Platypnoea–orthodeoxia syndrome

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Abstract

Platypnoea–orthodeoxia is a rare syndrome of postural hypoxaemia accompanied by breathlessness. The predominant symptom, dyspnoea induced by upright posture, can be debilitating and difficult to discern without thorough evaluation of the patient’s pattern of dyspnoea. The precise cause of the syndrome is unclear but patients develop right to left intracardiac shunting in the presence of normal right sided cardiac pressures. Initially, patients should have confirmation of orthostatic desaturation by erect and supine pulse oximetry. However, definitive diagnosis of an orthostatic intracardiac shunt is most readily established by echocardiography. The use of echocontrast with postural manoeuvres may facilitate the diagnosis. The treatment of choice is surgical closure of the intracardiac (usually interatrial) communication, which may result in dramatic symptomatic and haemodynamic improvement. Three cases (a 27 year old man and two women aged 63 and 72 years) are described that exemplify the presentation of this syndrome, and reflect the varied management strategies and outcomes of this condition. (Heart 2000;83:221–223)

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Platypnoea–orthodeoxia is an uncommon syndrome of dyspnoea induced by upright posture, with associated arterial hypoxaemia, which is subsequently relieved by recumbency.¹ The syndrome occurs when there is right to left shunting of blood, usually via an interatrial communication, in the presence of normal pulmonary artery pressure. This is an unusual situation given that most cases of significant right to left intracardiac shunting of blood are associated with increased right sided cardiac pressures.² The three cases described here exemplify the presentation of this syndrome, and reflect the varied management strategies and outcomes of this condition.

Case 1

A 27 year old man with a known atrial septal defect (ASD) presented with 24 hours of extreme dyspnoea at rest with a history of four months of worsening exertional dyspnoea and lethargy. The ASD had been diagnosed at 9 years of age but the patient had failed to attend follow up care. He had no other significant medical history but was a smoker with a 12 pack-year history of consumption. He had no symptoms suggestive of parenchymal lung disease, pulmonary infections, obstructive sleep disorders or thromboembolic disease. On physical examination his temperature was 37.5°C, he had a regular pulse at 88 beats/min, respiration rate 24/min, and blood pressure was 110/70 mm Hg. There was central cyanosis and digital clubbing with two subungual haemorrhages. Lung fields were clear to auscultation, and cardiac examination revealed fixed wide splitting of the second heart sound. There were no regurgitant heart murmurs or palpable precordial impulses. With the patient breathing room air, oxygen saturation was 92% supine and 84% in the upright position. Subsequent investigation revealed haemoglobin of 183 g/l, normal pulmonary ventilation perfusion scan, and normal respiratory function tests. Chest roentgenogram revealed a generalised increase in pulmonary vascular markings without dilatation of the main pulmonary artery or enlargement of the right atrium and ventricle. His ECG showed a right axis deviation with right bundle branch block. Blood cultures remained negative to day 5.

Echocardiography (transthoracic and subsequently transoesophageal) with bubble contrast study showed a significant right to left shunt at the atrial level through a secundum defect (fig 1). The calculated pulmonary artery pressure was normal (mean 14 mm Hg with a peak systolic pulmonary artery pressure of 19–23 mm Hg) with a moderately dilated right atrium and ventricle. Left ventricular size and function were preserved. There was a broad jet of inferior vena caval blood directed towards the ASD. No vegetations were seen.

Subsequent right and left heart catheterisation showed a large ASD (secundum type) with bidirectional shunting of blood. The pulmonary artery pressures were normal (22 mm Hg/8 mm Hg, mean 14 mm Hg) as was the coronary artery anatomy. The patient underwent open surgical closure of the ASD with a patch applied to the 3 cm diameter defect. He had an uneventful recovery and remained asymptomatic at four years’ follow up. Furthermore, the clubbing resolved as did the hypoxaemia.

