chest CT is the first option of imaging examination to diagnose suspected PH-FM, evaluate the extent of involvement in mediastinum structures (Figures 2B, 2C, and 4A to 4C), and detect hyperplasia of the collateral vessels. The results of this examination suggest the clinical classification of PH-FM and facilitate the determination of a therapeutic regimen. In particular, the multiplanar and virtually reconstructed images further illustrate the length and diameter of target lesions and provide the best projection view during interventional therapy, thereby subserving an important reference for making a strategy of intervention. In addition, contrast-enhanced chest CT is also crucial for evaluating right heart function. Moreover, metal artifacts of vascular stents on MRI influence the evaluation of intrastent restenosis; hence, CT pulmonary angiography and venography (CTPA and CTPV, respectively) could be the first choice of examination during follow-ups after interventional therapy. However, MRI is still the gold standard for evaluating right heart function.

POSITRON EMISSION TOMOGRAPHY/CT. Positron emission tomography/CT (PET-CT) is rarely used for the diagnosis and evaluation of PH-FM. The published reports on the use of PET-CT in FM diagnosis are all case reports and showed substantial variability in fluorodeoxyglucose uptake by mediastinal lesions (Figure 5).

PA AND PV ANGIOGRAPHY. PA angiography (PAG)/PV angiography (PVG) is the gold standard for evaluating vascular stenosis but not for tracing the etiology of pulmonary vascular stenosis. PH-FM is mainly manifested as localized stenosis at the origin of the second and third branches of the PA in PAG and at the first drainage branches of the PV in PVG around the hila (Figure 6, Video 1). However, PAG and PVG cannot accurately determine the cause of vascular stenosis, whether there are hyperplastic soft tissues or tumors in the mediastinum, whether the trachea/bronchus is involved, and whether calcification exists. In addition, the transeptal puncture entailed by direct PVG adds operational difficulties. Therefore, CTPA, CTPV, and MRI are the major diagnostic methods for PH-FM. Once the diagnosis is secured, either PAG or PVG or both are then used for evaluation before angioplasty and/or venoplasty.
PATHOLOGIC STUDY

Biopsy is the gold standard for pathologic diagnosis of FM and a critical approach to rule out malignancy. However, biopsy is not necessary for issuing a diagnosis of FM because of its invasive nature and the complications it may incur. The pathologic analysis reports nodular or diffuse hyperplastic fibrous tissues surrounding and infiltrating mediastinal structures. In addition, Peikert et al found that infiltration in proliferative fibrous tissues of the mixed lymphocytes, in which CD20-positive B lymphocytes accounted for a considerable portion, implicating an association of these B lymphocytes with FM (Figure 7).

HEMODYNAMICS

Right heart catheterization (RHC) is the gold standard for diagnosing PH. PH-FM mostly belongs to precapillary PH. Although some patients have PV stenosis, their PA wedge pressure measured by Swan-Ganz catheter is mostly normal. This is because the involved PVs are predominantly located at the proximal first drainage branches, which are usually smaller in size than the balloon of a Swan-Ganz catheter. This scenario is comparable to the apparently normal PA wedge pressure exhibited during PV occlusive disease. RHC can also measure the pressure gradient across the stenotic site in pulmonary vessels, evaluate the functional impairment of stenotic vessels, and provide detailed hemodynamic information for interventional modality.

CLINICAL CLASSIFICATION

PH-FM is categorized into 3 types based on the involved pulmonary vessels. Type I refers to the FM that mainly causes stenosis in the PAs, mostly accompanied by the stenosis of anatomically adjacent bronchi but not the PVs; type II refers to the FM that predominantly induces stenosis of the PVs without involvement of the PAs, and bronchus involvement is
rare; type III refers to the FM eliciting stenosis of the PAs, PVs, and bronchi (Figure 8). Furthermore, if other structures, such as the SVC and esophagus, are involved, the first letter of the involved structure is used as a subscript to specify the categorization. For example, type I plus involvement of the SVC is designated as type I_\text{SVC}^{49}. The prevalence of different types may depend on the triggering factors of the disease. In our center, most cases were triggered by TB. Among these cases, type III accounted for about 92%, and only 1% were type II (Y. Cao, unpublished data, 2021).

