A Rare Case of Recurrent Pneumonia in Heterotaxy Syndrome, Polysplenia/Left Isomerism

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Abstract

Heterotaxy syndrome (HS) or situs ambiguous refers to the abnormal arrangement of viscera across the body axis, and abnormalities arise depending on the isomerism of the right or left atrial appendage. The cause remains unexplained and is attributed to a combination of genetic mutations and environmental factors. It is a rare condition and may remain undiagnosed for a long time. In this report, we aim to highlight an unusual presentation and aggravation of an infection due to the underlying isomerism of the left atrial appendage. We discuss the case of a female patient who presented with symptoms of fever and cough. The patient underwent prolonged antibiotic treatment, and her recovery was slow. The presence of bilobed lungs, vertical left-bronchus, and polysplenia on CT scan explained the left-sided aspiration pneumonia. The hypofunctioning spleen (polysplenia) caused her to have a weak immunological response, necessitating prolonged antibiotic use. She was followed up over time and had a recurrence of pneumonia within a few months. The condition is associated with high morbidity and mortality, and the role of early diagnosis and reporting to prevent complications is paramount. The recurrent pneumonia observed in the patient also raises questions related to long-term antibiotic use and immunization in the case of polysplenia in this patient population.

Categories: Radiology, Infectious Disease, Pulmonology

Keywords: congenital heart defects, situs ambiguous, immunisation, left atrial isomerism, pneumonia, polysplenia, heterotaxy syndrome

Introduction

Heterotaxy syndrome (HS), also known as situs ambiguous, is a rare, complex birth anomaly that leads to the abnormal positioning of internal thoracoabdominal organs across the left-right axis of the body. The term “heterotaxy” is derived from the Greek words “heteros” meaning “other than” and “axis” meaning “arrangement” [1]. Situs ambiguous includes manifestations that cannot be classified into either situs solitus or situs inversus. While situs solitus refers to the normal arrangement of these structures, situs inversus is a condition characterized by the mirrored organization of the viscera. It occurs in one out of 10,000 live births, and a large number of these cases result in stillbirths [2]; it is more prevalent in males than females, with a male-to-female ratio of 2:1.

HS is classified based on the characteristics of the spleen or the morphology of atrial appendages. Isomerism refers to the aberrant bilateral symmetry of specific viscera: isomerism of the right atrial appendage (IRAA) or asplenia syndrome, and isomerism of the left atrial appendage (ILAA) or polysplenia [3]. Mutations of genes involved in the lateralization of body organs result in embryonic disruptions in the left-right axis determination. However, due to the heterogeneity in HS, the attempts to identify the precise genetic alterations have remained unyielding to date. The peculiar genotypic and phenotypic aspects of the syndrome result in varying presentations among different individuals. The bronchial-atrial disarrangement in IRAA causes more severe cardiac abnormalities than in ILAA [1]. Moreover, the extracardiac variants involving tracheoesophageal fistula, duodenal atresia, biliary tract malformations, and gastrointestinal malrotations are frequently seen with ILAA [4]. Congenital heart defects lead to high rates of mortality and morbidity. The widespread use and availability of CT scans have made it possible to diagnose heterotaxy as an incidental finding in healthy individuals. This syndrome is associated with diverse anomalies, and delayed diagnosis may lead to a worse prognosis. Prompt diagnosis and early remedial intervention are essential to reduce morbidity and mortality.

Left bronchial isomerism is associated with polysplenia in about three-fifths of the cases, and right bronchial isomerism is found in about seven-tenths of cases with asplenia [5]. Left isomerism is also characterized as bilateral left-sidedness, in which there is the duplication of left-sided viscera such as bilateral pulmonary atria, bilateral bilobed lungs, and bilateral hypophterial bronchi.

Case Presentation

A previously healthy 34-year-old female with a past medical history of behavioral health problems,
including impulse control disorder, bipolar disorder type 1, depression, and post-traumatic stress disorder (PTSD) was referred to the ER by the primary care physician due to the persistent complaint of low-grade fever and cough associated with meals. The patient also complained of nausea and vomiting after meals. There was no history of anorexia, weight loss, headache, dizziness, or blurry vision. She had a history of tracheoesophageal fistula repair and multiple abdominal surgeries for intestinal obstruction. On the basis of suspicion of aspiration pneumonia on a chest X-ray, she was referred to the emergency department. On physical examination in the ER, she was normotensive and afebrile with a blood pressure of 127/81 mmHg, heart rate of 86/minute, temperature of 98.8 °F, and respiratory rate of 16/minute with a saturation of 94% on room air. The lung examination demonstrated normal vesicular breathing with bronchial breathing and decreased air entry on the middle and lower left side of the chest. The rest of the examination was unremarkable.

