Initial isolated Takayasu's arteritis of bilateral pulmonary artery branches with severe pulmonary hypertension: a case report.

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Abstract

Background: Takayasu’s arteritis is a chronic systematic inflammatory disease. Isolated TA of pulmonary artery branches is very rare.

Methods: The patient received some laboratory tests (including blood routine examination, erythrocyte sedimentation rate, C-reactive protein, etc.) and imageological examinations (including echocardiography, computed tomographic pulmonary angiography, computed tomographic aortic angiography, right-sided heart catheterization, 18F-FDG PET, etc.) for diagnosis.

Results: Computed Tomographic Pulmonary Angiography (CTPA) and Right-sided Heart Catheterization (RHC) showed diffused stenosis in bilateral pulmonary arteries in the case. Pulmonary Artery Systolic Pressure (PASP) measured by echocardiography is 89 mmHg. Besides, an 18F-FDG PET showed the pulmonary arteries and the right ventricle uptake was high.

Conclusion: The symptoms of this patient have been controlled with the treatment of drugs, but pulmonary hypertension is not improved. Further treatment would be a great challenge.

Keywords: Takayasu's arteritis (TA), Pulmonary hypertension, Large vessel vasculitis. Abbreviations: CT: Computed Tomography; CTPA: Computed Tomographic Pulmonary Angiography; ESR: Erythrocyte Sedimentation Rate; FDG-PET: 18F-Fluorodeoxyglucose Positron Emission Tomography; PASP: Pulmonary Artery Systolic Pressure; RHC: Right-sided Heart Catheterization; TA: Takayasu's Arteritis.

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Introduction

Takayasu's Arteritis (TA) is a chronic systematic inflammatory disease, which easily affects aorta and its major branches, including aortic arch and its branches, descending aorta, renal artery, etc. [1]. As aortic secondary branch, pulmonary artery is often involved, but isolated TA of pulmonary artery is very rare. We will report a case of Initial isolated Takayasu's arteritis of bilateral pulmonary artery branches with severe pulmonary hypertension.

Case Description

A 35 years old Chinese woman was presented to the cardiovascular disease clinic with a 7 months history of chest tightness and palpitation after activities. During these 7 months, cough and sputum occurred occasionally and antibiotics were effective. However, chest tightness and palpitation after activities did not improve. And besides, the patient was healthy previously without medical history and relevant family history.

On examination, temperature was 36.7°C, blood pressure 121/76 mmHg in the right arm and 116/82 mmHg in the left, pulse 86 beats per min and room air saturation 98%. Radial and dorsalis pedis artery pulses were equal and palpable bilaterally. There was a systolic murmur in pulmonary valves area. The rest of her physical examination was unremarkable.

On laboratory tests, white blood cells were 11.2 × 10^9/L, C-reactive protein 1 mg/L, Erythrocyte Sedimentation Rate (ESR) 14 mm/h. Platelet count, hemoglobin, levels of electrolytes, and renal and liver function were normal.

Furthermore, an echocardiography showed right heart dilation and pulmonary hypertension (Pulmonary Artery Systolic Pressure, PASP 89 mmHg) and a lung Computed Tomography (CT) was suggestive of pulmonary infraction in middle lobe of right lung and inferior lobe of left lung. A Computed Tomographic Pulmonary Angiography (CTPA) showed varying degrees of stenosis and occlusion in bilateral lobar and segmental arteries (Figure 1). Right-sided Heart Catheterization (RHC) confirmed diffuse stenosis in bilateral pulmonary arteries. We monitored pressures in the distal left pulmonary artery, the proximal left pulmonary artery, the right ventricle, the right atrium and the superior vena cava, which were 24/9 mmHg, 89/17 mmHg, 100/4 mmHg, 9/5 mmHg and 10/6 mmHg separately by RHC. We also noticed no pressure gradient between the right ventricle and the pulmonary trunk. Besides there was no obvious stenosis in the aortic arch and its
branches, the thoracic aorta abdominal aorta, renal arteries, the iliac arteries and the femoral arteries in an examination of computed tomographic aortic angiography (Figure 2). The patient accepted an $^{18}$F-FDG PET, which showed the pulmonary arteries and the right ventricle uptake was high.