**DIAGNOSTIC FLOW CHART**

The clinical manifestations, FM dyad and triad, as well as PAS syndrome of ECG provide useful clues for diagnosis. V/Q scan is an important screening tool. Contrast-enhanced CT is the major examination for diagnosing PH-FM, identifying the involved...
mediastinal structures, and determining the clinical classification. RHC should be performed to measure hemodynamic parameters. PH-FM should be distinguished from other types of PH, especially from other types of chronic stenotic pulmonary vascular diseases, including chronic thromboembolic disease, CTEPH, PV thrombosis, PV stenosis after radiofrequency ablation, pulmonary vascular stenosis caused by vasculitis, pulmonary vascular stenosis after surgical repair of congenital heart disease, and compressive pulmonary vascular stenosis caused by mediastinal tumor. For patients with PH as the chief complaint, doctors should follow the diagnostic flow chart described in the guidelines for PH.69 For patients with mismatched perfusion defects, the possibility of PH-FM as one of the diagnoses should be considered; for PH patients with atelectasis, widened mediastinum, and/or refractory pleural effusion, PH-FM is highly suspected, and CTPA and CTPV are required to verify the suspicion. After confirmed diagnosis of PH-FM, clinical classification should be determined and the distribution of involved mediastinal structures demarcated, whereby the appropriate therapeutic strategy can be selected. Thorough medical histories should be taken, and the immune parameters, such as IgG4 levels and tuberculosis antibodies, should be tested for routinely to identify the disease triggers. Retroperitoneal fibrosis, sclerosing cholangitis, and other clinical conditions need to be examined (Figure 9, Central Illustration).

**TREATMENT**

**MEDITATION.** The general treatment for patients with PH-FM includes oxygen therapy, diuretics, and digoxin. Antifungal, anti-TB, and anti-inflammatory
drugs are ineffective, although there is an association of histoplasmosis and TB infection with the occurrence of FM.7,8,23,25,70 Unless there is evidence of active infection with Histoplasma, such as mediastinal granuloma or the H or M band in immunodiffusion tests, the administration of itraconazole may not be effective.71,72 For sarcoidosis- or IgG4-related FM, glucocorticoid therapy may be effective.11,73 In addition, the efficacy of hormone therapy varies among individuals with idiopathic FM.74 Furthermore, Westerly et al75 reported the administration of rituximab to 3 FM patients whose pathologic biopsy specimens showed abundant infiltration of CD20-positive B lymphocytes in proliferative fibrous tissues and in whom PET-CT revealed high uptake of fluorodeoxyglucose and active lesions in the mediastinum. Following rituximab administration, the patients’ symptoms were relieved, the sizes of the lesions shrank, and the metabolic activities of the lesions diminished. However, the safety and efficacy of rituximab as a therapeutic modality for PH-FM needs to be further investigated by multicenter, large-sample, randomized controlled clinical trials.

**SURGERY.** The purpose of surgical treatment is to relieve the compression of mediastinal structures by proliferative fibrous tissues and ameliorate the compression-related symptoms (Table 2). Common surgical procedures include mediastinal tissue resection, SVC bypass grafting, airway reconstruction, pulmonary revascularization, lobectomy, and segmentectomy.7,25,76-79 As a result of the complexity of FM pathology, surgical treatment has a mortality rate as high as 20%.7 Moreover, the clinical benefits of these surgeries are not yet clear. Studies have shown that 42% of FM patients subjected to surgical treatment relapse during follow-up and require other interventions.25 The cause of recurrence may be related to residual obstruction and progression of fibrous lesions in the mediastinum, and this awaits further investigation.
(A) Type I. The PA and bronchus are compressed by proliferative fibrous tissues. (B) Type II. Only the PV is compressed by proliferative fibrous tissues. (C) Type III. The PA, PV, and bronchus are compressed by proliferative fibrous tissues. BA = bronchus airway; FT = proliferative fibrous tissues; LA = left atrium; PA = pulmonary artery; PH-FM = pulmonary hypertension caused by fibrosing mediastinitis; PV = pulmonary vein.
ENDOVASCULAR INTERVENTIONS. Interventional therapy has been used to relieve symptoms caused by obstruction of the pulmonary vessels in PH-FM patients (Table 2, Video 1). It is the preferred treatment for such diseases.5,6,36,80-84 Different interventional modalities have been proposed according to the patient’s clinical classification. In general, PV intervention should be performed first. After the stenosed PV has regained patency, the PA intervention may be performed soon; if the PV occlusion cannot be removed, the PA intervention should be avoided. Also, if the PA occlusion is prejudged not to be alleviated, the PV intervention should not be performed. A study with the largest sample size so far showed that 59 stents in 47 pulmonary vessels (26 PAs and 21 PVs) were implanted in 30 patients with FM.36 Symptoms were relieved significantly after the procedure; however, during the median 115-month follow-up period, symptomatic restenosis occurred in 1 case of PA (5%) and 4 cases of PV (25%).36 Moreover, 8 patients with FM-induced PV stenosis underwent percutaneous pulmonary venoplasty, and their symptoms were ameliorated, but 50% of the patients experienced restenosis.83 Notably, owing to the stiff, rigid, and brittle pulmonary vessel wall formed during the pathophysiologic process of this disease, PH-FM incurs a higher rate of complications than other types of pulmonary vascular stenosis during interventions. The complications include vessel injury, lung injury, lung edema, stent displacement, stent underexpansion, and intrastent restenosis.5,6,36,80-87 Therefore, although interventional treatment can significantly ameliorate symptoms and improve the hemodynamics of the patients with PH-FM in small-sized case studies, its efficacy and safety should be thoroughly assessed in the future. Importantly, perioperative management should be strengthened to avoid fatal complications. In addition, intrastent restenosis, especially restenosis of the PV stent, is a prevalent and recalcitrant issue to which close attention should be paid following interventional modalities of PH-FM. The specific mechanisms underlying and preventative schemes for intrastent restenosis warrant further study.

