Neonatal repair of persistent fifth aortic arch and aortic coarctation

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There are 2 main clinical forms of persistent fifth aortic arch (PFAA), a systemic to systemic connection; that is, the double lumen aortic arch first described by Van Praagh in 1969, and the systemic to pulmonary connection variety.

We report a neonatal case of PFAA with severe hypoplasia of the fourth arch, and coarctation of both the fourth and fifth arches. The discrete coarctation segment relaxed with prostaglandin E1 (PGE1). We will review this patient and the literature specifically discussing PGE1-sensitive tissue.

**PATIENT AND METHODS**

Informed consent was obtained from the patient’s parent for this publication. Pediatric cardiology consultation was requested for a 2.9-kg term male neonate who failed a critical congenital heart disease screening test. The patient had weak femoral pulses and echocardiography showed normal intra-cardiac anatomy. Suprasternal imaging showed 2 arches connecting the ascending and descending aorta in superoinferior orientation on the same side of the trachea. The superior arch was severely hypoplastic (2 mm; z score, –6.3), the ductus arteriosus was patent but small, and Doppler interrogation showed typical coarctation flow pattern with mild gradients across both arches. The isthmus of the fourth arch was 2 mm (z score, –6.3), the ductus arteriosus was patent but small, and Doppler interrogation showed typical coarctation flow pattern with mild gradients across both arches. The isthmus of the fourth arch was 2 mm (z score, –6.3), and the fifth arch was 1.7 mm. After PGE1 administration, the ductus arteriosus was larger and there was normalization of coarctation flow pattern presumably from relaxation of PGE1-sensitive tissue in the terminal segment of both arches, the isthmus of the fourth and fifth arches now measured 2.5 and 2.7 mm, respectively. Computerized axial tomography scan confirmed the diagnosis (Figure 1). The patient underwent elective surgical repair via a midline sternotomy and selective cerebral perfusion. Intraoperative findings confirmed the diagnosis. All presumed ductal tissue was resected from the fourth and fifth arches, and they were connected in a side-to-side anastomosis forming 50% to 60% of the posterior circumference of the repaired arch. The posterior native tissue was connected to the descending aorta in an interdigitating fashion with anterior augmentation of this repair using a pulmonary homograft patch (Video 1). A small atrial septal defect was also closed. The patient had an uncomplicated postoperative course and the infant was discharged home in 2 weeks. At 1-year follow-up, the patient continues to have an excellent surgical result, normal femoral pulses, and a peak velocity across the arch of <1.5 m/s without diastolic continuation.

**DISCUSSION**

Our patient had PFAA with severe hypoplasia of the fourth arch, and juxtaductal coarctation of the fourth and fifth arches. PGE1 administration relaxed tissue in the isthmus of both arches and the Doppler waveform normalized suggesting ductal tissue. Our findings are in keeping with the recognition of PGE1-sensitive tissue in PFAA in the literature. Carroll and colleagues and Binsalamah and colleagues have shown histologically that the PFAA contains ductal tissue.

Several different surgical techniques have been successfully utilized to repair arch obstruction/interruption in the setting of PFAA with and without incorporating tissue from the PFAA. Both Lambert and colleagues and...
Binsalamah and colleagues\(^3\) successfully incorporated PFAA tissue into the surgical repair with good immediate and short-term results. Our institutional bias is to repair all patients with hypoplastic arch using patch aortoplasty techniques extrapolated from the Norwood experience. By utilizing the PFAA tissue, we were able to achieve 50\% to 60\% of the circumference composed of native tissue with growth potential compared with <25\% if the PFAA tissue was not included.

CONCLUSIONS
There is both clinical and histological evidence to show that ductal tissue migrates into the PFAA just like it does in coarctation of the fourth arch. The biggest clinical challenge is complete resection of ductal tissue, and if this can be achieved, it may be reasonable to incorporate PFAA tissue in the repair for its growth potential. Longer-term follow-up and experience from additional case reports will likely provide future guidance given the rarity of this entity.

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