Surgical correction of persistent truncus arteriosus on a 33-year-old male with unilateral pulmonary hypertension from migration of pulmonary artery band

Wen Ruan, Yee Jim Loh, Kenneth Wei Qiang Guo and Ju Le Tan

Abstract

Background: Persistent truncus arteriosus is a rare congenital condition with which survival into adulthood is dismal without surgery. This is the oldest patient reported to our knowledge demonstrating the feasibility of assessing operability in persistent truncus arteriosus with unilateral pulmonary stenosis, and performing full corrective surgery in adulthood.

Case presentation: We report a Chinese male with successful correction of Type I persistent truncus arteriosus at 33 years of age. He had unilateral pulmonary hypertension from migration of pulmonary artery band from the main to the right pulmonary artery, severe truncal valve regurgitation from previous infective endocarditis, and progressive congestive heart failure. Improvement of lung perfusion was demonstrated 21 months post operation.

Conclusion: This case demonstrated that in patients with persistent truncus arteriosus and two pulmonary arteries, pulmonary vascular disease or underdevelopment of one lung does not preclude a full corrective surgery so long as the other vascular bed is normal. It is important to emphasize the importance of assessing patient’s operability in totality.

Keywords: Persistent truncus arteriosus, Truncal valve regurgitation, Unilateral pulmonary hypertension, Case report
Case presentation

A 33-year-old male was diagnosed to have type I truncus arteriosus (TA) during infancy. He underwent palliative main PA banding at the age of four months. No following corrective surgery was performed at his family’s choice. He had an uneventful childhood. At 26, he presented with *Streptococcal Gordonii* endocarditis with severe truncal valve regurgitation. This was complicated with septic emboli to his lungs and kidneys. At the same time, he was found to have left sided pulmonary hypertension due to migration of the PA band to the right PA (RPA) on the computed tomography of the chest (Fig. 1). He survived from a stormy hospital stay which required intubation and mechanical ventilation, and completed 5 weeks of antibiotics. A decision was made not for surgery at that point of time in view of the high surgical risk and uncertain surgical outcomes at this age [7].

He remained relatively well in New York Heart Association (NYHA) functional class I for 3 years, subsequently, his functional class deteriorated to NYHA II. On annual echocardiographic follow up, there was evidence of increase in size and decrease in systolic function of the left ventricle (LV). At 32 years old, he was hospitalized for worsening heart failure. On physical examination, he was thinly built with pectus carinatum, finger clubbing, raised jugular venous pressure and pitting edema over bilateral lower limbs. He had displaced apex beat, first and second heart sounds were present. There were ejection systolic murmur and early diastolic murmur at the left parasternal edge. The oxygen saturation was 90 % on room air. Blood investigation were unremarkable except for elevated hemoglobin of 17.9 g/L and serum creatinine of 105 mmol/L. Electrocardiogram revealed right ventricular (RV) hypotrophy with strain pattern. Chest radiograph showed an enlarged heart with prominent pulmonary trunk.

Trans-thoracic echocardiogram showed type I TA (Fig. 2). The LV was severely dilated with ejection fraction of 40–45 %. The truncal valve annulus was dilated with severe regurgitation. The RV systolic pressure was 116 mmHg with severe RV hypertrophy. Invasive haemodynamic study revealed progression of pulmonary vascular disease compared to 6 years ago. (Table 1) However, the RPA was still protected with no significant rise in PA pressure. Calculated left and right pulmonary vascular resistances (PVR) were 10.2 Wu and 2.6 Wu respectively. At the same time, lung perfusion scan showed marked deterioration of perfusion activity to the left lung (4.3 % vs 54 %) compared to 6 years ago (Fig. 3a and b).

In view of the worsening symptoms, cardiac size and function, as well as progression in asymmetrical pulmonary vasculature remodeling, decision was made for total correction at the age of 33. Surgery was performed via a standard median sternotomy and pericardiotomy. The establishment of cardiopulmonary bypass included ascending aorta, superior and inferior vena cava cannulation and core cooling to 25 °C. Circulatory arrest was achieved by fibrillating the heart. The bilateral branch PAs were snared at the commencement of cardiopulmonary bypass and aortic cross-clamping was applied. Truncus was transected, and the PA orifice and the coronary arteries were identified. A single dose of direct antegrade cardioplega (custodial) was introduced down the coronary ostium with spontaneous cardiac arrest.