INTERVENTIONAL MODALITIES TO SVC STENOSIS AND BRONCHIAL STENOSIS. Interventional modalities can relieve SVC syndrome caused by FM-induced SVC stenosis, but restenosis is prone to occur after intervention (Table 2).23,36 For bronchial stenosis caused by FM, the stent strategy is difficult and perilous and, hence, is more challenging than the strategy for other benign lesions in the mediastinum (Table 2).34 Collectively, there is limited experience in the treatment of PH-FM, and no guideline or expert consensus has been established. Therefore, it is recommended that patients with PH-FM be referred to an experienced pulmonary vascular center for effective and safe treatment after careful evaluation by multidisciplinary approaches.
Central Illustration: Pulmonary Hypertension Caused by Fibrosing Mediastinitis

Etiology
- Tuberculosis
- Histoplasma
- Idiopathic
- Others

Symptoms
- Dyspnea
- Cough
- SVCS
- Hemoptysis
- Pleural effusion

Imaging clues
- Chest X-ray
- V/Q scan
- ECG

FM dyad/triad
- Predominant MPA
- Pleural effusion

Mismatches perfusion defects

PAS syndrome
- 

Diagnosis
- Evaluation
- Classification

CTPA
- RHC

Type III

Type I

Type II

Treatment
- Endovascular therapy
- Surgery
- Drugs

Effective Restenosis (PV)
- Uncertain
- High risk
- Variable

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AW = airway; CTPA = computed tomography pulmonary angiography; CTPV = computed tomography pulmonary venography; ECG = electrocardiogram; L = left; PA = pulmonary artery; PAS = pulmonary artery stenosis; PV = pulmonary vein; R = right; RHC = right heart catheterization.
HIGHLIGHTS

- PH-FM, as a type of rare condition in group 5 PH, has a poor prognosis because of a lack of effective therapeutic modalities and frequent misdiagnosis and underdiagnosis.
- The most prevalent trigger of FM is infection of Histoplasma capsulatum in the United States and infection of Mycobacterium tuberculosis in China.
- Imaging findings, including mismatched perfusion defects in the V/Q scan, FM dyad, and FM triad are important diagnostic clues, and clinical classification facilitates decision making in diagnosis and therapeutics.
- Because of the limited efficacy of drug therapy as well as the uncertain effectiveness and high risk of surgical treatment, endovascular intervention modality is currently the preferred therapeutic option, although procedure-related complications and intrastent restenosis after PV intervention need to be addressed.

PROGNOSIS

Because of the limited reports with limited sample sizes in the past, the prognosis of patients with PH-FM is not clear. Peikert et al conducted a retrospective study of 80 patients with FM admitted to the Mayo Clinic from 1998 to 2007; 2 patients (2.5%) died of FM during the follow-up period of a median of 68 months. Another study summarized the clinical data of 71 patients with FM and showed that the mortality rate was as high as 30% based on incomplete follow-up data.

In 2015, Seferian et al followed up 27 patients with PH-FM and found that survival rates at 1, 3, and 5 years postdiagnosis were 88%, 73%, and 56%, respectively.

