Type I aortopulmonary window presenting with very early onset Eisenmenger’s syndrome

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DESCRIPTION
A 2-year-old child with a history of recurrent chest infection early in infancy presented with clubbing, cyanosis, oxygen saturation of 86% in room air, left parasternal heave, short ejection systolic murmur at left second space and loud pulmonary component of second heart sound suggestive of severe pulmonary hypertension.

X-ray revealed cardiomegaly with pulmonary artery dilation and peripheral vasoconstriction (figure 1). Echocardiography revealed biventricular hypertrophy with moderate tricuspid regurgitation and pulmonary artery systolic pressure of 90 mm Hg. A large aortopulmonary (AP) window (1 cm) was demonstrated in the parasternal short axis view with bidirectional shunting (figures 2–4, videos 1–3). Cardiac catheterisation to assess for the reversibility of the shunt revealed that there was irreversible pulmonary hypertension, hence the child was managed medically with medications to decrease pulmonary arterial pressure like bosentan.

Aortopulmonary window is a rare disease with an incidence of 0.1% and is classified by Mori et al into three types. Type I, which is the most common type and seen in our case, is described as a defect midway between the semilunar valve and the pulmonary bifurcation. Echocardiography is the investigation of choice and can accurately diagnose and prognosticate the disease and cardiac catheterisation is usually required for ambiguity in diagnosis or for the assessment of shunt reversibility.

Survival depends on the size of the defect and the pulmonary vascular resistance. In case of large defect, if left untreated, the majority of patients die in infancy with heart failure and only few survive.
till adolescence or adulthood. Early closure of defect either surgically (preferred in most cases) or by device (small restrictive defect) is the treatment of choice and outcome after successful and early closure is excellent.

Competing interests None.

Patient consent Obtained.

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Algorithms

Aortopulmonary window is a rare congenital heart disease which predisposes to Eisenmenger’s syndrome much earlier than other left to right shunts. A certain percentage of patients are prone to directly transform to Eisenmenger’s without a preceding stage of heart failure due to large left to right shunt.

The cases require early diagnosis preferably with the help of a careful echocardiogram in association with other clinical clues and easily available investigations like X-ray and pulse oximetry. Cardiac catheterisation could be performed preoperatively to assess for shunt reversibility.

Early surgical (preferred) or device closure (if suitable) of this defect is recommended before development of the irreversible pulmonary vascular obstructive disease.

REFERENCES