Heterotaxy Asplenia Syndrome

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ABSTRACT

Introduction: Heterotaxy asplenia syndrome (right isomerism or bilateral right-sidedness), which is associated with a centrally located liver, absent spleen, and two morphologic right lungs usually associated with dextrocardia and cyanotic congenital heart disease.

Case Presentation: We report the case of dextrocardia with heterotaxy syndrome who presented with respiratory distress soon after birth. A chest X-ray showed dextrocardia, cardiomegaly with pulmonary congestion. The findings of an electrocardiogram and echocardiography confirmed dextrocardia and an abdominal Ultrasound showed liver central more on right side with mild hepatomagly, no spleen visualized, stomach on left side, pancreas both kidneys urinary bladder normal with free minimal fluid.

Conclusions: Heterotaxy asplenia syndrome a rare group of congenital malformations that must be fully evaluated. There is need for a complete and elaborate diagnostic work up of suspected cases by using various imaging modalities so that they are not missed. Doctors should encourage routine medical examination for their patients which could help identify this anomaly, thereby preventing wrong diagnosis and possibly death due to delay in management.

Keywords: Dextrocardia; heterotaxy; asplenia syndrome.

INTRODUCTION

Heterotaxy asplenia syndrome (right isomerism or bilateral right-sidedness), which is associated with a centrally located liver, absent spleen, and two morphologic right lungs usually associated with dextrocardia and cyanotic congenital heart disease. (1)

CASE PRESENTATION

Full Term male Baby delivered to Gravida 4 Para 3 mother with no history of any maternal illness presented with respiratory distress. Baby cried well after birth with good apgar score but developed respiratory distress at 22 hours of age with tachypnea and grunting respiration baby was started on oxygen 2Liter/min by nasal pronge. On examination baby had birth weight of 3.25 kg with stable vital signs SPO2 96%. Baby had respiratory distress, tachypnea and chest retractions, air entry equal, no crepitations with no dysmorphic features. CVS: All peripheral pulses felt, no radio femoral delay, heart sounds better heard on right side, heart sounds normal with III/VI systolic murmur in the right sternal border Abdomen: Soft and lax, liver 3 cm below right costal margin, bowel sounds normal, normal male genitalia. CNS: normal.
**Investigations:** CBC biochemistry within normal limits.

**Chest X-ray:** dextrocardia, cardiomegaly with pulmonary congestion. ECG showed inverted P waves in lead 1.

Abdominal Ultrasound showed liver central more on right side with mild hepatomegaly, no focal lesions seen, Gall bladder normal location and morphology, no spleen visualized, stomach on left side, pancreas both kidneys urinary bladder normal with minimal fluid in abdomen.

Echocardiography revealed situs ambiguous, dextrocardia, IVC connected to large common atrium, Right atrial isomerism, only one coronary sinus seen. Pulmonary veins connected to large common ambiguous atrium and 2 small muscular VSD, large perimembranous VSD, single common AV valve with pulmonary hypertension, right ventricle larger than left and both vessels arising from right ventricle with transposition of great vessels. No aortic stenosis or pulmonary stenosis, pulmonary valve larger than aortic valve, ascending aorta and proximal aortic arch normal, distal arch and preductal descending aorta narrowed. PDA moderate size with left to right shunt. Normal coronaries and postductal descending aorta.

Progress - Baby developed increasing respiratory distress, and CO2 retention so ventilated on day 4 of life and referred to higher centre for surgical management.

**DISCUSSION**

Heterotaxy syndromes are rare occurring in 1/10,000 live births (Lin et al, 2002) with male female ratio 2:1. (1,2) If the visceroatrial situs cannot be readily determined, a condition known as situs ambiguous or heterotaxia syndrome. (2) The two major variations are (1) asplenia syndrome (right isomerism or bilateral right-sidedness), which is associated with a centrally located liver, absent spleen, and two morphologic right lungs, and (2) polysplenia syndrome (left isomerism or bilateral left-sidedness), which is associated with multiple small spleens, absence of the intrahepatic portion of the inferior vena cava, and bilateral left lung morphology (i.e., in both lungs). The heterotaxia syndromes are usually associated with conduction defects and severe congenital heart lesions: ASD, VSD, atrioventricular septal defect, single functional ventricle, transposition of great arteries, pulmonary stenosis or atresia, and anomalous systemic venous or pulmonary venous return. (1,2)

In our patient situs ambiguous was associated with asplenia and dextrocardia. Cardiac anomalies identified on echocardiography were dextrocardia, IVC connected to large common atrium, Right atrial isomerism, only one coronary sinus
seen. Pulmonary veins connected to large common ambiguous atrium and 2 small muscular VSD, large perimembranous VSD, single common AV valve with pulmonary hypertension, right ventricle larger than left and both vessels arising from RV with transposition, pulmonary valve larger than aortic valve, ascending aorta and proximal aortic arch normal, narrowing of distal arch and preductal descending aorta. PDA moderate size with left to right shunt. Normal coronaries and post ductal descending aorta. In the vascular anomaly IVC and aorta were both on left side of vertebral column. However, dextrocardia with situs inversus is associated with a lower incidence of congenital heart disease (0 to 10%). Presentation of cause varies depending on associated malformation. (1-4) Situs inversus may be associated with other congenital anomalies such as duodenal atresia, asplenism, multiple spleens, ectopic kidney, horseshoe kidney and various pulmonary and vascular abnormalities. (5,6) Situs inversus totalis that is associated with primary ciliary dyskinesia is known as Kartagener syndrome. (7) Patients with primary ciliary dyskinesia have repeated sinus and pulmonary infections. (8) The arrangements of the position of the abdominal viscera in dextrocardia may be normal (situs solitus), reversed (situs inversus), and indeterminate (situs ambiguous or isomerism) in 32 to 35%, 35 to 39% and 26 to 28% of cases respectively. (8)

Dextrocardia with a normal abdominal situs has a high incidence of associated congenital cardiac anomalies including among others, transposition of the great vessels and ASDs (9) and VSDs (9) in 90 to 95% of cases. Diagnosis of dextrocardia is usually confirmed by several modalities which include chest radiography, ECG, echocardiography, computed tomography, magnetic resonance imaging and abdominal ultrasonography. Echocardiography is one of the modalities for making the diagnosis. Of interest, this patient had situs ambiguous with complex cardiac anomalies. This case is reported because of situs ambiguous, dextrocardia and asplenia with early symptomatic presentation due to complex pattern of cardiac malformation.

CONCLUSIONS

Heterotaxy asplenia syndrome a rare group of congenital malformations that must be fully evaluated when noticed because in rare instances it may result in fatal outcome. There is need for a complete and elaborate diagnostic work up of suspected cases by various imaging modalities so that they are not missed. Surgeons, radiologists and radiographers should look out for this anomaly during preoperative and surgical management of their patients. Doctors should encourage routine medical examination for their patients which could help identify this anomaly, thereby preventing wrong diagnosis and possibly death due to delay in management.

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