

Isolated left lung bronchiectasis: a rare presentation of H type tracheoesophageal fistula

Jiledar Rawat, Gurmeet Singh, Sudhir Singh

Department of Pediatric Surgery, King George's Medical University, Lucknow, UP, India

Correspondence to: Dr. Sudhir Singh. Associate Professor, Department of Pediatric Surgery, King George's Medical University, Lucknow, UP, India. Email: drsudhir_singh25@yahoo.in.

Abstract: H type tracheoesophageal fistula (TEF) is a very uncommon congenital entity. Diagnosis of these cases are usually delayed and cases are treated as respiratory tract infections. We here reporting a four years old girl, presented to us with recurrent respiratory tract infection. On examination of baby decreased chest movement, and on auscultation of chest there was decreased breath sounds on left side was found. Chest X rays suggest esophagus was grossly dilated, and air filled. Contrast esophagogram suggestive of H type tracheoesophageal fistula and bronchiectasis changes on left lung. Contrast Enhanced Computer Tomography suggest bronchiectasis changes on left lung. The diagnosis of H type tracheoesophageal fistula (TEF) was made by rigid bronchoscopy. Successful surgical repair of H type TEF required right thoracotomy approach as fistula was quite low laying. Tracheo-bronchoscopy (especially rigid) is an important investigation due to the better visualization, in doubtful cases and to locate the exact location of fistula. H type TEF above T2 should be approached through cervical approach whereas lower fistula should be always approached by right posterolateral thoracotomy. Low laying fistula is extremely rare finding along with bronchiectasis changes in left lung. Therefore, this case report describes a rare case and review of literature.

Keywords: H type tracheoesophageal fistula (TEF); bronchiectasis; child

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Introduction

Tracheoesophageal fistula (TEF) constitute about 1 in 100,000 live births with H-type TEF is 4–5% of all, and has a variable clinical presentation and approach of surgical correction (1). The presentations are recurrent respiratory symptoms due to aspiration pneumonitis while feeding, cyanosis, and abdominal distension and usually diagnosed in late infancy or childhood (2). The common surgical approach is right cervicotomy, if the fistula is above the T2 vertebra and rarely thoracotomy required if fistula is below (3). We hereby report a case of H-type TEF presented to us at the age of 4 years with recurrent chest infection and isolated left side lung bronchiectasis require a thoracic approach for surgical correction as it was quite low. We present the following case in accordance with the CARE reporting checklist. Written informed consent was obtained from the patient for publication of this study and any accompanying images.

Case presentation

A 4-year-old female child presented with symptoms of recurrent pneumonitis since birth. There was a history of nasal and oral regurgitation of feeds, and required recurrent admission for pneumonia and left side bronchiectasis. The patient was referred to our hospital for the non-improvement of problem. On examination, respiratory rate was 45 per minute, decreased chest movements on the left side. On chest auscultation, there were decreased breath sounds on the left side. There were no other anomalies detected. Chest X-rays performed (*Figure 1A*) showed esophagus was grossly dilated, and air-filled, left side of



Figure 1 (A) Chest X-rays PA view showing the esophagus was grossly dilated, and air-filled, left side of chest show tram tracking the appearance of dilated lower lobe bronchi suggestive of bronchiectasis. (B) Contrast-Enhanced Computer Tomography showing bronchiectatic changes on the left lung.



Figure 2 Rigid bronchoscopy showing H type TEF (tracheoesophageal fistula).

chest show tram tracking appearance of dilated lower lobe bronchi suggestive of bronchiectasis. Esophagogram suggestive of H type fistula as contrast seen in the trachea. Contrast-Enhanced Computer Tomography (*Figure 1B*) was suggestive of bronchiectasis changes in the left lung field. Rigid bronchoscopy was performed under general anaesthesia and a diagnosis of H-type TEF was made at the level of T3 thoracic vertebra with purulent fluid was found in the left bronchus, aspirated and lavage done (*Figure 2*). TEF was repaired through the right thoracotomy extra pleural approach in the same setting of anaesthesia. The girl improved and was asymptomatic in the post-procedure duration. A follow-up of 8-month patient doing well with occasional cough. Radiography of thorax (including plain chest X-ray, contrast oesophagogram, and computer tomography of the thorax) demonstrates resolution of pulmonary bronchiectasis and there was a decrease in oesophageal diameter.

Discussion

The condition H type TEF, may present mild symptoms of repeated aspiration and pneumonia to life-threatening events. The symptoms may occur in the neonatal period to the oldest reported patient was 79 years old (4). The main symptoms of an H-TEF include recurrent pneumonitis, cyanosis, and choking on feeding and abdominal distension. Presentation like left lung isolated bronchiectasis in four years old girl child was not reported till date as found in our case. In our case, the fistula was quite low and towards left bronchus resulting in repeated aspiration and bronchiectasis. On plane chest X-ray, diagnosis is confirmed by an air-filled dilated esophagus due to constant inflation air from TEF in the esophagus (Figure 1A). In the case of an air-filled dilated esophagus, a possibility of TEF should always be considered. The mainstay of diagnosis is a water-soluble contrast oesophagogram which should be performed early in case of suspicious H-type fistula, and bronchoscopy should be done if the contrast esophagogram not show the fistula (5). There is a risk of aspiration pneumonia in contrast esophagogram. Rigid bronchoscopy is a better procedure to see the posterior wall of the trachea and can be gently probed to see the opening of a collapsed fistula. On rigid bronchoscopy, fistula can be cannulated with a guidewire, which aid identification of the fistula and repair. Rigid bronchoscopy helps determine the level of fistula, presence of unusual variants, e.g., double fistula, trifurcation fistula, presence and severity of tracheobronchitis, and position of the aortic arch (5). Most surgeons agreed that fistula at or above T2 should be approached through the cervical approach and fistula below this should be ligated by the thoracic approach (3,6-8). In our case, the fistula was at the T3 and we approached the fistula through the right thoracotomy. When the fistula was surgically divided our approach is to divide the TEF on the esophageal side to have sufficient amount tissue for good repair over posterior wall of the windpipe. The esophageal side has a tensionfree closure due to the adequate width of the lumen and the interposition of vascularized tissue between repaired fistulas can prevent recurrence. Reported complications were fistula recurrence, esophageal stricture tracheomalacia, and left recurrent laryngeal nerve paralysis (9,10). Mortalities were usually rare if the condition is diagnosed and treated early.

Conclusions

The presentation of H type TEF may varied. Rigid bronchoscopy is diagnostic of H type TEF and surgical repair is curative of these anomalies.

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Footnote

Reporting Checklist: The authors have completed the CARE reporting checklist. Available at http://dx.doi.org/10.21037/jxym-20-57

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi. org/10.21037/jxym-20-57). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee(s) and with the Helsinki Declaration (as revised in 2013). Written informed consent was obtained from the patient for publication of this study and any accompanying images.

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