Pulmonary arterial hypertension in systemic sclerosis: diagnostic pathway and therapeutic approach

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CASE HISTORIES

Case 1
A 63 year old white woman was admitted to hospital in February 2000 owing to rapidly increasing dyspnoea on exertion (NYHA III), chest pain, and fever. The presence of sclerodactyly with loss of distal parts of the fingers, and megacapillaries at capillaroscopy, allowed the diagnosis of limited cutaneous systemic sclerosis (fig 1). Physical examination also disclosed crackles in the lower lobes of the lungs, systolic murmur with loud pulmonary second sound, and feet oedema.

Laboratory tests included high erythrocyte sedimentation rate (145 mm/1st h), C reactive protein (2530 mg/l, normal <100 mg/l), presence of antinuclear antibodies (1/1280 and speckled pattern on indirect immunofluorescence), mild anaemia and thrombocytopenia (platelet count of 115 x 10^9/l), decreased arterial oxygen and carbon dioxide pressures. A chest x ray examination showed cardiomegaly, dilatation of the main pulmonary artery (fig 1), centralisation of pulmonary vascular flow, and reticular opacities in the lower parts of the lungs. Electrocardiography (ECG) at admission did not show any significant abnormalities. Doppler echocardiography disclosed a pulmonary artery systolic pressure (PASP) of 86 mm Hg, suggesting a severe pulmonary artery hypertension (PAH). Pulmonary function tests showed a reduction of forced vital capacity (FVC, 45% of predicted), and a high resolution chest computed tomography (CT) scan showed interstitial lung disease (ground glass and interstitial fibrosis) (fig 1). Furthermore, the patient developed pulmonary embolism, confirmed with perfusion scintigraphy (fig 1), with an increase in pulmonary vascular flow, and reticular opacities in the lower parts of the lungs. Electrocardiography (ECG) at admission did not show any significant abnormalities. Doppler echocardiography disclosed a pulmonary artery systolic pressure (PASP) of 86 mm Hg, suggesting a severe pulmonary artery hypertension (PAH). Pulmonary function tests showed a reduction of forced vital capacity (FVC, 45% of predicted), and a high resolution chest computed tomography (CT) scan showed interstitial lung disease (ground glass and interstitial fibrosis) (fig 1). Furthermore, the patient developed pulmonary embolism, confirmed with perfusion scintigraphy (fig 1), with an increase in pulmonary vascular flow up to 115 mm Hg. Anticardiolipin antibodies were found to be positive.

The patient was treated with warfarin (international normalisation ratio (INR) between 2 and 3) and a high dose of calcium channel blocker, (nifedipine 120 mg/day), an angiotensin converting enzyme (ACE) inhibitor (perindopril 4 mg/day), diuretics (furosemide 20 mg/day, spironolactone 100 mg/day), and oxygen (1–2 litres/min). Owing to interstitial lung disease, confirmed at bronchoalveolar lavage (no infection detected, total cell number 200 x 10^9/l, neutrophils 4%, eosinophils 0%, lymphocytes 23%, macrophages 73%), she also received cyclophosphamide (CYC) pulses (1.0 g/m2/ month for 6 months) and prednisone 15 mg/day. Right cardiac catheterisation confirmed severe PAH with a positive test for resistance responders: mean right atrial pressure = 20 mm Hg at rest and 19 mm Hg after the epoprostenol test, PASP 56 mm Hg at rest and 50 mm Hg after the epoprostenol test, mean pulmonary capillary wedge pressure 13 mm Hg, reduced cardiac output of 2.50 litres/min that became 4.40 litres/min after the epoprostenol test, and pulmonary vascular resistance of 22.4 U/W at rest and 11.3 U/W after the epoprostenol test.

DISCUSSION

Remarks on case 1

Key questions
- Are any further investigations necessary?
- Is it appropriate to initiate anticoagulation?
- Was O2 necessary?

Case 2
In April 2002 a 69 year old white woman with diffuse systemic sclerosis (SSc), diagnosed in 2001, was admitted to hospital after a 6 month history of dyspnoea on exertion (NYHA II/III), fatigue, chest pain, and dry cough. Physical examination showed diffuse telangiectasias, cyanosis of the lips and fingers, mild oedema of the feet, crackles in lower lobes, pansystolic murmur (4/6 L) on the apex, with accentuation of the pulmonary component of the second heart sound; the liver was enlarged with mild ascites.

Laboratory testing showed an erythrocyte sedimentation rate of 31 mm/1st h; prothrombin time 20 seconds (normal <15 seconds), and activated partial prothrombin time 41 seconds ( normal 26–36 seconds). Antinuclear antibodies were positive (1/5120). A chest x ray showed cardiomegaly without interstitial involvement. FVC was normal (84%), with a significant decrease of carbon monoxide transfer factor TCO (36%), severe hypoxaemia and respiratory alkalosis (partial pressure of oxygen 62.6 mm Hg; partial pressure of carbon dioxide 28 mm Hg; HCO3^− 20.7 mmol/l; arterial oxygen saturation 90%). Chest high resolution CT showed an enlarged right atrium and ventricle, dilatation of the main pulmonary artery, centralisation of pulmonary vascular flow with pulmonary congestion, bilateral pleural effusion, and pericardial effusion. Bibasilar mild ground glass opacities were detected. A Doppler echocardiogram showed enlargement of the right heart with tricuspid insufficiency, paradoxical movement of the cardiac septum, mild pericardial effusion, and raised PASP (70 mm Hg).