**Investigations**

A CT scan was performed in the ER, which demonstrated two cavitary lesions in the lingula and left lower lobe, raising concerns for aspiration pneumonia. The CT scan also showed bilateral bilobed lungs (Figure 1), heart apex directed towards the left side, left bronchus coursing below the left pulmonary artery (Figure 2), vertical bronchi bilaterally (Figure 3), midline liver (Figure 4), right-sided polysplenia, multiple aberrant nodules of splenic tissue in the right upper quadrant (Figure 5), the azygos continuation of inferior vena cava (Figure 6), and aberrant right subclavian artery (Figure 7), characteristic of heterotaxy. Tuberculosis test (QuantiFERON-TB Gold), beta-D glucan test, and coronavirus disease 2019 (COVID-19) polymerase chain reaction (PCR) nasal swab were sent to rule out tuberculosis, fungal pneumonia, and COVID-19 pneumonia. Blood cultures were also sent.
FIGURE 3: Coronal chest CT image demonstrating left vertical bronchus
CT: computed tomography

FIGURE 4: Axial abdominal CT image showing midline liver
CT: computed tomography
FIGURE 5: Axial and coronal abdominal CT images
A. Axial abdominal CT image. B. Coronal abdominal CT image demonstrating polysplenia

CT: computed tomography

FIGURE 6: Lateral abdominal CT scan image showing the azygous continuation of inferior vena cava

CT: computed tomography
FIGURE 7: Axial chest CT scan

A. Aberrant right subclavian coursing posterior to the esophagus. B. The first branch of the aortic arch is the right common carotid artery, then the left common carotid artery, then the left subclavian, and finally the right aberrant subclavian.

CT: computed tomography

Treatment

The initial diagnosis was "lung abscess secondary to aspiration pneumonia." Hence, she was empirically started on piperacillin-tazobactam and vancomycin. A pulmonology consult was obtained and bronchoscopy was planned in case of a worsening condition. The QuantiFERON-TB Gold test, beta-D glucan test, and COVID-19 PCR nasal swab all came back negative. After 48 hours, the creatinine was deranged and dose adjustment for vancomycin was required. After obtaining negative blood cultures on the fifth day, vancomycin was discontinued and piperacillin-tazobactam was continued.

Outcome and follow-up

The patient’s cough subsided and creatinine improved. She had no further episodes of fever. She was discharged on day 10 after optimizing her medications. At discharge, she was advised to follow up with pulmonology along with referral to the cardiology clinic for an ECG and echocardiography to perform an assessment of heart disease associated with HS.

Repeat chest X-ray performed one month after the clinical improvement revealed the resolution of cavities. About three months after the initial episode of pneumonia, she again presented to the ER with complaints of cough and lethargy. Chest CT scan demonstrated a new cavitary lesion on the left lower side of the chest. She was managed with the same antibiotics as before and discharged after seven days. Currently, she is doing better and awaiting cardiac workups such as ECG and echocardiogram.

Discussion

HS is linked with a higher risk of community-acquired severe bacterial infections and increased mortality rate, regardless of the presence or absence of the spleen [6]. It is clear that the absence of splenic function due to asplenia in right atrial isomerism (RAI) predisposes to a higher risk of bacteremia. However, data regarding infection risk in left atrial isomerism (LAI) with polysplenia (qualitative dysfunction of the spleen) is scarce and there is no agreement on the ways to evaluate the splenic function in polysplenia. A systematic review on the incidence of HS associated with bacteremia in children concluded that 79% of cases of bacteremia occurred in HS with anatomic asplenia and 21% in polysplenia. The mortality rate was similar in both groups; 68% in asplenia vs. 60% in polysplenia [7]. This highlights that the hyposplenism in either of the settings carries a higher risk for infections. Moreover, the guidelines for the assessment and antibiotic management for sepsis or infections associated with HS are lacking.

In this patient, the occurrence of recurrent aspiration pneumonia suggests that the causative factor was polysplenia or the presence of multiple nodules of the spleen, which is associated with abnormal splenic function. This suggests that the hypofunctioning spleen or functioning hyposplenism in such patients makes them more susceptible to infections due to their weak immunological response. Unrecognized ciliary dysfunction causing poor secretion clearance also contributes to it. Moreover, the presence of bilobed lungs and vertical left bronchus (similar to the right) gives an equal predisposition to left-sided aspiration pneumonia.