**Figure 1.** Total occlusion in the right upper lobar (showed by an arrow), and varying degrees of stenosis in bilateral lobar and segmental arteries.

**Figure 2.** No obvious stenosis in the aortic arch and its branches, the thoracic aorta abdominal aorta, renal arteries, the iliac arteries and the femoral arteries.

Drugs therapy was chosen for this patient to ameliorate of her symptoms and improve pulmonary hypertension. The treatment program included four parts of drugs, which were for pulmonary hypertension (beraprost sodium 120 μg daily for three times, which is a kind of prostaglandins and bosentan 125 mg daily for two times, which is a kind of endothelial receptor antagonists), for heart failure (furosemid 20 mg daily, which a kind of diuretics), for thrombus (aspirin 0.1 g daily, which is a kind of antiplatelet agents) and for TA (prednisone 60 mg daily with tapering weekly, which is a kind of glucocorticoids and cytotoxic agents). The patient has been followed up in our clinic for one and a half year. The symptoms have been improved, but recent echocardiographies showed that PASP still sustained 90mmHg approximately.

**Discussion**

Takayasu’s arteritis is a rare disease of unknown etiology which occurs more often in women under the age of 40 years; it is characterized by medium and large vessel vasculitis [1]. Isolated TA of pulmonary artery is very rare and easy to misdiagnose [2]. The bases of our diagnosis with TA in this patient are: 1. young woman, <40 y old; 2. diffuse stenosis in bilateral pulmonary arteries; 3. suggestion of $^{18}$F-FDG PET [3,4]; 4. no evidence of other vasculitis.

Obviously, pulmonary hypertension of this patient is caused by pulmonary arteritis. Generally, pulmonary artery reconstruction (including balloon dilatation, stenting and bypass) can be used to improve pulmonary hypertension and patient’s symptoms [2,5]. But for this patient, pulmonary artery reconstruction doesn’t apply to the diffuse stenosis in bilateral pulmonary arteries. There was no cure for TA. Improvement of symptoms and delaying progression of TA are the most important targets of treatment. For TA with pulmonary hypertension, the targets of treatment are to reduce pulmonary pressure, control right heart failure, delay progression of TA and improve patient’s prognosis. Some kinds of drugs are suggested to be used for pulmonary hypertension, including prostaglandins, endothelin receptor antagonists, guanylate cyclase agonists, and type 5 phosphodiesterase inhibitor [2,6]. If feasible, pulmonary artery reconstruction is an option [5]. For another, treatment of TA is even more important, which includes glucocorticoids and immunosuppressants [1,7]. Glucocorticoids in combination with immunosuppressants should been considered for refractory TA. For this patient, prednisone combining with cytoxan has been used for controlling TA. By tapering, dose of prednisone has been reduced from 60 mg to 5 mg daily.

The symptoms of this patient have been controlled with the treatment of drugs, but pulmonary hypertension is not improved, and her PASP is still 90mmHg approximately. If pulmonary hypertension of the patient worsens, we will have no effective treatment in the condition of pulmonary artery reconstruction to be inapplicable. Recently, biological agents, such as tumor necrosis factor-alpha inhibitors, interleukin-6 inhibitor, tocilizumab and abatacept, maybe an optional therapy [8,9]. But more randomized controlled trials are still needed to verify their effectiveness. Moreover, lung transplantation, as the last resorts, maybe effective in this patient [6]. However, there is no report about lung transplantation to treat TA with severe pulmonary hypertension currently in the world.
Informed Consent

Informed consent was obtained from the patient for publication of this case report and any accompanying images. The study was accepted by the Ethics Committee of the hospital and they encouraged publishing the article (Chinese scientific medical ethic review No.2017-108-01).

References

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