Journal of Mind and Medical Sciences

Volume 8 | Issue 1

Article 22

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Iozsa, Dan-Alexandru; Puscasu, Maria; Cirstoveanu, Catalin Gabriel; Constantin, Vlad Denis; Dumitriu, Anca Silvia; Paunica, Stana; and Ionescu, Nicolae Sebastian () "A pedunculated esophageal chondromatous hamartoma in a child," *Journal of Mind and Medical Sciences*: Vol. 8 : Iss. 1 , Article 22. DOI: 10.22543/7674.81.P161166

Available at: https://scholar.valpo.edu/jmms/vol8/iss1/22

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This case presentation is available in Journal of Mind and Medical Sciences: https://scholar.valpo.edu/jmms/vol8/ iss1/22

https://scholar.valpo.edu/jmms/ https://proscholar.org/jmms/ ISSN: 2392-7674

A pedunculated esophageal chondromatous hamartoma in a child

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ABSTRACT

Esophageal tumors are uncommon in pediatric population and most of them are benign. Esophageal hamartomas have been reported extremely rare in children. These can present as intramural tumors of the esophageal wall or as polyps. Dysphagia is the main symptom described in their case, but other specific symptoms are also reported. Such symptoms encountered in clinical practice are represented by obstructive apnea episodes and bradycardia, poor weight gain, epigastric or retrosternal pain, hematemesis or melena, dysphonia or tracheal sounds. Diagnosis can be delayed due to the insidious onset and non-specific symptoms, therefore patient's compliance to follow-up and broad, careful evaluation are mandatory. Preoperative imagistic assessment is extremely important for a precise definition of the tumor's anatomical relations, especially when facing large tumors located in the posterior mediastinum. Herein we report a case of a large chondromatous polypoid hamartoma of the esophagus in a 9 years old boy, emphasizing over the diagnosis and surgical challenges we have met, along with comments on illustrative similar cases reported in the literature.



Category: Case Presentation

Received: September 11, 2020 Accepted: December 16, 2020

Keywords:

chondromatous hamartoma, esophageal tumor, dysphagia, children

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Introduction

The posterior mediastinal masses in the pediatric population have a neurogenic origin in 90% of the cases [1]. Less than 5% of all tumors detected during childhood or adolescence originate in the gastrointestinal tract, from which a small part arises from the esophagus, 70% of them being benign [2].

Regarding their configuration in relation with the esophageal lumen, esophageal polypoid lesions are very unusual endoscopic findings in children, being found by Septer et al. in only 0.14% of the 9,438 pediatric patients who underwent routine upper endoscopies [3].

Hamartomas are non-neoplastic developmental malformations. They are made of a variety of cytologically normal, mature cells and tissues representative for the anatomical region they arise from. The architectural pattern is chaotic, displaying a dominant component. Hamartomas can be found arising in any part of the body, but most of them are identified in the lungs, liver, pancreas and spleen, and extremely rarely in the esophagus, in both pediatric and adult population [4].

Because of their rarity, being reported in isolated cases or small series, literature is not consistent enough regarding the management of the suspected cases of esophageal tumors in children.

We report the case of a large pedunculated esophageal chondromatous hamartoma in a young boy. Further on, we bring into discussion the history, the clinical presentation, the work-up, the surgical approach and the histopathologic diagnosis of this extremely rare entity.

Case Report

A 9-year-old boy was admitted to our clinic for stridor and mild dysphagia. The informed consent was signed by

To cite this article: Dan-Alexandru Iozsa, Maria Puscasu, Catalin Gabriel Cirstoveanu, Vlad Denis Constantin, Anca Silvia Dumitriu, Stana Paunica, Nicolae Sebastian Ionescu. A pedunculated esophageal chondromatous hamartoma in a child. *J Mind Med Sci.* 2021; 8(1): 161-166. DOI: 10.22543/7674.81.P161166

the patient's parents, after having discussed the reasonable disclosure and the therapeutic possibilities [5]. The onset of the patient's symptoms was 2 years prior to the present admission, when he had begun to exhibit recurrent upper airway infections and chronic coughing, being treated by the local family physician without any additional work-up.

Progressive dysphagia and stridor subsequently came into the clinical picture, so he presented to the Pediatrics Department, where asthma was suspected, and a pulmonary X-ray was done, revealing a large mass presumably in the posterior mediastinum. Therefore, the evaluation was continued through an esophagogram (Figure 1), which identified a large cervical-thoracic esophageal mass. In order to better define the tumor, including its relationship with the esophageal wall, MRI scans (Figure 2) and CT scans (Figure 3) were done.



Figure 1. The esophagogram revealing a large cervical-thoracic esophageal mass.



Figure 2. The MRI aspect of the esophageal tumor.



Figure 3. CT scan showing the esophageal tumor.

A preoperative esophagoscopy was performed, showing the intraluminal protrusion of the esophageal tumor, retained food remnants, but no additional information was obtained in order to describe the mass, since its size was making the evaluation very difficult (Figure 4).

The patient underwent surgery via right thoracotomy. After the mediastinal inspection, oblique esophagotomy was done. A massive cervical-thoracic pedunculated intraluminal tumor ($102 \times 52 \times 48 \text{ mm}$) was noted in the cervical-thoracic esophagus lumen, which was significantly dilated from the T4 level upward to the lower cervical region (Figure 5). The mass was found to have arisen from the esophageal wall at the distal cervical segment.

