Peripartum cardiomyopathy in uncorrected tetralogy of fallot: a case report

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ABSTRACT

Tetralogy of Fallot (ToF) is the most common cyanotic congenital heart with incidence about 10% of all congenital heart disease. Natural survival to the fourth decade is extremely rare (only about 3%), but there is a tendency of increasing number of women with cyanotic congenital heart disease living 3 to 4 decades and are becoming pregnant. Because of significant physiology adaptation and changes, pregnancy and delivery process are troublesome for most unhealthy women, including those with uncorrected ToF. For ToF patients, it remains an important cause of maternal morbidity (62.5%), and even mortality (10%) and has significant effects on fetal outcome. Discussed below a case of pregnancy in a 30 year old woman with uncorrected ToF, dyspnoea class II of New York Heart Association, during week pregnancy with complication of postpartum cardiomyopathy. Through dedicated medical care, patient's condition improvement can be seen. Prognosis of pregnancy in patient with uncorrected ToF is poor without optimal obstetrical and medical management.

Keywords: Uncorrected tetralogy of fallot, Pregnancy, Postpartum cardiomyopathy (PPCM)

INTRODUCTION

Tetralogy of fallot is the most common cyanotic congenital heart disease, with incidence about 10% of all congenital heart disease. The defects found in patient with ToF are caused by a single developmental defect: an abnormal anterior and cephalad displacement of the infundibular (outflow tract) portion of the interventricular septum. Four anomalies arising from this defect are (1) pulmonary stenosis, (2) right ventricular hypertrophy (3) overriding aorta (4) nonrestrictive ventricular septum defect.1

To reach adulthood, most patients with ToF will need surgery, either palliative or reparative. However, few patients will present as adults without correction of ToF. Natural survival into the fourth decade is extremely rare (only about 3%). Hemodynamic changes during pregnancy are profound with plasma volume overload, tachycardia and slight fall in peripheral vascular resistance. Cardiac output has to increase 50% above baseline during second stage of labour and at time of delivery.2

Pregnancy is troublesome for most unhealthy women, including those with uncorrected ToF. Due to fall in peripheral vascular resistance there is increase in right to left shunt, with subsequent increase in cyanosis.

During parturition blood loss may lead hypotension and eventually the right to left shunt. Thus, ToF patient notice a deterioration during pregnancy and parturition. These remain an important cause of maternal morbidity
(62.5%), and mortality (10%) and has significant effects on fetal outcome.3

CASE REPORT

A 30 year old woman G1P1L0A0 referred to cardiology department at 36 weeks and 2 days of gestation. She suffered from some congenital heart disease since childhood and was advised to undergo repair but could not do so, due to financial reason. Now she presented with dyspnoea NYHA class II symptom. Her pulse was 80/minute, blood pressure 112/70 mm of mercury, respiratory rate - 20/minute, temperature 98.4°F, cyanosis with grade II clubbing of finger, with oxygen saturation of 89% at hospitalization. On auscultation S1 was normal with single S2 and loud ejection systolic murmur over pulmonary area with normal respiratory examination.

Laboratory examination showed hemoglobin 13 g/dL, hematocrit 62%, WBC 10,200/dl, platelet count 2,20,000/cc. Other blood chemistry, urine analysis, and serum electrolytes were within normal limit. ECG showed sinus rhythm, QRS rate 78/minute, right QRS axis deviation 90, normal PR interval, incomplete right bundle branch block, and no significant ST- T changes.

Echocardiogram (Figure 1) showed levocardia and hypertrophy of inter ventricular septum and right ventricle, large VSD, bidirectional shunt, overriding aorta >50%, severe. Pulmonary stenosis, fair PA size, left aortic arch, no coarctation of the aorta, no collateral from descendent aorta with LV ejection fraction 60% finding consisting with Tetralogy of Fallot.