CONCLUSIONS

In summary, the pathogenesis of PH-FM is still unclear, and there is no curable modality for it. The clinical manifestations of PH-FM lack specificity, and the rates of clinical misdiagnosis and underdiagnosis are high. For patients with PH, especially those with mismatched perfusion defects, SVC syndrome, FM dyad, and FM triad, doctors should consider the possibility of PH-FM. Contrast-enhanced chest CT can confirm the initial diagnosis and determine the clinical classification. In view of the limited efficacy of drug therapy as well as the uncertain effectiveness and high risk of surgical treatment, the endovascular interventional modality is currently the preferred therapeutic option, although procedure-related complications and intrastent restenosis after PV intervention remain concerns (Central Illustration).

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REFERENCES

1. Simonneau G, Montani D, Celemajer DS, et al. Haemodynamic definitions and updated clinical classification of pulmonary hypertension. Eur Respir J. 2019;53:1801913.
2. Seferian A, Steriade A, Jais X, et al. Pulmonary hypertension complicating fibrosing mediastinitis. Medicine (Baltimore). 2015;94:e1800.
3. Davis AM, Pierson RN, Loyd JE. Mediastinal fibrosis. Semin Respir Crit Care Med. 2002;23:135-143.
4. Parish JM, Rosenow EC 3rd. Mediastinal granuloma and mediastinal fibrosis. Semin Respir Crit Care Med. 2002;23:135-143.
5. Welby JP, Fender EA, Peikert T, Holmes DR Jr, Bjamason H, Knavel-Koepsel EM. Evaluation of outcomes following pulmonary artery stenting in fibrosing mediastinitis. Cardiovasc Interv Radiol. 2021;44:384-391.
6. Fender EA, Widmer RJ, Knavel Koepsel EM, et al. Catheter based treatments for fibrosing mediastinitis. Catheter Cardiovasc Intervent. 2019;94:878-885.
7. Mathisen DJ, Grillo HC. Clinical manifestation of mediastinal fibrosis and histoplasmosis. Ann Thorac Surg. 1990;54:1053-1057.
8. Hu Y, Qiu JX, Liao JP, Zhang H, Jin Z, Wang GF. Clinical manifestations of fibrosing mediastinitis in Chinese patients. Chin Med J (Engl). 2016;129:2697-2702.
9. Lagerstrom CF, Mitchell HG, Graham BS, Hammon JW Jr. Chronic fibrosing mediastinitis and superior vena caval obstruction from blastomyces. Ann Thorac Surg. 1992;54:764-765.
10. Chatterjee D, Bal A, Singhal M, Vijayaragya R, Das A. Fibrosing mediastinitis due to Aspergillus with dominant cardiac involvement: report of two autopsy cases with review of literature. Cardiovasc Pathol. 2014;23:354-357.
mimicking connective tissue disease. Am J Med Chest 2019;39:651

Mediastinitis and bronchial anthracosis: a case series and a review of the literature. J Int Med Res 2021;49:3000605211010073.

Panagopoulos N, Leivaditis V, Kranitis P, Panagopoulos N, Leivaditis V, Kranitis P, Rokkas CK, Papadopoulos K, Dimitrakos P. Diffuse mediastinal fibrosis and bronchial anthracosis: a systemic immune-mediated disorder. a case series and a review of the literature. Clin Rev Allergy Immunol. 2017;52:446-459.

Morrone N, Gama e Silva VL, Dines DE, Payne WS, Bernatz PE, Pairolero PC. Idiopathic mediastinal fibrosis. J Thorac Cardiovasc Surg 2022;163:3-12.

Christenson ML, Martinez-Jimenez S, Munoz P, Mullington J, Broida J. Idiopathic mediastinal fibrosis: a systemic immune-mediated disorder. a case series and a review of the literature. J Int Med Res 2021;49:3000605211010073.

Ramakantan R, Shah P. Dysphagia due to mediastinal fibrosis in advanced pulmonary tuberculosis. AJR Am J Roentgenol. 1990;154:61-63.

Paragopoulos N, Leivaditis V, Kranitis P, Panagopoulos N, Leivaditis V, Kranitis P, Rokkas CK, Papadopoulos K, Dimitrakos P. Diffuse mediastinal fibrosis and bronchial anthracosis: a systemic immune-mediated disorder. a case series and a review of the literature. J Int Med Res 2021;49:3000605211010073.

Majumdar S, Shub C, Lie JT. Constrictive pericarditis associated with combined idiopathic retroperitoneal and mediastinal fibrosis. Mayo Clin Proc. 1984;59:300-304.