Recurrent episodes of pneumonia in our young patient suggested the presence of impaired immune response and ciliary dysfunction. The cause remains the absence of IgM memory B cells predisposing individuals to repeated infections with encapsulated organisms like Streptococcus pneumoniae, Neisseria meningitidis, and...
have revealed that two-thirds of problems in left isomerism are manageable with biventricular repair. The In contrast to right isomerism, cardiac problems are milder. Among the prevalent cardiac diseases, studies the esophagus causing dysphagia and can develop aneurysmal dilatation called diverticulum of Kommerell. is about 80%). The aberrant right subclavian artery is another feature and can result in the compression of interrupted inferior vena cava with the azygos or hemiazygos continuation of inferior vena cava (incidence “isomeric atria” is used for both. As a result, both atria receive the pulmonary veins. It is associated with normally on the left side. The atrial appendages on both sides resemble the left atrium, and the term LAI refers to the similar appearance of the structures on both sides of the body to the ones that develop normally on the left side. The atrial appendages on both sides resemble the left atrium, and the term “isomeric atria” is used for both. As a result, both atria receive the pulmonary veins. It is associated with interrupted inferior vena cava with the azygos or hemiazygos continuation of inferior vena cava (incidence is about 80%). The aberrant right subclavian artery is another feature and can result in the compression of the esophagus causing dysphagia and can develop aneurysmal dilatation called diverticulum of Kommerell. In contrast to right isomerism, cardiac problems are milder. Among the prevalent cardiac diseases, studies have revealed that two-thirds of problems in left isomerism are manageable with biventricular repair. The remaining presents with univentricular atrioventricular connection [1]. The sinus node, normally located in
the right atrium, is absent, leading to sinus nodal dysfunction and frequent episodes of atrial flutter and fibrillation. Such patients are more susceptible to conduction defects like atrioventricular block, intraventricular conduction delay, and atrioventricular nodal reentry tachycardia [16,17]. Moreover, due to a concordance between cardiac and pulmonary defects, LAI has bilaterally symmetrical bilobed lungs and hyperatrial bronchi. Hyperatrial bronchi refer to bronchi originating and coursing inferior to the pulmonary artery.

Extracardiac manifestations are various and more common in LAI. Polysplenia is usual. Similar to asplenia, it is hypofunctioning and responsible for producing a weak immune response. It is the quality of function that is relevant rather than the quantity of spleen. However, studies on this are scarce, and no definite evidence is available. The risks of infection may be decreased than in asplenia; still, it can result in lethal sepsis [4]. In most cases, the liver is midline and transverse. In such patients, the presence of jaundice must raise suspicion for biliary atresia, which occurs in 10% of cases [18]. The stomach has variable locations and is usually found on the right side. Tracheoesophageal fistula, duodenal atresia, annular pancreas, and anal atresia (in atresia) are other abnormalities of the gastrointestinal tract. Intestinal malrotation with or without volvulus is a widely prevalent finding in LAI, and some form of it is found in around 70% of cases [4]. The capricious structural defects in the gastrointestinal tract generate atypical chest pains, aspiration pneumonia, and recurring intestinal obstruction. Genito urinary defects that result include horse-shoe shaped, hypoplastic, dysplastic, or absent kidney and urinary tract abnormalities, predisposing to recurrent infections, hydronephrosis, and deranged renal function.

**Investigations and management**

Any patient presenting in the ER with fever requires a complete blood count with differentials, C-reactive protein, and blood culture. The most significant investigation in patients presenting with infections and HS including polysplenia should involve assessing the functioning of the spleen and can help in identifying the bacterial source behind the bacteremia and deciding the type and duration of antibiotics for the management. Ultrasound or CT scan cannot be utilized solely to assess splenic functioning. Some other tests should be employed, which could have been beneficial in our patient. These include the detection of Howell-Jolly bodies in the peripheral blood smear by light microscopy or quantification via flow cytometry. Technetium-labeled scintigraphy or pitted erythrocyte count measures the erythrocytes not cleared by the spleen. Thus, the percentage of Howell-Jolly bodies in the blood quantifies the degree of impairment of the spleen and can be a pivotal test for defining the severity and recurrence of the disease, ultimately helping to decide on the duration and prophylaxis related to antibiotics.

In addition, some other tests such as ECG, transthoracic echocardiography, cardiac catheterization, chest X-ray, abdominal ultrasound, upper gastrointestinal contrast studies, and CT/MRI scan should be done in all newly diagnosed HS patients after the discharge. Angiography is challenging to perform in LAI; thus, a percutaneous approach through neck veins is used. Detailed family history, along with genetic testing, may be required in a few cases. The routine prenatal screening ultrasonography and fetal echocardiography form the hallmarks for the prenatal diagnosis. Fetal MRI may be required in some cases.