We preferred vaginal delivery over caesarian section for delivery mode. Multidisciplinary team, including anesthesiologists, cardiologists, cardio thoracic surgeons, obstetrics and pediatrics, managed the patient. Foetal examination was normal. She was kept in propped up position with continuous O2 inhalation. IV fluid and propranolol 20 mg tablet BD was given. Antibiotic prophylaxis with amoxicillin was administered before episiotomy with cut short of second stage of labour. Delivery was carried out without complications and was uneventful. The baby weighted 2200g and was stable. During labour and postpartum period we monitored the mother with blood pressure, heart rate, oxygen saturation and temperature three times daily. After delivery her general condition was fair. Uterus was well contracted. Her postnatal period was uneventful. She forced hospital to discharge with assurance of further follow up in cardiology OPD and counseled regarding need of cardiac surgery. Risk of future pregnancy explained and advised a barrier method for contraception.

She lost follow up and came after 1 month with dyspnoea NYHA class III, investigations showed hemoglobin 13.5g/dL, hematocrit 60.8%, WBC 6200/cc, platelet count 2.80,000/cc. Arterial blood gas analysis showed pH: 7.307, pCO2: 24.3 mmHg, pO2: 50.6 mmHg, saturation of O2: 87%, HCO3-: 12.6 mEq/l; showing uncompensated metabolic acidosis, urine analysis, and serum electrolytes were within normal limit. She was afebrile, her pulse rate was 128/minute, blood pressure 100/70 mm of mercury, respiratory rate 28/minute, bilaterally symmetric chest movement with crepitation on right lower zone, and baseline cyanosis. On auscultation S1 was normal, single S2, gallop rhythm and ejection systolic murmur over pulmonary area.

ECG (Figure 2) showed sinus tachycardia, QRS rate 114/minute, right QRS axis deviation 90, left atrial enlargement with normal PR interval, incomplete right bundle branch block, and T wave inversion in lead I, aVL, V2 to V6.

Echocardiography this time showed congested IVC, TOF physiology but there was striking finding with LV ejection fraction <35%. Serial Trop I was markedly positive with decreasing trend, with significant ECG findings, and X ray we came to conclusion of TOF with postpartum cardiomyopathy. We stabilized patient with
diuretics. There is very small and controversial data regarding long-term management of uncorrected TOF patient with postpartum cardiomyopathy. We advised our patient diuretics and after stabilization low dose of ACE inhibitor and B blocker with close clinical and echocardiography follow up. Counseled about their cardiovascular risk and advised against pregnancy.

Administration fluid before procedure is recommended to avoid hypotension. In third phase of delivery, it is recommended to limit the bleeding amount. These patients should be given prophylactic antibiotic because they are at high risk of infective endocarditis.7

The pregnancy with uncorrected TOF carries major risks of maternal morbidity, including heart failure, arrhythmia, endocarditis, thromboembolism, even maternal mortality. In present case, patient presented with postpartum cardiomyopathy (PPCM). There is strongly suggested a role for vascular dysfunction, hormonal insults, and underlying genetics in PPCM. LV ejection fraction at presentation best predicts rate of recovery of PPCM.

The risk of PPCM relapse in patients with persistent LV dysfunction is much higher in subsequent pregnancy than in those who have normalized LV function. Most patients will recover by 6 months, with little change in EF thereafter. Treatment is focused, as with other forms of systolic failure, on controlling volume status, neutralizing maladaptive neurohormonal responses, and preventing thromboembolic and arrhythmic complications. Neurohormonal blockade with ACE inhibitors or angiotensin receptor blockers can be used postpartum.8

Education about reparative procedure after pregnancy and contraception method to prevent further pregnancy before reparative process is needed to be given. The number of TOF patients are steadily increasing each year so the number of TOF patients with pregnancy. There is a strong need for carefully coordinated multidisciplinary efforts to manage these patients and maintain long-term follow-up.

CONCLUSION

The management of such patients with proper monitoring, bed rest, beta blocker, diuretics and ACE inhibitor significant improvement can be seen. Corrective surgery after stabilization should be done. Without proper cardiac care and obstetrical management, prognosis of pregnancy in patient with uncorrected TOF is poor.

Postpartum these patients should be vigilantly monitored both clinically and by ECG, X ray, echocardiography for postpartum cardiomyopathy like complications. Such patients should be counselled regarding their

Figure 3: X ray Chest PA view showing cardiomegaly.
cardiovascular risk and advised against further pregnancy. Further studies are required for management of such patients.

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