Successful Reversal of Pulmonary Hypertension in Eisenmenger Complex

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We present a case of a 19-year old female with systemic pulmonary artery (PA) pressure due to a congenital ventricular septal defect (VSD) and atrial septal defect (ASD). She was pink at rest and cyanotic on exercise. Lung biopsy revealed grade IV pulmonary vascular changes. As a preliminary step PA was banded to increase right-to-left shunt and decrease aortic (Ao) saturation with consequent decrease in PA saturation. After one year, when she was no longer cyanotic, even on exercise, lung biopsy revealed total regression of pulmonary vascular changes. As a definitive procedure VSD and ASD were closed and PA was debanded. Cardiac catheterization one week postoperatively showed PA pressure to be 50% of systemic pressure. We postulate that reversal of pulmonary vascular changes were due to lowered PA saturation. We further believe that lower PA pressure could have contributed to this regression of pulmonary vascular changes. We performed the same procedure in six more patients with similar positive clinical response. This new concept brings renewed hope to many children who otherwise are candidates for heart lung transplantation.

Case Report

We present a case of a 19-year-old female with severe pulmonary hypertension due to a congenital VSD, ASD with dextrocardia. She was pink at rest and cyanotic on exercise. Cardiac catheterization showed an aortic (Ao) saturation = 97%. PA-saturation = 87%. AoP = 130/80mmHg (mean=96mmHg); pulmonary artery pressure (PAP) = 125/
45mmHg (mean=70mmHg) unresponsive to increased inspired PO₂. Lung biopsy showed grade IV pulmonary vascular changes (fig. 1). The PA was banded to decrease aortic saturation down to 60% (room air). Thus increasing right-to-left shunt, resulting in low PA saturation (20 to 30%).

The patient left the operating room with Ao saturation of 75% and the PA saturation of 30% on room air. Systolic PA pressure was 50% of systemic. During the following year, Ao saturation increased to normal levels indicating a decrease in pulmonary vascular resistance. Cardiac catheterization at that time showed: Ao sat. = 96%, PA sat. = 88%; AoP = 127/70mmHg (mean = 90mmHg) and PAP = 68/16mmHg (mean = 41mmHg). Lung biopsy showed total regression of previous lesions (fig. 2). Subsequently PA was debanded and VSD and ASD closed. Before closing the chest, AoP was 120/80mmHg (mean = 90mmHg) and PAP was 50/20mmHg (mean = 31mmHg). At the time of discharge cardiac catheterization showed similar results.

**Discussion**

It is possible that end stage congenital left-to-right shunt lesions with fixed PA hypertension can be corrected by a two-stage procedure. We believe that the reason for the regression of lung lesions is due to lowering O₂ saturation of the pulmonary artery or in combination with lower PA pressure. This lower PA saturation results in dilatation of the pulmonary vascular bed and a decrease in pulmonary vascular resistance, causing regression of fixed pulmonary lesions. We believe that high oxygen content in the PA is harmful to the arterioles, causing increased PA vascular resistance. We also postulate that PA vascular tree has O₂ sensors that mediate vascular response, contrary to the response to oxygen at alveolar level. That is: oxygen through the alveoli decreases PA vascular resistance and oxygen given through the PA increases PA vascular resistance. Then we will be able to treat also primary PA hypertension (with drugs to regulate these O₂ sensors). Applying this new concept may avoid heart-lung or lung transplantation for this special indication in the future.

We have performed the first stage of this procedure in six more patients with similar positive outcome and they are waiting for the definitive procedure.

**References**