In the case of hypofunctioning spleen, appropriate and timely immunization and antibiotic prophylaxis are crucial. All the data regarding it stems from the sickle cell disease (SCD) population. Therefore, all patients are recommended prophylactic antibiotic treatment with penicillins from the time of diagnosis until they reach at least five years of age. For adults, lifelong antibiotic therapy is recommended after sepsis or severe infections. Moreover, immunization against all encapsulated organisms such as pneumococcal, meningococcal, and Haemophilus influenzae should be done. Seasonal influenza vaccination is also mandated. The role of prophylactic antibiotics in asplenia is firmly established and follows the guidelines for SCD. The increased risk of bacteremia with polysplenia, though present, is not highly prevalent. There are no studies that have evaluated the risk amelioration of bacterial infections in the polysplenia population. Therefore, antibiotic prophylaxis is not routinely recommended [6].

Early diagnosis and recognition of abnormalities are essential for the effective management and prevention of complications in all HS patients. Therefore, other complications should be evaluated for management as well. Surgical correction is the fundamental step in managing HS, though, in LAI, anatomic disturbances can present a hurdle. In biliary atresia, repair by Roux-en-Y Kasai operation or a liver transplant may be required. Intestinal malrotation requiring a pre-emptive laparoscopic LADD procedure remains controversial; however, since emergency laparotomy is associated with significant morbidity, offering the procedure to those with malrotation remains reasonable [19,20]. Biventricular repair is done for cardiac abnormalities, while the Fontan procedure is performed in case of univentricular cardiac defects [21]. Most procedures are palliative in nature, as normal anatomy cannot be fully and successfully restored by surgical procedures.

**Prognosis**

The prognosis in LAI is greatly influenced by the type and number of associated cardiac problems, and in the absence of such defects, it is quite promising. The mortality is approximately 50% in LAI, mostly attributed to the presence of heart block [22]. Bacterial sepsis in HS is associated with a mortality rate of 44% [23]. Prenatal diagnosis is especially useful in asplenia patients as around 10% of cases can lead to stillbirths.
as compared to 6% in polysplenia (where the main cause remains bradyarrhythmias). One study has demonstrated that the postnatal death rate in live-borns asplenia babies is 43% vs. only 15% in polysplenia babies [24]. The major cause in both remained cardiac. The morbidity is significantly high owing to the propensity for several complications together with deranged anatomy. This disease confers a dilemma in both its diagnosis and management and remains a challenging condition to be dealt with.

Conclusions

We discussed an unusual presentation that led to the diagnosis of a rare underlying condition termed HS/left isomerism. Hyposplenism can occur in HS patients regardless of spleen anatomy or presence. The bacteremia and recurrent pneumonia associated with polysplenia in such patients raise questions regarding prophylactic antibiotics and immunization similar to asplenia, which require further studies involving measures such as the quantification of splenic function. More studies to determine the specific immunological disruptions occurring in HS are also required. The question of whether the guidelines for the management of hyposplenism in polysplenic HS should be similar to those for non-heterotaxic hyposplenism and heterotoxic asplenic patients is pertinent.

This condition can have variable presentations and remain undiagnosed for a long time unless congenital heart defects are severe enough to warrant at-birth investigations. This adds to the morbidity and mortality associated with it. To prevent this, early diagnosis and management are essential. In cases of high suspicion and those with a family history, early CT scan and echocardiography can provide timely treatment and prevention of complications. A multidisciplinary team consisting of internists, gastroenterologists, cardiologists, radiologists, surgeons, and nurses is essential to provide timely and imminent care.

