Mitral valve Replacement in a 4 yrs old Child: First Time in Bangladesh

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Abstract:
Congenital mitral valve incompetence is a rare and complex congenital heart disease in children. We report a case of a 4-year-old child admitted to hospital with fever, dyspnea on exertion or feeding and repeated respiratory infection for last 3 years. The transthoracic echocardiogram revealed grossly dilated left atrium and left ventricle and severe mitral regurgitation due to cleft in anterior mitral leaflet. Per-operatively mitral valve annulus was found very much dilated; leaflet thinned out and rudimentary posterior mitral leaflet. Morphology of mitral valve was totally distorted, leaflets were diminutive and beyond repairable. Mitral valve replacement was done with 25 mm Edward Life Science porcine tissue heart valve with total preservation of subvalcular structure and the patient showed dramatic symptomatic improvement and later follow up revealed good LV function with alleviation of symptoms. This is a rare and unusual case of congenital mitral valve disease with better prognosis after surgical replacement with tissue valve.


Key Words: Congenital heart disease, mitral regurgitation, mitral valve replacement.

Introduction:
Congenital malformations of mitral valve are rare, complex and frequently associated with other congenital malformations of the heart or the aorta that may hide or be hidden by the valve malformation. Congenital mitral incompetence is even rarer. Congenital mitral valve incompetence may lead to repeated pulmonary infections, cardiac failure and eventually failure to thrive. In practice, repeated episodes of cardiac failure and progressive enlargement of systolic diameter indicates a deterioration of systolic function. This must be recognized and should indicate an operation, whatever the age. Despite improving surgical techniques, treatment of heart valve disease in children remains controversial. Growth of the child and adequate anticoagulation level are the main concerns when valve replacement is preferred in the pediatric age. In patient with congenital mitral valve disease, reconstructive surgery is the primary goal. However, in cases with severely dysplastic valves or failed repair, mitral valve replacement is the only option.

Case report:
A 4 years old child admitted on Metropolitan medical centre, Dhaka, Bangladesh with the complaints of fever, dyspnea on exertion or feeding and repeated respiratory infection requiring repeated hospitalization for last 3 years. Physical examination revealed a regular pulse 100 beat/min, blood pressure 100/60 mm of Hg, and bibasal crackles on chest auscultation. Her first heart sound was developed in mitral area and second heart sound was normal but there is a blowing pan systolic murmur present at the apex. Electrocardiogram showed heart rate 100/min, regular with feature of left atrial enlargement and left ventricular hypertrophy. The chest X-ray showed pulmonary congestion and a moderate cardiac enlargement. Color Doppler echocardiogram revealed grossly dilated left atrium and left ventricle, severe mitral regurgitation due to cleft in anterior mitral leaflet (Figure-1) and PASP 86mm Hg.

Fig.-1: Cleft in anterior mitral leaflet (echocardiographic view)

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Patient was operated on 18.09.2013. Under general anesthesia with all aseptic precautions standard median sternotomy was done. After thymus dissection pericardiotomy was done. Cardiopulmonary bypass was established with bicaval cannulation with aortic cannulation. Heart was arrested by giving crossclamp and antegrade cardioplegia under mild hypothermia (32°C). Left artiotomy done. Mitral valve annulus was very much dilated; leaflets were thinned out, rudimentary posterior mitral leaflet. Morphology of mitral valve totally distorted and it was beyond repairable. Mitral valve replacement was done with 25 mm Edward Life Science porcine tissue heart valve with total preservation of subvalvular structure (figure-2). A vent was given through right superior pulmonary vein to left atrium. Left artiotomy closed. Patient weaned from cardiopulmonary bypass without any difficulty. X-clamp time was 28 min and Total bypass time was 60 min. After achieving proper haemostasis, chest was closed leaving one mediastinal drain tube (retrosternal 24Fr) attached under water seal drain bag. The patient was shifted to the ICU with minimum inotropic support. Patient was extubated on the same day, shifted to general word on 3rd POD and discharged home on 10th POD with advice of taking oral anticoagulant Warfarin Sodium 1mg daily only for 3months followed by antiplatlet therapy. Patient remains asymptomatic in the subsequent follow up (figure-3) and leading to a normal life.

**Discussion:**
Isolated congenital mitral regurgitation is caused by defects in the leaflet tissue (isolated cleft), absent or redundant chordate tendineae and abnormal size or location of papillary muscle. Our patient have congenital mitral regurgitation due to cleft in anterior mitral leaflet with diminutive mitral leaflet tissue. At all ages, patients with mitral regurgitation present with various degrees of failure to thrive, dyspnea on feeding or exertion. An enlarged left ventricular impulse may be present and a high-frequency, high-intensity holosystolic murmur is heard at the apex extending into the axillae. Isolated congenital mitral regurgitation is often only moderate in severity in early life and about half of the patient with it do not show development of important symptoms or come to operation until older than about age 5 years. The sign and symptom of our 4 years old patient started from 1 year and severity of symptoms was gradually increasing with her age.

The electrocardiogram shows left atrial and left ventricular enlargement in patient with mitral regurgitation and chest radiography demonstrate cardiomegaly. Our patient also had left atrial and left ventricular enlargement on electrocardiogram and cardiomegaly on chest X-ray.

Transthoracic echocardiography provides nonspecific evidence as to enlargement of left
ventricle, left atrium and right ventricle. Real time 3-dimensional echocardiography demonstrated the cleft with its wide and depth, the extent of cleft edge fibrosis and retraction, and the presence of accessory chordae and their attachment to the septum. Our patient have only cleft on anterior mitral leaflet with rudimentary posterior mitral leaflets.

Mitral valve replacement might be necessary in children with extremely dysplastic valves and severe hemodynamic impairment or after failed repair. Mortality and mitral reoperation are common after mitral valve replacement in children and outcomes can be predicted based on patient’s age and other associated factors. A study showed that bileaflet prosthesis larger than 23 mm have the lowest operative risk. Valve related complications are thromboembolism, hemorrhage, structural deterioration and non-structural dysfunction. A study done on comparing on 24 children on 2 to 18 years of age who have received mechanical valve compared with 24 children who received porcine. Major complications were seen in 50 percent of mechanical valve group and 13 percent of porcine valve group. Superiority of tissue heart valve on children is established. A study on 64 children suggested that despite the disappointment occurrence of premature tissue valve failure in young population, valve replacement in children currently safer. Our patient has extreme dysplastic mitral valve and cleft anterior mitral leaflet and rudimentary posterior mitral leaflet with significant hemodynamic impairment. So, repair was not feasible for her. We replaced the mitral valve with 25mm porcine tissue heart valve without any post-operative complication which supports these data.

A study showed that the durability of bioprosthetic heart valve is severely limited by early calcification and therefore mechanical heart valves are the preferred option, albeit at the expense of lifelong anticoagulation. Now a days, because of better anticalcific treatment, the durability of tissue valve is increased. So, considering the difficulty in maintaining anticoagulation in remote rural area of Bangladesh tissue heart valve was preferred in children.

Reference: