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IP Journal of Surgery and Allied Sciences

Journal homepage: https://www.jsas.co.in/



Case Report

Association of gastric duplication cyst with isolated oesophageal atresia and one additional floating rib — A rare presentation

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ARTICLE INFO

Article history: Received 08-12-2022 Accepted 17-12-2022 Available online 23-01-2023

Keywords:
Duplication cyst
Enteric duplication
Gastric duplication cyst
Oesophageal atresia
and Floating rib

ABSTRACT

A gastric duplication cyst is not a common anomaly of the alimentary tract. The most common site of alimentary tract duplication is the ileum. We found a non-communicating gastric duplication cyst, along with the greater curvature of the stomach during feeding gastrostomy for isolated Oesophageal atresia. We are reporting a gastric duplication cyst associated with isolated Oesophageal atresia and left side one additional floating rib. This is the third case report of gastric duplication cyst with Oesophageal atresia in English literature.

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1. Introduction

Enteric duplication cyst is a rare entity. It can be found from mouth to rectum throughout the gastrointestinal tract. Ileum is the most common(30-35%) site of enteric duplication. Incidence of duplication in alimentary tract is 20% in oesophagus, 13% in colon, 10% in jejunum, 7% in stomach and 5% in duodenum. ^{1,2} Most common presentation of Gastric duplication is non-communicating, cystic pattern. Most common site of gastric duplication is along the greater curvature of stomach. ³ Gastric duplication represents 4-7% in whole enteric duplication. Near about 67% of gastric duplication are diagnosed up to the first birthday of baby. ⁴ Here, we are reporting acase of cystic duplication of stomach association with isolated Oesophageal atresia and one additional floating rib. This is an extremely rare

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association.

2. Case Report

An 1850 gram male child, delivered at full-term, vaginal without any obstetric complication in hospital. He had complaints of frothing from mouth and regurgitation of feed. He was referred to our hospital after optimization. He was admitted here. On clinical examination baby has upper abdominal fullness and excessive frothing from mouth. Red rubber catheter (10 french) introduced to rule out Oesophageal atresia but could not be negotiated beyond 10 cm. There was no pallor and no jaundice and no other visible abnormality. Babygram with red rubber catheter showed gas less abdomen except one irregular shape area at left hypochondriac region. There was one additional floating rib present left side (Figure 2). Other skeletal system was normal. Our differential diagnosis was isolated Oesophageal

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atresia with pyloric atresia. USG abdomen showed two heterogeneous cystic structures with thick walls (3.8x3.0 cm and 1.7x1.6 cm) were noted at left hypochondriac region. These lesions were abutting and compressing the stomach. Spleen and kidney were seen separately and normal. These findings were concluded foregut duplication or extra lobar pulmonary sequestration. After getting routine blood reports baby was taken for surgery on live day second. Baby was operated under general anaesthesia with endotracheal intubation in supine position and bolster beneath the both shoulders to make slight extension of neck. After painting and draping, mid line upper abdominal incision was given. Stomach was delivered through wound. There was a palpable cystic swelling at middle of greater curvature of stomach (Figure 1). This cystic swelling was excised and stomach was repaired and feeding gastrostomy done. Cyst was not communicating with stomach cavity. Abdominal wall repaired and feeding gastrostomy tube was fixed from abdominal wall. Left cervical esophagostomy was made to drain out the saliva from oral cavity. Excised cystic swelling was sent for histopathological examination. Histopathology of specimen was showing gastric mucosa with muscle layers, which proposed the gastric duplication cyst.



Fig. 1: Showed duplication cyst along the greater curvature.

Sham feeding was started on post-operative day two. Feeding through gastrostomy tube was started on seventh post-operative day. Baby tolerated feed through gastrostomy



Fig. 2: X-ray showed gasless abdomen indicating isolated oesophageal atresia and one additional floating rib left side.