Right cardiac catheterisation confirmed severe PAH with a positive test for resistance responders: mean right atrial pressure = 20 mm Hg at rest and 19 mm Hg after the epoprostenol test, PASP 56 mm Hg at rest and 50 mm Hg after the epoprostenol test, mean pulmonary capillary wedge pressure 13 mm Hg, reduced cardiac output of 2.50 litres/min that became 4.40 litres/min after the epoprostenol test, and pulmonary vascular resistance of 22.4 U/W at rest and 11.3 U/W after the epoprostenol test.

ABBREVIATIONS

ACE, angiotensin converting enzyme; CYC, cyclophosphamide; ECG, electrocardiography; FVC, forced vital capacity; PAH, pulmonary artery hypertension; PASP, pulmonary artery systolic pressure; SSC, systemic sclerosis
Facts in this case
- Pulmonary function tests indicated a "restrictive pattern" with a reduction of the FVC.
- A chest CT scan indicated interstitial fibrosis and alveolitis.
- Doppler echocardiography disclosed severe pulmonary hypertension.
- The immune profile identified anticardiolipin antibodies that together with thrombocytopenia and pulmonary thromboembolism allowed the diagnosis of antiphospholipid syndrome.
- ECG did not show any modifications. In fact, ECG may be an inadequate tool for ruling out the presence of pulmonary hypertension, in particular in early phases.
- A standard chest x ray showed typical signs of PAH such as "dilated pulmonary arteries" and "increased cardiothoracic ratio".
- This is a case of limited cutaneous SSc. In such a case, when the patient complains of breathlessness, interstitial involvement or vascular lung disease, or both, should be suspected, both leading to PAH and respiratory failure. In this case, PAH was secondary to interstitial lung disease, as indicated by a reduced FVC and the CT scan and was exacerbated by a pulmonary thromboembolic event due to antiphospholipid syndrome. The relationship between PAH and interstitial disease is confirmed by the results of the treatment with CYC, which significantly improved the FVC and PASP.

Comment
This case shows that some important investigations, that might have added significant information for a more precise management, are missing:
- Carbon monoxide transfer factor (TLCO) is missing. It measures gas exchange at the level of the alveolocapillary membrane and identifies impairment of the transfer of CO that may be due to interstitial fibrosis or vascular disease. In particular, the FVC/TLCO ratio may be helpful in a differential diagnosis for PAH complicating SSc, with an FVC/TLCO ratio >1.4 indicating a vascular pulmonary hypertension. Accordingly, an isolated decrease of TLCO, with a normal FVC, is a reliable marker for the development of PAH in SSc. In SSc, a severe reduction in TLCO (<40% of predicted values) is indeed an adverse prognostic factor. For this reason, pulmonary function tests with TLCO are a fundamental investigation in assessment of the patient with SSc.
- Right heart catheterisation with vasodilator test. When the clinical assessment and the echo Doppler measurement indicate PAH then right heart catheterisation is mandatory. It allows the "mean pulmonary artery pressure" to be determined. Also, assessment of other indices such as right atrial and ventricular pressures, pulmonary capillary wedge pressure, and cardiac output is essential for differential diagnosis of PAH secondary to myocardial disease and has important prognostic value. Catheterisation makes it possible to verify if PAH is postcapillary, which is seldom the case in patients with SSc. A vasodilator test is necessary to evaluate the response of pulmonary vasculature in order to identify those patients who may benefit from the use of calcium channel blocker or other drugs, although the frequency of true responders is low in scleroderma and the benefits of high dose calcium channel blockers must be balanced against potential toxicity.
- Helical spiral CT angiography is indicated for further diagnosis and localisation of thromboembolic disease.
Dyspnoea on exertion, hilar congestion at chest-x-ray, hypoxaemia

Clinical suspect of PAH

Negative

Heart echo colour Doppler

Increased PASP

To be confirmed with

Catheterisation

PFT

HRCT

Negative

FVC

TLCO

PAH

ILD

= or /

FVC

TLCO

Vasoreactivity positive

Vasoreactivity negative

Anticoagulation

CCB

Prostanoids

Bosentan

O2

Recurrent

Comment

In this case, some investigations are still missing:

- Anticardiolipin antibodies, β2-glycoprotein I, and lupus anticoagulant were not determined. Note that in this case the activated partial prothrombin time was prolonged; this may indicate an underlying antiphospholipid syndrome, as in case 1.
- Perfusion scintigraphy of the lungs/helical CT angiography is mandatory to exclude thromboembolic disease.
- An exercise test (6 minute walk test) may be helpful as in case 1.

In this case, basal therapy is needed: first of all it is necessary to anticoagulate (reach an INR of between 2 and 3), use oxygen (2 litres/min) that may be highly helpful, as well as diuretics. If during catheterisation the patient shows a positive response to vasodilatation, high dose calcium channel blockers may be used. If these fail to lower PAH, in patients in NYHA class III, bosentan or prostanoids may be used; in NYHA class IV, epoprostenol alone or in combination with bosentan may then be employed. When all these treatments fail to control PAH, atrioseptostomy or, eventually, heart-lung transplantation may be considered.

THE LESSONS

- In SSC, PAH is a severe complication that may rapidly lead to the patient’s death; the early identification of PAH is mandatory to design a management plan to reduce pulmonary pressure (see fig 2).
- Cardiac Doppler echocardiography should be used regularly, including in asymptomatic patients; if a rise in pulmonary pressure is detected or, despite a negative Doppler echocardiography, a strong clinical suspicion for PAH exists, right catheterisation with vasoreactivity tests is mandatory to confirm PAH and to design the treatment.
- In cases of non-responsiveness to pharmacological vasoactive treatments, a surgical approach (atrio-septotomy) should be considered.
- In the presence of alveolitis, CYC treatment may be useful in order to control interstitial fibrosis (see fig 2), which contributes to the genesis of PAH.

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