Albers EL, Pugh ME, Hill KD, Wang L, Lloyd JE, Doyle TP. Percutaneous vascular stent implantation as treatment for central vascular obstruction due to fibrosing mediastinitis—a feasibility and safety analysis. Vasc Endovascular Surg. 2018;52:202-206.

Kern R, Peikert T, Eidel E, et al. Bronchoscopic management of airway compression due to fibrosing mediastinitis. Ann Am Thorac Soc. 2017;14:1353-1355.

Hanley PC, Shub C, Lie JT. Constrictive pericarditis associated with combined idiopathic retroperitoneal and mediastinal fibrosis. Mayo Clin Proc. 1984;59:300-304.

Albers EL, Pugh ME, Hill KD, Wang L, Lloyd JE, Doyle TP. Percutaneous vascular stent implantation as treatment for central vascular obstruction due to fibrosing mediastinitis. Circulation. 2011;123:1391-1399.

Fernandez FG, Denlinger CE, Patterson GA, Kreisel D, Krupnick AC. Massive bilateral chylohemorrhages complicating mediastinal granulomatous disease. Ann Thorac Surg. 2009;88:1012-1013.

Shimizu H, Ishikawa S, Yamamoto T, et al. Effectiveness of steroid treatment for hoarseness caused by idiopathic fibrosing mediastinitis: report of a case. Surg Today. 2006;36:382-384.

Cochrane A, Warren R, Mullerworth M, Manolas E. Fibrosing mediastinitis with coronary artery involvement. Ann Thorac Surg. 1991;51:652-654.

Saepena P, Tesar PJ. Mediastinal fibrosis causing myocardial ischemia. Ann Thorac Surg. 2005;80:2368-2370.

Lewczuk J, Ajlan AW, Piszko P, Jaga S, Mikulewicz M, Wrabec K. Electrocardiographic signs of right ventricular overload in patients who underwent pulmonary embolism event(s). Are they useful in diagnosis of chronic thromboembolic pulmonary hypertension? J Electrocardiol. 2004;37:219-225.
58. Rodriguez E, Soler R, Pombo F, Requejo I, Montero C. Fibrosing mediastinitis: CT and MR findings. Clin Radiol. 1998;53:907-910.

59. Rholl KS, Levitt RG, Glazer HS. Magnetic resonance imaging of fibrosing mediastinitis. AJR Am J Roentgenol. 1985;145:259-259.

60. Levitt RG, Glazer HS, Gutierrez F, Moran J. Magnetic resonance imaging of spiral vein graft bypass of superior vena cava in fibrosing mediastinitis. Chest. 1986;90:676-680.

61. Farmer DW, Moore E, Amparo E, Webb WR, Gamsu G, Higgins CB. Calcific fibrosing mediastinitis: demonstration of pulmonary vascular obstruction by magnetic resonance imaging. AJR Am J Roentgenol. 1984;143:1189-1191.

62. Freed BH, Collins JD, Francois CJ, et al. MR and CT imaging for the evaluation of pulmonary hypertension. J Am Coll Cardiol Img. 2016;9:715-732.

63. Kaya H, Rider K, Cho AH, Schwartz A, Alrehaili G, Ahari J. The role of PET scan in monitoring the progression of fibrosing mediastinitis. Clin Imaging. 2016;40:177-179.

64. Takalkar AM, Bruno GL, Makanjula AJ, El-Haddad G, Lilien DL, Payne DK. A potential role for F-18 FDG PET/CT in evaluation and management of fibrosing mediastinitis. Clin Nucl Med. 2007;32:703-706.

65. Lee KY, Yi JG, Park JH, Kim YJ, So Y, Kim JS. Fibrosing mediastinitis manifesting as thoracic prevertebral thin band-like mass on MRI and PET-CT. Br J Radiol. 2007;80:e141-e144.

66. Rossi SE, McAdams HP, Rosado-de-Christenson ML, Franks TJ, Galvin JR. Fibrosing mediastinitis. Radiographics. 2001;21:737-757.

67. Weed HG. Pulmonary “capillary” wedge pressure not the pressure in the pulmonary capillaries. Chest. 1991;100:1138-1140.

68. Wiedemann HP. Wedge pressure in pulmonary veno-occlusive disease. N Engl J Med. 1986;315:1233.

69. Galle N, Humbert M, Vachiery JL, et al. 2015 ESC/ERS guidelines for the diagnosis and treatment of pulmonary hypertension: the Joint Task Force for the Diagnosis and Treatment of Pulmonary Hypertension of the European Society of Cardiology (ESC) and the European Respiratory Society (ERS); endorsed by: Association for European Paediatric and Congenital Cardiology (AEPC), International Society for Heart and Lung Transplantation (ISHLT). Eur Heart J. 2016;37:67-119.