Additional Information

Disclosures

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References

1. Kim SJ: Heterotaxy syndrome. Korean Circ J. 2011, 41:227-32. 10.4070/kcj.2011.41.5.227
2. Sempou E, Khokha MK: Genes and mechanisms of heterotaxy: patients drive the search. Curr Opin Genet Dev. 2019, 56:34-40. 10.1016/j.gde.2019.05.003
3. Mishra S: Cardiac and non-cardiac abnormalities in heterotaxy syndrome. Indian J Pediatr. 2015, 82:1135-46. 10.1007/s12098-015-1925-x
4. Kothari SS: Non-cardiac issues in patients with heterotaxy syndrome. Ann Pediatr Cardiol. 2014, 7:187-92. 10.4103/0974-2069.140854
5. Anderson RH, Brown NA, Meno C, Spicer DE: The importance of being isomeretic. Clin Anat. 2015, 28:477-86. 10.1002/ca.22517
6. Chiu SN, Shao PL, Wang JK, et al.: Severe bacterial infection in patients with heterotaxy syndrome. J Pediatr. 2014, 164:99-104. 10.1016/j.jpeds.2013.08.051
7. Loomba RS, Geddes GC, Basel D, et al.: Bacteremia in patients with heterotaxy: a review and implications for management. Congenit Heart Dis. 2016, 11:537-47. 10.1111/chd.12595
8. Kruetzmann S, Rosado MM, Weber H, et al.: Human immunoglobulin M memory B cells controlling Streptococcus pneumoniae infections are generated in the spleen. J Exp Med. 2003, 197:859-45. 10.1084/jem.20022020
9. Chiu SN, Shao PL, Wang JK, et al.: Low immunoglobulin M memory B-cell percentage in patients with heterotaxy syndrome correlates with the risk of severe bacterial infection. Pediatr Res. 2016, 79:271-7. 10.1038/pr.2015.221
10. Shapiro AJ, Tolleson-Rinehart S, Zariwala MA, Knowles MB, Leigh MW: The prevalence of clinical features associated with primary ciliary dyskinesia in a heterotaxy population: results of a web-based survey. Cardiol Young. 2015, 25:752-9. 10.1016/j.cly.2014.05.003
11. Belmont JW, Mohapatra B, Towbin JA, Ware ML: Molecular genetics of heterotaxy syndromes. Curr Opin Cardiol. 2004, 19:216-20. 10.1097/00001573-200405000-00005
12. Lin AE, Ticho BS, Houde K, Westgate MN, Holmes LB: Heterotaxy: associated conditions and hospital-based prevalence in newborns. Genet Med. 2000, 2:157-72. 10.1097/00001573-200005000-00002
13. Ryan AK, Izipisiá Belmonte JC: Establishing a left-right axis in the embryo. JUBMB Life. 2000, 50:1-11. 10.1080/15216540050176520
14. Shiraiishi I, Ichikawa H: Human heterotaxy syndrome - from molecular genetics to clinical features, management, and prognosis. Cire J. 2012, 76:2066-75. 10.1253/circjc.cj-12-0957
15. Ticho BS, Goldstein AM, Van Praagh R: Extracardiac anomalies in the heterotaxy syndromes with focus on anomalies of midline-associated structures. Am J Cardiol. 2000, 85:729-34. 10.1016/s0002-9149(99)00849-8
16. Ozawa Y, Asakai H, Shiraga K, Shindo T, Hirata Y, Hirata Y, Inuzuka R: Cardiac rhythm disturbances in heterotaxy syndrome. Pediadi Cardiol. 2019, 40:909-15. 10.1007/s00246-019-02087-2
17. Czosek RJ, Haaning A, Ware SM: A mouse model of conduction system patterning abnormalities in heterotaxy syndrome. Pediatr Res. 2010, 68:275-80. 10.1203/PDR.0b013e3181ee0028
18. Mirza B, Iqbal S, Sheikh A: Biliary atresia associated with polysplenia syndrome, situs inversus abdominus, and reverse rotation of intestine. APSP J Case Rep. 2012, 5:14.
19. Pickett CR, Dicken B, Rebyeka IM, Ross DR, Ryerson LM: Heterotaxy syndrome: is a prophylactic Ladd procedure necessary in asymptomatic patients?. Pediatr Cardiol. 2013, 34:59-63. 10.1007/s00246-012-0385-6
20. Yu DC, Thiaagarajan RR, Laussen PC, Laussen JP, Jaksic T, Weldon CB: Outcomes after the Ladd procedure in patients with heterotaxy syndrome, congenital heart disease, and intestinal malrotation. J Pediatr Surg. 2009, 44:1089-95. 10.1016/j.jpedsurg.2009.02.015
21. Agarwal R, Varghese B, Jerudian V, Moses J: The heterotaxy syndrome: associated congenital heart defects and management. Indian J Thorac Cardiovasc Surg. 2021, 37:67-81. 10.1007/s12055-020-00955-y
22. Thomas C, Sawyer SN: Prenatal findings in left atrial isomerism and an overview of heterotaxy syndrome. J Diagn Med Sonogr. 2017, 2:148-54. 10.1177/8756479317743196
23. Prendiville TW, Barton LL, Thompson WR, Fink DL, Holmes KW: Heterotaxy syndrome: defining contemporary disease trends. Pediatr Cardiol. 2010, 31:1052-8. 10.1007/s00246-010-9764-9
24. Escobar-Díaz MC, Friedman K, Salem Y, et al.: Perinatal and infant outcomes of prenatal diagnosis of heterotaxy syndrome (asplenia and polysplenia). Am J Cardiol. 2014, 114:612-7. 10.1016/j.amjcard.2014.05.042