Heterotaxy syndrome: a case report

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Abstract

The present report describes the findings at chest computed tomography angiography of a 28-year-old female patient with heterotaxy syndrome. This syndrome consists of a variety of anomalies of position and morphology of thoracoabdominal organs which do not follow the situs solitus or situs inversus arrangement. Imaging studies play a fundamental role in the individualization of the approach to the patient.

Keywords: Heterotaxy syndrome; Situs inversus; Spleen – abnormalities.

INTRODUCTION

Heterotaxy syndrome (from the Greek heteros – different and taxis – arrangement) consists of several position and morphology abnormalities of thoracoabdominal organs, which do not fit in the habitual positioning order of organs laterality (situs solitus) or its mirror image (situs inversus)1–3. It is a rare entity, and its severity is mainly due to cardiac manifestations.

Terms such as asplenia (right atrial isomerism) or polysplenia (left atrial isomerism) are falling into disuse on account of the wide spectrum of anatomical findings, with no pathognomonic characteristic existing which would allow the subclassification under such terms3.

CASE REPORT

Female, 28-year-old patient with clinical signs of dyspnea since her childhood, World Health Organization functional class II. At the age of two, the patient was submitted to catheterization, with the following findings: dextrocardia, functionally single atrium, corrected transposition of great vessels, persistence of the arterial canal and interventricular communication. In her adulthood, echocardiogram presented left atrial isomerism and dextrocardia, interruption of the inferior vena cava, atrioventricular septal defect (partial presentation), patent arterial duct with aortopulmonary shunt. Five years ago, significant pulmonary hypertension was diagnosed.

Four months ago the patient presented sudden worsening of dyspnea associated with chest pain, mild cyanosis, systolic murmur more audible in the pulmonary focus, with fixed split of S2, physiological vesicular murmurs and oxygen saturation at 70%.

A chest computed tomography angiography was requested, and demonstrated dextrocardia and cardiomegaly with predominance of the right chambers, atrial septal defect, increased caliber of the pulmonary trunk (35 mm) with no sign of pulmonary thromboembolism, ectatic ascending aorta with descending segment at the left, infrahepatic vena cava interruption, andazygos vein engorgement from the infra-diaphragmatic portion (Figures 1 and 2). Other findings included polysplenia and bilobed lungs, with the main bronchi coursing inferiorly to the pulmonary arteries, and the liver in centrally located, characterizing heterotaxy syndrome (Figures 3 and 4).

DISCUSSION

The habitual and orderly arrangement of the organs in the human body is determined early in the embryonic development and is based on genetic information4. The loss of such orderly arrangement may characterize situs inversus or a disordered and variable arrangement (heterotaxy syndrome). Heterotaxy syndrome presents an approximate incidence of 1:10,000 births and is slightly more prevalent in men, at a ratio of 2:13,5.

Cardiac abnormalities are present in 50% to 100% of patients6 and, generally, such abnormalities are accountable for the severity and mortality of patients with such a syndrome. Heterotaxy syndrome encompasses a wide range of extracardiac variants, including: urinary system, biliary tract and hepatic abnormalities, duodenal atre-
Figure 1. Chest computed tomography angiography demonstrating dextrocardia, cardiomegaly and atrial septal defect (arrow).

Figure 2. Chest computed tomography angiography showing ectatic ascending aorta, (arrow head) with descending segment at left (small arrow), engorgement of the azygos vein (large arrow), besides increased caliber of the pulmonary artery trunk (figure at right).

Figure 3. Chest computed tomography angiography showing bilobed lungs, with main bronchi coursing inferiorly to the pulmonary artery (arrow).

Figure 4. Chest computed tomography angiography demonstrating polysplenia (arrow) and liver centrally located.
sia, gastrointestinal malrotation and several cardiovascular alterations\(^6\).

The presentation previously called “asplenia” generally presents duplication of structures located in the right side of the body, with trilobed lungs, left atrium with morphology corresponding to the right atrium, centrally positioned liver, left-sided aorta and inferior vena cava, besides intestinal malrotation.

In most such cases, death is an early event\(^3\) as a consequence of complications from congenital heart defects, notably the presence of a single atrioventricular chamber. Other reported complications include immunological changes and intestinal volvulus\(^7\).

In the presentation previously called “polysplenia” generally there are duplication of the structures in the left side of the body, presenting bilobed lungs, right atrium anatomically identical to the left atrium, liver also centrally positioned, absence of the hepatic segment of the inferior vena cava with continuity through the azygos or hemiazygos vein, besides intestinal malrotation. Cardiac alterations are less frequently found and milder, a fact that explains a higher prevalence of such findings in individuals of more advanced ages\(^5\).

The multiplicity and diversity of findings in such a syndrome make the individualization of cases extremely valuable, as most of them do not perfectly fit in any classification. Thus, the radiological evaluation is indispensable for the identification and planning of the approach to patients presenting cardiac and immunological complications or surgical conditions, allowing for the evaluation of the alterations present in each patient\(^8\).

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**REFERENCES**