Unoperated tetralogy of Fallot: case report of a natural survivor who died in his 73rd year; is it ever too late to operate?

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The case history is described of a man in his 73rd year who was one of the oldest surviving patients with uncorrected tetralogy of Fallot (ToF) before succumbing with renal failure. Factors contributing to his longevity included small pulmonary arteries and presumed slow development of subpulmonary obstruction together with moderate concentric left ventricular hypertrophy—features previously seen in long term survivors. Less than 3% of all patients with uncorrected ToF survive beyond their 40s but late operative repair is still a valuable option. Practicalities of renal dialysis in the presence of an intracardiac shunt are considered.

Tetralogy of Fallot (ToF), first described in 1888, comprises an interventricular septal defect, right ventricular outflow tract obstruction, an overriding aorta, and right ventricular hypertrophy (RVH). It is the most common form of cyanotic congenital heart disease (10% of all cases). This case describes a man in his 73rd year who was one of the oldest surviving patients with an uncorrected ToF before succumbing with renal failure. We review the factors contributing to his longevity and consider whether surgery in his 66th year would have been appropriate.

CASE PRESENTATION

The patient, described as having a weak heart in childhood, never attended school or played sports. Diagnosis of ToF was not made until the age of 31 (1961) but he defaulted from further investigations. In 1996 (aged 66 years) he was reassessed while being investigated for proteinuria (urea = 5.7 mmol/l; Cr = 98 μmol/l). He was asymptomatic. He was mildly cyanosed and clubbed with pectus carinatum. Blood pressure was 140/80 mm Hg. He had an ejection systolic murmur in the pulmonary area. Echocardiography disclosed a subaortic VSD with a left to right shunt of 1.94 m/s, minimal aortic override (diameter 3.76 cm), small volume, hypertrophied well contracting LV, RVH, dilated RA, bicuspid pulmonary valve with a small pulmonary artery (1.4 cm), subpulmonary stenosis with an RVOT gradient of 112 mm Hg, and moderate PR. Electocardiography confirmed sinus rhythm, voltage criteria for RVH; RBBB with a QRS duration of 100 ms. Chest radiography showed a right aortic arch. Holter monitoring disclosed one asymptomatic 4 beat burst of SVT. Exercise testing terminated at four minutes because of fatigue (O2 saturation fell from 91% to 80%). The decision was made for medical management rather than surgical correction in view of his advanced age, asymptomatic status, and personal wishes. At the age of 69 the patient complained of regular fast palpitations and associated dyspnoea. The echocardiogram showed severely hypertrophied and impaired RV function, huge RA (6.8×8.2 cm), normal LV dimensions with moderate concentric hypertrophy as before, and a dilated LA (5.82 cm). Creatinine was 118 μmol/l. He was treated with frusemide, spironolactone, amiodarone, and aspirin. His renal function deteriorated in his 72nd year (urea = 18.5 mmol/l; Cr = 174 μmol/l), and he needed thyroxine for amiodarone induced hypothyroidism. He was never polycythaemic. In his 73rd year he had new onset AF, at which time his urea was 43.9 mmol/l and his creatinine was 436 μmol/l. He was cardioverted and treated with warfarin and aspirin. Four weeks later, AF recurred and despite cardioversion he remained hypotensive and unwell and died three days later. At postmortem examination the pericardial sac was obliterated with a dense fibrous pericarditis. The heart was enlarged (565 g) with massive enlargement of the right atrium and right ventricle. There was pronounced subpulmonary stenosis with an effective outflow tract of 10 mm; the pulmonary valve was bicuspid and the annulus was only 13 mm. The LA was moderately dilated and the LV was concentrically hypertrophied; there was no infarction. The VSD was 2 cm in diameter and abutted the aortic valve ring (fig 1). There was 90% stenosis of the RCA and recent haemorrhage into a proximal plaque, an 80% LAD stenosis, and a blocked circumflex. There were no systemic pulmonary anastomoses; lungs were oedematous. Macroscopically, the kidneys showed cortical atrophy.

**Figure 1** Postmortem findings of 2 cm ventricular septal defect (VSD), overriding aorta, and left ventricular hypertrophy (LVH).

Abbreviations: ToF, tetralogy of Fallot; RVH, right ventricular hypertrophy; LA, left atrium; RA, right atrium; LV, left ventricle; RVOT, right ventricle outflow tract; VSD, ventricular septal defect; LAD, left anterior descending
but postmortem autolysis prevented microscopic interpretation.

**DISCUSSION**

Without operation, few patients with ToF reach adulthood with an average life expectancy of 12 years. Ten per cent may survive to their 30s but only 3% reach their 40s or older. There are three main reasons for the longevity in natural survivors with unoperated ToF. Firstly, a hypoplastic pulmonary artery with slow development of subpulmonary obstruction. The pulmonary annulus was 13 mm in this patient, comparable to that reported in the other published cases—(age 45, PVD 10 mm), (age 77, PVD 9 mm). A second common feature is that of LVH as seen in this patient (16 mm); presumably this acts by delaying of shunting from the right to left ventricle. LVH may be a late development in the natural history of Fallot and any beneficial effect may not be seen until adult life. In this case hypoplasia of the pulmonary artery (seen in up to 50% of all Fallot’s) may have been sufficient early on to reduce pulmonary blood flow; then the subpulmonary stenosis needs to have also developed slowly over decades while his left ventricular hypertrophied concomitantly. Such a fine haemodynamic balance is clearly rare. The third finding in other cases has been extracardiac shunts including patent ducus arteriosus—reported in the oldest survivor who died aged 77—or systemic to pulmonary artery shunting via internal mammarys. There are few studies on the outcome and benefit of late surgical repair. The Mayo series followed up 30 patients who had total correction of ToF between the ages of 40 to 60 years. The operative mortality was 3% with long term survival rate at 5 years and 10 years postoperatively of 92% and 74% respectively. In this case, surgical repair would have been possible at the age of 66 years with low operative mortality and is likely to have prevented his subsequent systemic venous congestion. Nevertheless his personal wishes were not for operation— but given similar circumstances, operation would be recommended. Preoperative catheterisation also would have disclosed his coronary artery disease, which was unrecognised and untreated and probably had a role in his terminal decline. It is always important to consider detailed reinvestigation of any new elderly patient given a label of inoperable congenital heart disease as new approaches may be available to palliate the condition. He remained in sinus rhythm for a long time given his atrial dimensions; this undoubtedly improved his longevity and is probably a testament to amiodarone. His rapid decline was heralded by the change to atrial fibrillation resulting in prerenal failure—which was not a consequence of his longstanding congenital heart disease. Dialysis in patients with intracardiac shunts requires antibiotic prophylaxis at the time of line insertion, use of a blood filter downstream from the bubble trap on the venous limb of the dialysis circuit to avoid paradoxical embolism together with avoidance of rapid fluid removal, which may precipitate hypotension and exacerbate shunt reversal.

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**Learning points**

- Few patients with tetralogy of Fallot survive into adulthood without operation
- Unoperated survivors have three common features: hypoplastic pulmonary artery with moderately slow development of subpulmonary obstruction, left ventricular hypertrophy, or systemic-pulmonary artery collateral for pulmonary blood flow.
- Late operative intervention is associated with low mortality—it may never be too late to operate.
- Full re-investigation of older patients labelled as “inoperable” congenital heart disease may sometimes permit late palliation.
- Renal dialysis in the presence of an intracardiac shunt requires bubble filters to prevent paradoxical embolism.

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