Case 2

A 63 year old woman presented with a six week history of worsening exertional dyspnoea with
reduction in exercise tolerance to less than 50 metres. She denied any associated anginal symptoms and remarked how the breathlessness abated quickly with recumbency. Six months previously, she had undergone coronary artery bypass grafting for anginal symptoms refractory to medical treatment. Her postoperative course had been complicated by a right occipital and thalamic infarct (confirmed by magnetic resonance imaging) with a residual left homonymous hemianopia, left hemisensory disturbance, and significant tremor with gait abnormality. On physical examination, values for blood pressure, pulse, and respiratory rate were within normal limits. There was no cyanosis or digital clubbing and heart sounds were of normal intensity with physiological splitting and without audible murmur. The lung fields were clear to auscultation and the remainder of the examination was normal apart from the neurological abnormalities. Oxygen saturation was 98% with the patient supine and 74% while standing. Routine haematology and biochemical blood analysis were normal, as was the resting ECG and chest roentgenogram. Arterial blood gases performed on room air showed a significant hypoxaemia (PO₂, 68 mm Hg) with an alveolar to arterial gradient of greater than 40 mm Hg. Pulmonary ventilation perfusion scan, non-invasive peripheral vascular studies of the lower limb venous system, and pulmonary function tests were normal.

Transoesophageal echocardiography showed a mobile interatrial septum with no definitive defect. Echocardiography confirmed the presence of right to left shunting of blood with the appearance of left sided bubbles soon after injection. The calculated pulmonary artery pressure was normal (mean 12 mm Hg, peak systolic pulmonary artery pressure 19–23 mm Hg) and left ventricular systolic function was preserved. There was moderate tricuspid regurgitation with a broad jet of central regurgitant flow directed at the interatrial septum. Subsequent right atrial contrast confirmed pulsatile shunting from the right atrium to the left atrium across the superior limb of the foramen ovale membrane. Coronary angiography revealed no stenosis in the graft vessels.

Following consultation with a cardiac surgeon, the patient declined any surgical intervention given her previous complicated cardiac surgery. Her symptoms have not progressed in two years of follow up, however the ability to perform many activities of daily living have been significantly curtailed by exertional and postural dyspnoea.

Case 3
A 72 year old woman with a 10 year history of schizophrenia, for which she had been an inpatient for one month with acute psychosis, was noted by a duty nurse to appear breathless and cyanosed on walking to the shower. Her oxygen saturation was 76% on room air at the time of this observation and she was returned to bed. Review by the duty medical officer showed values for blood pressure, pulse, and respiratory rate were within normal limits. Oxygen saturation in the supine position was 98%. Cardiorespiratory examinations were unremarkable apart from a soft 2/6 ejection systolic murmur along the left sternal border, which radiated into the carotids. The patient was non-communicative owing to her psychosis and as such no symptoms were obtainable. Routine haematological and biochemical blood analysis were normal as was her resting ECG. Chest roentgenogram revealed a tortuous proximal aorta with clear lung fields. Computed tomography pulmonary angiography was normal apart from a 5 cm diameter aneurysm of the ascending aorta. Syphilis serology was negative. Non-invasive peripheral venous studies of the lower limbs were also negative.

The observed breathlessness and cyanosis were reproducible on subsequent mobilisation and consequently the patient underwent transthoracic echocardiography with bubble contrast.
Platypnoea–orthodeoxia syndrome is a rare condition characterized by dyspnea and arterial hypoxemia resulting from right to left shunting through a patent foramen ovale. This shunting occurs in about 25% of the general population and is more common in patients with a history of lung disease such as severe chronic obstructive lung disease, pulmonary hypertension, or aortic stenosis. Various theories have been advanced to explain why patients with a patent foramen ovale may develop this syndrome. Explanations include preferential arterial inflow in the erect position, unequal diastolic compliance between the right and left sides of the heart, and positional dyspnea and hypoxemia out of proportion to any underlying cardiopulmonary disease.

In case 1, this can result in a dramatic symptomatic improvement, as well as eliminating the site of a possible paradoxical embolus. However, in some patients—and before surgical correction—right and left cardiac catheterisation will be required.

The treatment of choice is surgical closure of the intracardiac communication. As demonstrated in case 1, this can result in a dramatic symptomatic improvement, as well as eliminating the site of a possible paradoxical embolus. However, in some patients, such as case 2, surgery carries a significant risk.

Platypnoea–orthodeoxia is a rare condition with the potential for surgical cure. Non-invasive investigations can readily establish the diagnosis and should be performed when this condition is suspected.