Intraoperative findings were consistent with imagings: The truncal valve was tricuspid, the leaflet was damaged beyond repair by the endocarditis. The truncus root was very dilated and overriding into the RV (>50 %). The left coronary artery (LCA) orifice originated at a higher level than the right coronary artery (RCA) and had a short intramural course, arising from the posterior facing...
sinus. RCA arised from the anterior facing sinus. No abnormal course of the LCA.

The coronary buttons were cut out. Left ventricular outflow tract was established by a valve conduit (size 31 mm, Metronic, Inc, Minneapolis, MN) and the coronary buttons were anastomosed. The ventricular septal defect (VSD) was closed with Dacron patch. The RV to PA continuity was established by aortic homograft conduit (size 25 mm). As the stenosis of RPA from previous PA band was at the orifice of the RPA, this was dealt with during the suturing of the RV-PA conduit. Autologous pericardial "hood" was used to augment RV outflow tract to homograft anastomosis proximally. The distal end of the homograft was cut at an angle and then incised to enlarge the anastomotic suture line. Prior to completion, a trans-esophageal echo was performed to detail the VSD closure, bileaflet prosthesis function, and patency of PV-PA conduit with minimal regurgitation. Challenges associated with operation: 1) Bleeding. Patient developed massive bleeding originating from the Bental anastomosis, required going back on-pump and reinforcing the suture line towards the end of the surgery. Chest was kept open for 48 h with the support of intra-aortic balloon pump (IABP) until day 4 post operation. 2) Arrhythmias. After removal of IABP, the patient developed atrial fibrillation and was started on intravenous amiodarone. He then developed ventricular standstill and emergency arterio-venous Extracorporeal Membrane Oxygenation (ECMO) had to be inserted. He made good recovery after ECMO was explanted on day 7 post operation.

Patient was followed up after discharge and remained in NYHA II. At 21 months after the primary procedure, lung perfusion scan suggested improvement of the left lung perfusion activity from 4.3 to 20.2 % (Fig. 3c).

### Table 1: Haemodynamic Data Obtained during Cardiac Catheterization and Cardiac MRI

|                      | Year 2006 |                     | Year 2012 (pre-op) |                     |
|----------------------|-----------|---------------------|--------------------|---------------------|
|                      | Pressure (mmHg) | Oxygen Saturation(%) | Pressure (mmHg) | Oxygen Saturation(%) |
| Superior vena cava   | Mean 6    | 68.4                | Mean 5            | 62.1                |
| Inferior vena cava   | Mean 6    | 64.9                | Mean 4            | 70.5                |
| Mid right atrium     | 7/7/6     | 62.6                | 7/5/4             | 60.2                |
| Right ventricle      | 83/4/7    | 61.6                | 102/3/11          | 55.6                |
| Left ventricle       | 110/3/14  | 83.3                | 104/8             | 74.3                |
| Truncus              | 102/41/70 | 81.7                | 110/40/68         | 89.4                |
| Main pulmonary artery| 108/48/72 | 83.5                | 108/40/53         | 88.9                |
| Left pulmonary artery| 89/46/66  | 81.2                | 110/34/67         | 88.1                |
| LPA flow (fr MRI), ml/s |           |                     | 98                 |                     |
| LPA PVR (fr MRI), Wu |           |                     | 6.62               |                     |
| Right pulmonary artery (RPA) | 20/14/16  | 86.7                | 24/16/20          | 89.8                |
| RPA flow (fr MRI), ml/s |           |                     | 67                 |                     |
| RPA RVR (fr MRI), Wu |           |                     | 1.97               |                     |

Pressure data are expressed as systolic, diastolic and mean pressures, respectively. PVR, pulmonary vascular resistance

*LPA PVR = mean LPA pressure-LVEDP/flow LPA*
Discussion

There were several challenges when approaching this case: Firstly, truncal valve regurgitation is the major life-threatening complication of truncus arteriosus. It affects postoperative outcomes in patients with this cardiac anomaly. In our case, due to severe truncal valve damage and annular dilatation (>40 mm), it was difficult to find a suitable prosthesis (largest size 37 mm). Although several truncal valve reparative techniques have been reported, it remains challenging due to the unsatisfactory results [8]. 2) Origin of the truncal valve predominantly from the RV, causing under development of the aortic arch and preferential flow into PAs over that into the aorta. This is associated with a higher risk of creation of subaortic stenosis post-surgery. 3) Plicating the aortic root to suit the prosthesis size will almost require a root remodelling surgery with plication done right down to the base of the root. In such cases, the valves should be normal for maximal benefit. However the truncal valve was damaged from previous infective endocarditis. Also as the patient is very young. Plication of the root may run a risk of future aortopathy and dilation requiring a re-operation. Hence, we decided that an isolated truncal valve repair was not a long term solution for this patient. Taking into consideration, there is still unresolved unilateral pulmonary hypertension and progression of pulmonary vascular disease.