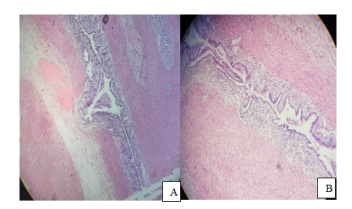
Fig. 3: Showing follow up photograph of baby having feeding gastrostomy tube and cervical oesophagostomy.

S. No. Author name with Site of gastric Histopathological finding Gender Associated presenting year duplication Oesophageal atresia (EA) 1 Hemalatha et al. 1980⁵ Tubular duplication EA distal TEF Pyloric type gastric mucosa along the greater with aberrant pancreatic curvature tissue with smooth muscle tissues 2 Piplani R.et al.2018⁶ Cystic duplication of EA distal TEF Gastric lining including all Male greater curvature of layers of the gastrointestinal the stomach tract with Heterotopic pancreatic tissue 3 Reporting case Dr.Dinesh Cystic duplication of Isolated EA All smooth muscle layers Male

Table 1: Table 1- table containing reported cases of gastric duplication associated with oesophageal atresia

greater curvature of

the stomach



Kumar Barolia et. al.]

Fig. 4: H & E stained histopathology slides of excised duplication cyst-showed gastric mucosa and muscle layers.

tube and his journey was uneventful after surgery in six month follow up (Figure 3).

Showing follow up photograph of baby having feeding gastrostomy tube and cervical oesophagostomy.

3. Discussion

Gastric duplication is a rare entity. Greater curvature is the most common site of gastric duplication. Antrum is the dominant site of greater curvature. Cystic form is the major presentation. ^{4,7,8} In our case duplication cyst was present along the greater curvatureat antrum non-communicating.

Cardinal features of enteric duplication are (a) developed smooth muscle coat (b) lining epithelium comparable to intestinal mucosa (c) sharing common blood supply (d) communicating or non-communicating with alimentary tract. Gastric duplication can present with palpable swelling in upper abdomen. Some time it may present with recurrent non-bilious vomiting. ^{4,9} Gastric duplication cyst can be treated by drainage in jejunum mostly in non-communicating cyst. Formation of common cavity of cyst and stomach can be done in communicating cyst. Complete excision of duplication cyst and repair is the treatment of

choice. 10

Spataru R.I. et al. (2015) tabulated all the Reported cases of esophageal atresia associated with foregut duplications in their article. Only seventeen cases of esophageal atresia associated with foregut duplications were reported. Oesophageal duplication was the common site of fore gut duplication. Gastric duplication was less common in association with oesophageal atresia. ¹¹ Only one case was reported till 2015. This first case was reported by Hemalatha et al. in 1980. They reported tubular gastric duplication along greater curvature associated with oesophageal atresia and tracheoesophageal fistula (EA-TEF). ⁵

and containing gastric

mucosa

Second case of gastric duplication with oesophageal atresia was reported by Piplani R. et al. (2018). The gastric duplication wascystic along greater curvature associated with oesophageal atresia and tracheoesophageal fistula (EATEF). ⁶

Here we are reporting the third case of gastric duplication along the greater curvature associated with oesophageal atresia without tracheoesophageal fistula (EA without TEF). This is the case of gastric duplication with associated with oesophageal atresia without tracheoesophageal fistula (isolated oesophageal atresia) and additional one rib.

4. Conclusion

Gastric duplication with association of oesophageal is an extremely rare entity. To the best of our knowledge, this is the first case report in English literature of isolated Oesophageal atresia with association of gastric duplication cyst and one additional floating rib.

5. Source of Funding

None.

6. Conflict of Interest

None.

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Cite this article: Singh AP, Barolia DK, Bathia HV, Parmar VH, Mehta BA, Mehta S. Association of gastric duplication cyst with isolated oesophageal atresia and one additional floating rib — A rare presentation. *IP J Surg Allied Sci* 2022;4(4):145-148.