**Case Report**

**Medicine Science 201:.(.)….**

**Down Syndrome is not contraindication for surgical treatment of complex cardiac anomalies: two case report**

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Received 26 November 2016; Accepted 09 December 2016
Available online 22.12.2016 with doi: 10.5455/medscience.2016.05.8568

**Abstract**

Approximately half of the individuals with Down syndrome have congenital heart disease requiring surgical repair. A holistic approach to treatment of these individuals should be focused on the management of physiological and cognitive functions. For cases requiring surgical treatment, adequate level of physiological functions is not expected without surgical treatment. Inadequate physiological functions may cause ineffective physical and cognitive functions. Early diagnosis and surgical treatment is crucially important for these individuals. Individuals with Down Syndrome, if not treated surgically because of high operative mortality, the decrease in functional capacity and quality of life is unavoidable. In both of our cases, surgical treatment was delayed by their families. Families should be informed about the success of the early surgical treatment for cardiac problems in Down syndrome.

**Keywords:** Down Syndrome, cardiac surgery, family awareness

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**Introduction**

Down Syndrome is characterized with the 21st chromosome being 3 units. The phenotype of Down’s Syndrome consists of intellectual impairment, short stature, heart diseases, digestive disorders, and orthopedic anomalies associated with abnormal physical and neurological findings [1].

The most common congenital heart diseases in individuals with Down syndrome are atrioventricular septal defects and conotruncal defects. On the other hand, 59-72% of all atrioventricular septal defect cases are Down syndrome cases. Fallot Tetralogy (TOF) anomalies are common in Down syndrome [2]. 50% of Down syndrome births can be associated with primum ASD, 65-70% with secundum ASD, and 40-50% with sinus venosus defect [3].

The leading cause of mortality in the first two years of life in Down syndrome cases is cardiac malformations [1]. Conditions with the highest mortality rate in Down Syndrome are those associated with tetralogy of Fallot [4].

The aim of this study is to raise awareness of the importance of early diagnosis and treatment of the congenital heart diseases that affect morbidity and mortality in Down Syndrome. We would like to emphasize the importance of performing surgical therapy in the presence of complex anomalies accompanying Down syndrome.

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**Case Report 1**

A 24-year-old male patient referred to cardiology polyclinic with complaints of palpitation and dyspnea. The physical examination revealed that the mildly obese patient had Down Syndrome. There was a 2/6 systolic murmur at the pulmonary area. ECG showed normal sinus rhythm. Cardiomegaly was present on the telecardiography. Echocardiographic examination revealed a sequential type defect with a diameter of 2.3 cm and left to right shunt in the interatrial septum. The QP / QS ratio was measured as 4.1. The patient was recommended for operation for large secundum ASD diagnosis. After describing the risks to the patient's family, patient was taken to the operation with standard open heart surgery techniques (normothermic cardiopulmonary bypass with membrane oxygenator). Large secundum ASD was closed with pericardial patch. The patient who had a postoperative period without any problems. He was discharged on the 5th postoperative day. The patient, had surgery 5 years ago, is currently at FC1 according to the New York Heart Association (NYHA) functional capacity classification. Echocardiographic controls showed intact interatrial septum.

**Case Report 2**

A 19-year-old woman, known to be Down's syndrome, was referred to the cardiology polyclinic for further examination and treatment. She had shortness of breath, complaints of cyanosis on her lips and quick fatigue after effort. In the story taken from her family, when she cries, her lips turn purplish. She developed growth retardation comparing to her siblings. She could not run like her friends. She wanted to squat after slight effort.
Perioral cyanosis, clubbing on the fingers and systolic murmur of 3/6 degrees along the left side of the sternum was found at the physical examination of the patient. ECG had sinus rhythm, right ventricular hypertrophy. Echocardiographic examination revealed that dilated right aurium and ventricle, intact interatrial septum, a 1.5 cm diameter VSD in the subaortic region, shunt from left to right and right sided deviation of the aorta. Hypertrophy on the right ventricular outflow tract and valvuler and subvalvular pulmonary stenosis were found. There was 80 mm Hg gradient across the pulmonary valve. The patient was diagnosed as fallot tetralogy. The operation was suggested. All the risks of the operation was discussed with the family. After the necessary preparations, the patient was taken to the operation room. Total Correction Surgery (closing of the VSD and removing the pulmonary stenosis) was performed (mild hypothermic cardiopulmonary byass with membrane oxygenator).Postoperative follow-up period was uneventful. The patient was discharged on the 8th day with necessary suggestions. At the postoperative 4.th year the patient is in NYHA FC I, interventricular septum intact at echo examinations and 20 mmHg gradient was detected on pulmonary valve. The medical follow-up and treatment of the patient continues.

Discussion

It is known that about half of individuals with Down's syndrome have congenital heart disease requiring surgical repair. The holistic treatment approach of these individuals focuses on the management of physical, physiological and cognitive functions. In cases where surgical repair is required, it is unlikely that physiological functions will be adequate without treatment. Inadequate physiological functions may lead to the failure to obtain the desired efficiency in physical and cognitive functions. This is an important situation that should be aware of the professionals and families living in the rehabilitation process with a patient with down syndrome.

Studies have shown that the risks involved in surgical approaches to the treatment of congenital heart diseases in the Down's syndrome population are not different from the risks encountered in the surgery of patients without Down's syndrome [4]. Along with improvements in cardiovascular surgery, survival rates after surgery are increasing steadily due to early surgery, improvements in myocardial protection methods, and better postoperative care [5]. Today’s accepted information is the families should be encouraged as much early surgical treatment as possible by the surgical team. Late-acting surgeries likely to cause further impairment of valve function and development of residual pulmonary hypertension.

Findings from a study of adult Down syndrome who had surgery in an experienced center suggest that Down syndrome may have heart surgery with a very low mortality and morbidity risk [6]. We think that the duration of hospitalization in our cases with short duration like 5 and 8 days is a significant factor to reduce the mortality when considering the postoperative complications due to long hospital stay.

Early diagnosis and early surgery in the treatment of congenital heart diseases in individuals with Down syndrome is important for the prognosis of the disease. It is inevitable that individuals with Down Syndrome suffer from a significant decrease in their functional capacities and a negative impact without surgery. Their quality of life becomes worse if they are not operated on the basis of a life threatening risk of heart surgery [7]. However, as seen in both of our cases, surgical treatment was delayed considerably. Families should be informed about the success of surgical treatments for cardiac problems in Down Syndrome.

As a result, solutions are possible, no matter how complex the problems are. In order to inform and direct this issue, we have great responsibility becoming the health professionals against to the families of individuals with Down Syndrome.

References