Secondly, there was no combined heart-lung transplant program available in our country. A ventricular assist device was not helpful due to uncorrected VSD, RV failure and raised pulmonary vascular resistance. However, the long term results of full PTA corrective surgery at a later age (above 1 year old) are largely unknown.

A thorough search of literature revealed a few small series of surgical repair in older children, and isolated case reports in adults. Marcelletti et al. reported a series of 96 patients with surgical repair of TA (73 % type I TA, 18 % has unilateral PA stenosis by nature or banding) from 1967 to 1975. The 30-day mortality for those underwent repair after 2 years old were about 21 %. The relation between pulmonary resistance and surgical risk was only studied for those with presence of both right and left PAs, and it was higher in patients with indexed PVR (iPVR) >8 W.u.m2 pre-operatively [9]. Stegmann et al. reported 8 cases of type I TA repair in patients’ age from 2 month to 4.5 years (2 of which had PA banding before) from 1976 to 1981. Only one 3.5-year-old child died postoperatively because of RV failure. In this child the iPVR had risen to 13 W.u.m2 despite PA banding [10]. In recent years, Talwar et al. reported 9 children with PTA (7 with type I PTA). Among the 9 patients, 2 had PA stenosis (1 at the origin of PA trunk from the TA and the other at the origin of RPA). These 9 patients underwent corrective surgery at mean age of 3 (range, 1 to 12 years) between 2000 and 2012. The hospital mortality was 22.2 % (2 deaths of Type 1 TA, 1 due to intractable pulmonary arterial hypertension and 1 due to sepsis). The rest had good functional recovery and no mortality after a mean 39 months (range, 3 month to 138 months) follow up [11]. Zhang et al. reported 12 patients (83 % Type 1 PTA) with only 1 patient having PA stenosis. They underwent repair of PTA at mean age 6.4 (range 1.2 to 19 years) between 2006 and 2010. There was no early mortality or requirement of re-operation after a mid-term follow up of mean 2.4 years (1.1 to 5.3 years). All patients had iPVR of less than 8 W.u.m2 [12]. These findings suggested that late correction of TA should be undertaken prior to the development of pulmonary vascular disease. On the other hand, successful surgical correction in adults were only reported in a 31-year-old male with PTA and severe pulmonary vascular obstructive disease [13], and a 28-year-old male with atypical PTA (The artery to the middle lobe of right lung originating from the descending aorta with stenosis at the origin). Although there were severe hypertension in the
right chambers (100/50/70 mmHg), his lung was protected until adulthood due to the pulmonary trunk stenosis at the origin [14].

Thirdly, difficulty in assessment of operability: With truncus arteriosus defects, the possible inequalities of pressure and flow between the two PAs made precise calculation of pulmonary resistance difficult. To solve this problem, a combined right heart catheterization for pulmonary pressures, and a cardiac MRI for flow in main and branch PA was used in this case. As pressure distal to the PA band was not elevated, pulmonary vasoreactivity test was not performed. In the era when cardiac MRI was not available, McFAUL et al. presented 27 patients with TA and previous PA banding (range, 6 weeks to 14 years). They managed to demonstrate that for patients who had two PAs, the presence of at least one adequate PA having low distal pressure or arteriolar resistance is the minimal criterion on which operability is based [15].

Conclusions
This case demonstrated that in adult PTA patients with two PAs, pulmonary vascular disease or underdevelopment of one lung does not preclude a full corrective surgery so long as the other vascular bed is normal. It is important to emphasize the importance of assessing patient’s operability in totality. This patient is the oldest PTA patient reported in literature to have undergone a complete repair at the age of 33, having had relatively protected right lung vasculature in infancy by migration of a proximately placed PA band. This, together with the patient’s improvement of lung perfusion scan postoperatively, is in congruent with other reports of successful corrective surgery in older PTA patients with either bilateral [9] or unilateral PA [16].

Consent
Written informed consent was obtained from the patient for publication of this case report and an accompanying images. A copy of the written consent is available for review by the Editor-in-chief of this journal.