70. Loyd JE, Tillman BF, Atkinson JB, Des Prez RM. Medial fibrosis complicating histoplasmosis. Medicine (Baltimore). 1988;67:295-310.

71. Wheat LJ, Freifeld AG, Kleinman MB, et al. Clinical practice guidelines for the management of patients with histoplasmosis. 2007 update by the Infectious Diseases Society of America. Clin Infect Dis. 2007;45:807-825.

72. Urschel HC Jr, Razzuk MA, Netto GJ, Disiere J, Chung SY. Sclerosing mediastinitis: improved management with histoplasmosis titer and ketoconazole. Ann Thorac Surg. 1990;50:215-221.

73. Takahashi S, Akiyama M, Suzuki K, Otomo K, Takeuchi T. IgG4-related fibrosing mediastinitis diagnosed with computed tomography-guided percutaneous needle biopsy: two case reports and a review of the literature. Medicine (Baltimore). 2018;97:e10935.

74. Kobayashi Y, Ishiguro T, Takaku Y, Kagiyama N, Shimizu Y, Takayanagi N. Clinical features of fibrosing mediastinitis in Japanese patients: two case reports and a literature review. Intern Med. 2021;60(23):3765-3772.

75. Westerny BD, Johnson GB, Maldonado F, Utz JP, Specks U, Peikert T. Targeting B lymphocytes in progressive fibrosing mediastinitis. Am J Respir Crit Care Med. 2014;190:1069-1071.

76. Garrett HE Jr, Roper CL. Surgical intervention in histoplasmosis. Ann Thorac Surg. 1986;42:711-722.

77. Ojeifo O, Gilotra NA, Kemp CD, et al. Mediastinal fibrosis of the pulmonary artery secondary to tuberculosis. Ann Thorac Surg. 2015;100:e49-e50.

78. Hamoud ZT, Rose AS, Hage CA, Knox KS, Rieger K, Kesler KA. Surgical management of pulmonary and mediastinal sequelae of histoplasmosis: a challenging spectrum. Ann Thorac Surg. 2009;88:399-403.

79. Brown ML, Cedeno AR, Edell ES, Hagler DJ, Schaff HV. Operative strategies for pulmonary artery occlusion secondary to mediastinal fibrosis. Ann Thorac Surg. 2009;88:233-237.

80. Doyle TP, Loyd JE, Robbins IM. Percutaneous pulmonary artery and vein stenting: a novel treatment for mediastinal fibrosis. Am J Respir Crit Care Med. 2001;164:657-660.

81. Fontaine AB, Borsa J, Hoffer EK, Bloch RD, So C. Stent placement in the treatment of pulmonary artery stenosis secondary to fibrosing mediastinitis. J Vasc Interv Radiol. 2001;12:1107-1111.

82. Satpathy R, Aguila V, Moheddin SM, Khan IA. Fibrosing mediastinitis presenting as pulmonary stenosis: stenting works. Int J Cardiol. 2007;118: e85-e86.

83. Ponamgi SP, DeSimone CV, Lenz CJ, et al. Catheter-based intervention for pulmonary vein stenosis due to fibrosing mediastinitis: the Mayo Clinic experience. Int J Cardiol Heart Vasc. 2015;8:103-107.

84. Duan Y, Zhou X, Su H, et al. Balloon angioplasty or stent implantation for pulmonary vein stenosis caused by fibrosing mediastinitis: a systematic review. Cardiovasc Diagn Ther. 2019;9:520-528.

85. Cao Y, Duan Y, Su H, et al. Fast pulmonary edema induced by percutaneous transluminal pulmonary angioplasty. J Am Coll Cardiol Intv. 2019;12:e111-e113.

86. Duan YC, Su HL, Zhu Y, et al. Rescue of pulmonary artery intra-stent re-stenosis by unzipping an under-sized stent in an adult patient with fibrosing mediastinitis. Chin Med J (Engl). 2021;134:1880-1882.

87. Vacouby S, Meador M, Mossad E. Lung reperfusion injury in patients after balloon angioplasty for pulmonary artery stenosis. J Cardiothorac Vasc Anesth. 2014;28:502-505.

KEY WORDS: clinical classification, fibrosing mediastinitis, FM, FM dyad, FM triad, interventional therapy, pulmonary hypertension

APPENDIX: For supplemental videos and the search strategy, please see the online version of this paper.