Progressive primary pulmonary hypertension in a patient born at high altitude

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SUMMARY A girl, born at high altitude, subsequently developed pulmonary hypertension which did not improve on return to sea-level and proved fatal at the age of 19 years.

The clinical and necropsy features were those of primary pulmonary hypertension, including a plexiform arteriopathy. There is a possibility of hypoxia in early life triggering progressive pulmonary vascular changes.

Although the pathogenesis of primary pulmonary hypertension remains obscure, it has long been established that exposure to the hypoxic stimulus of high altitude produces an increase in pulmonary arterial pressure in man and this increase may well be completely reversible.1 2 The characteristic vascular changes of primary pulmonary hypertension are rarely seen in the indigenous high altitude dwellers of the Andes, and there appears to be considerable variation in the susceptibility of the individual to develop pulmonary hypertension at altitude.3 4 Several cases of pulmonary hypertension have been described in young children whose parents had recently settled at altitude in an area near Leadville and Climax in Colorado, and it was found that they improved on moving to a lower altitude and did not exhibit the usual progressive course of primary pulmonary hypertension.5

In this paper, we describe the clinical presentation and necropsy findings of progressive pulmonary hypertension in a girl who was exposed to high altitude in the Peruvian Andes for the first 15 months of life.

Case report

HISTORY

The patient, a 16-year-old girl, presented in 1974 with a history of tiredness and progressive exertional dyspnoea of three years' duration. Her history during early life was of particular interest.

She had been born of British parents at an altitude of 13 000 feet in the Peruvian Andes. She had had a normal full-term delivery but at the age of 6 months she was failing to gain weight at a normal rate. At the age of 1 year, it was noted that her appetite and rate of weight gain improved while on holiday at sea-level. She returned to altitude and at the age of 14 months had an episode of acute respiratory distress. She returned to sea-level the following month and her general well-being improved almost immediately. She came to Scotland with her parents at the age of 2 years. Her overall growth and development were slow during childhood, but she caught up with her peers at puberty. At no time did she take oestrogen-containing preparations or slimming drugs.

CLINICAL FINDINGS

She was of normal stature and there was no central cyanosis. A giant a wave in the jugular venous pulse was present and there was an obvious right ventricular heave. Auscultation disclosed an accentuated pulmonary second sound with a pulmonary ejection click and a grade 4/6 pulmonary early diastolic murmur.

INVESTIGATIONS

The electrocardiogram showed sinus rhythm with gross right ventricular hypertrophy and right axis deviation. The chest x-ray film showed prominence of the main pulmonary artery. Right heart catheterisation was carried out and the pulmonary artery pressure was found to be 125/55 mmHg (mean 90 mmHg) at rest. The right ventricular pressure was 125/0 mmHg and the mean pulmonary wedge pressure 5 mmHg. No shunt was detected on oximetry.
A provisional diagnosis of primary pulmonary hypertension was made and she was started on long-term Warfarin to prevent pulmonary arterial thrombosis. Her symptoms, however, slowly deteriorated, and within two years she was breathless on minimal exertion. Right heart catheterisation was repeated in 1977 and the findings were similar to those of 1974, the right ventricular systolic pressure being 150 mmHg. The diagnosis remained unchanged. Later in 1977, despite the addition of digoxin and propranolol, she died suddenly, aged 19 years, three years after presentation.

NECROPSY FINDINGS

Gross appearances
The heart weighed 400 g and showed severe hypertrophy of the right ventricle which measured approximately 35 mm in thickness 1 cm below the pulmonary valve. The tricuspid valve ring was dilated and the pulmonary valve appeared regurgitant. Both coronary arteries were normal in distribution and were patent throughout their length. The main pulmonary arterial trunk was thick-walled, severely dilated, and moderately atheromatous. There was no evidence of a septal defect and the ductus arteriosus was closed. The liver weighed 1200 g and had the typical “nutmeg” appearance of long-standing venous congestion. Nothing of significance was noted in the other organs. The cause of death was acute right ventricular failure secondary to pulmonary hypertension.

Microscopical examination
The most striking findings were florid hypertensive changes involving the pulmonary vasculature. Many of the muscular pulmonary arteries (100 to 1000 μm in external diameter) showed prominent hypertrophy with crenation of the internal elastic lamina and several of them were occluded by intimal fibroelastic tissue (Fig. 1). In addition, plexiform and occasional angiomatoid lesions were present (Fig. 2). The elastic pulmonary arteries (1000 μm in diameter) were atheromatous. Of particular note was the similarity of the elastic fibre pattern of the pulmonary trunk to that of the aorta. An ischaemic infarct was present in the upper lobe of the right lung. The carotid bodies showed no hyperplasia of the major cellular components.

Discussion
It is probable that the disease known as primary pulmonary hypertension will prove to have more than one cause. A familial occurrence has been well established and impaired fibrinolytic mechanisms have been documented in one family with the disease, suggesting that recurrent miliary emboli...
Proliferation of endothelial cells and fibrous tissue were associated with fibres to thought of plexiform arteriopathy, from finding suggests hypertension. Pulmonary glandin.'6

Such changes as in the fetus and branching, as in the aorta. Such a finding suggests that pulmonary hypertension had persisted from birth.17 In addition, there was florid plexiform arteriopathy, as seen in primary pulmonary hypertension. Such changes are also associated with intracardiac shunts, persistent ductus arteriosus, and hepatic cirrhosis, all of which were excluded in this case.

Although the vascular lesions of most patients with primary pulmonary hypertension are consistent with a vasoconstrictive pathogenesis,18 the disease appears to be rare among the Quechua people who have lived in the Andes for countless generations. Khoury and Hawes4 in 1963 described 11 cases of primary pulmonary hypertension in children of recent settlers in an area near Leadville and Climax, Colorado, at an altitude above 10 000 feet. Two of the children died, but the remaining nine recovered when they were moved to lower altitude. Interestingly, the plexiform arteriopathy particularly associated with primary pulmonary hypertension was not described in these cases.

The present report records the occurrence of primary pulmonary hypertension in a young girl who was born at an altitude of 13 000 feet, and it is suggested that exposure to this hypoxic environment at birth may have been responsible for the development of pulmonary vascular changes which progressed to the florid plexiform vascular changes usually associated with primary or idiopathic pulmonary hypertension.

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References


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