Abbreviations
ECMO: extracorporeal Membrane Oxygenation; IABP: intra-aortic balloon pump; LCA: left coronary artery; LV: left ventricle; PA: pulmonary artery; PTA: Persistent truncus arteriosus; PVR: pulmonary vascular resistance; RCA: right coronary artery; RV: right ventricle; TA: Truncus arteriosus; VSD: Ventricular septal defect.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
JLT, KWOG carried out the clinical diagnosis and participated in provision of clinical information and reviewed the manuscript. YJL carried out the surgical procedure, participated in provision of clinical information and reviewed the manuscript. WR participated in collecting clinical and diagnostic information, repairing and revising the manuscript. All authors read and approved the final manuscript.

Acknowledgements
We greatly appreciate the assistance of the staff of the Department of Cardiology and Cardiothoracic surgery, National Heart Centre Singapore.

Author details
1 Department of Cardiology, National Heart Centre Singapore, Level 12, 5 Hospital Drive, Singapore 169609, Singapore. 2 Department of Cardiothoracic Surgery, National Heart Centre Singapore, Level 12, 5 Hospital Drive, Singapore 169609, Singapore.

Received: 25 December 2015 Accepted: 23 March 2016
Published online: 29 March 2016

References
1. Gatzoulis MA, Webb GD, Daubney PEF. Diagnosis and management of adult congenital heart disease. 2011. p. 271–6.
2. Marcelletti C, McGoon DC, Mair DD. The natural history of truncus arteriosus. Circulation. 1976;54(1):108–11.
3. Rajasinghe HA et al. Long-term follow-up of truncus arteriosus repaired in infancy: a twenty-year experience. J Thorac Cardiovasc Surg. 1997;113(5):869–78. discussion 878–9.
4. Kharwar RB et al. Persistent truncus arteriosus: a rare survival beyond the first decade. J Am Coll Cardiol. 2014;63(17):1807.
5. Abid D et al. Unrepaired persistent truncus arteriosus in a 38-year-old woman with an uneventful pregnancy. Cardiovasc J Afr. 2015;26(4):296–8.
6. Klewer SE BD, Atkins DL. Truncus arteriosus. In: Moller JH, editor. Surgery of congenital heart disease, Pediatric Cardiac Care Consortium 1984–1995. Philadelphia: Futura; 1998. p. 271–85.
7. Chiaw TH, San TR, Le TJ. An adult with truncus arteriosus and unilateral pulmonary hypertension. Congenit Heart Dis. 2007;2(6):433–7.
8. Imamura M, Drummond-Webb JJ, Sarris GE, Mee RB. Improving early and intermediate results of truncus arteriosus repair: a new technique of truncal valve repair. Ann Thorac Surg. 1999;67:1142–6.
9. Marcelletti C et al. Early and late results of surgical repair of truncus arteriosus. Circulation. 1977;55(4):636–44.
10. Stegmann T et al. Surgical correction of truncus arteriosus type I. Thorac Cardiovasc Surg. 1982;30(3):163–6.
11. Talwar S, Saxena R, Choudhary SK, Saxena A, Kothari SS, Juneja R, Airan B. Persistent truncus arteriosus repaired beyond infancy. Indian J Thorac Cardiovasc Surg. 2012;28(3):171–6.
12. O’Byrne ML et al. Morbidity in children and adolescents after surgical correction of truncus arteriosus communis. Am Heart J. 2013;166(3):512–8.
13. Oka Y et al. Truncus arteriosus in a thirty-one-year-old man with severe obstructive pulmonary vascular disease: a case report of successful surgical correction (author’s transl). Nihon Kyobu Geka Gakai Zasshi. 1980;28(1):1322–9.
14. Lopes LM et al. Atypical truncus arteriosus operated at 28 years of age: importance of differential diagnosis. Arq Bras Cardiol. 2011;97(2):29–32.
15. McFaul RC et al. Truncus arteriosus and previous pulmonary arterial banding: clinical and hemodynamic assessment. Am J Cardiol. 1976;38(5):526–32.
16. Nair DD et al. Truncus arteriosus with unilateral absence of a pulmonary artery. Criteria for operability and surgical results. Circulation. 1977;55(4):461–7.
17. Durmovicz AG et al. Unilateral pulmonary hypertension as a result of chronic high flow to one lung. Am Rev Respir Dis. 1990;142(1):230–3.
18. Davies P. How do vascular cells respond to flow? NIPS. 1989;42:22–5.