The complete excision and the esophageal preservation were possible without the cervical approach. Dissection revealed that the tumor has developed in the esophageal wall – no invasion was noted in the stretched mucosa and muscularis of the esophagus. The histopathological diagnosis was chondromatous polypoid hamartoma (Figure 6).



Figure 4. The endoscopic aspect of the lesion.



Figure 6. The pathological exam – gross and microscopic aspect of the esophageal chondromatous hamartoma.

The one-year follow-up did not reveal any complications, the follow-up esophagogram showed the persistence of a mild esophageal dilatation (Figure 7), but no swallowing disorders were noted. A significant 6 kg weight gain was observed – from 39 Kg (percentile 86.7) to 45 (percentile 92.4).



Figure 7. The follow-up esophagogram showing the persistence of a mild esophageal dilatation.



Figure 5. The diagram and the intraoperative aspect of the large, intra-luminal tumor.

Discussions

The division process of the primitive foregut underlies many ways of embryo pathogenesis. The trachealbronchial bud arises from the ventral part and grows caudally separating from the future esophagus. A disturbed tracheal-esophageal separation process sets up the origin of the congenital tracheal-esophageal fistulas or foregut duplication cysts – which, in exceptional cases, may associate esophageal atresia [6,7]. In the same course, respiratory tissue sequestration in the esophageal wall during the tracheal-esophageal separation process may explain the origins of congenital esophageal stenoses due to tracheal-bronchial remnants or chondromatous hamartomas (either polypoid or intramural) if the heterotopic tissue is organized in tumor-like lesions [8-10].

Dysphagia is the main manifestation of the esophageal tumor overall. However, clinical presentations may vary depending on their benign or malignant nature. A careful pre- and intraoperative assessment is necessary to prevent medical errors, and the therapeutic decision should be taken following the discussion of the alternatives with the child's legal representative and signing the informed consent [11,12]. In children, benign tumors, having a slowly growing pattern, often lead to airway compression and micro-aspirations, therefore a chronic respiratory distress picture (recurrent airway infections, wheezing, shortness of breath) is common. By contrast, some benign tumors generate the total obstruction of the digestive tract since birth [11]. Malignant esophageal tumors of childhood present signs of nutritional deficiency such as weight loss, anemia or dehydration, more often secondary to dysphagia [2]. Decreased serum melatonin may be a relevant biomarker for malignancy [12]. Our patient's history is similar to the case presented by Samad, et al., where the diagnosis of a pedunculated lipoma of the esophagus has been delayed for 2 years, the tumor's clinical presentation dissimulating asthma [13]. On the other hand, Coury, et al.

report a neonate whose initial symptoms (dyspnea and cyanosis) were disguised by the presumption of severe gastro-esophageal reflux and peripheral pulmonary stenosis and the tumor's diagnosis was delayed until the 40th day of life, when due to recurrent obstructive apnea episodes and bradycardia (despite of nasal-jejunal feeding), a laryngoscopy was done revealing a polypoid cervical esophageal hamartoma prolapsing intermittently in the upper airway [14,15]. Other clinical signs reported in the esophageal tumors of childhood are poor weight gain, epigastric or retrosternal pain, hematemesis or melena, dysphonia or tracheal sounds [2]. The malabsorption of micronutrients related to dysphagia, with low serum Zn/Cu ratio may lead to attention deficit spectrum disorders [16]. In our opinion, considering the broad spectrum of subtle signs and the frequency of the gastroesophageal reflux disease symptoms in pediatric population, endoscopic studies of the esophagus and airways should be more seriously taken into consideration as an early diagnostic tool, in case of an inadequate response therapy, in order to exclude unlikely, but threatening conditions.

Dysphagia in children should be carefully assessed, mainly because of the aspiration risk. The primary evaluation tools are video fluoroscopy and fiber optic endoscopy. All swallowing aspects should be checked, from the oro-pharyngeal phase to the esophageal passage of the barium solution [17]. On the other hand, from our experience, we consider that all children (especially toddlers), presenting with respiratory symptoms, should undergo systematically plain X-ray evaluations. These can identify any mediastinal masses much earlier or exclude life-threatening situations such as unsuspected esophageal foreign bodies (for example, button batteries). However, if esophageal masses remain undetected at the initial evaluation, parents' compliance and return for follow-up plays an essential role in the early detection in children and the health professional should never hesitate to consider barium swallowing examination and both endoscopic and digestive endoscopic tools in case of doubtful results or persistent symptoms. Additional means, such as CT and MRI scans, may be required for a better mass description as a differential diagnosis, anatomical relations or preoperative planning [18]. In selected cases, endoscopic ultrasounds and fine needle aspiration biopsies might be considered in the preoperative evaluation and treatment planning, which has the ad

Esophageal hamartomas are extremely rare and there is no commonly accepted guidance for their treatment. The vast underlying molecular network connecting these disorders opened a vast area of research [19]. The surgical attitude may vary widely depending on their aspect – intramural or polypoid, their volume and depth of invasion into the esophageal layers. Throughout the years, minimally invasive methods have gained ground when considering the surgical field [20]. Endoscopic resection using snare coagulation or simple ligation is possible in carefully selected cases [8], with all the advantages of the minimally invasive approach: quick recovery, minimal scaring, reduced pain and postoperative complications [21,22]. If the nature of the tumor is not suitable for a minimally invasive approach, esophagotomy and enucleation may be attempted [23,24], but this may not be possible all the time, therefore esophagectomy and end-toend anastomosis have to be done [25]. In these cases, an ICG tissue angiography is a feasible and reliable technical support in the evaluation of the anastomotic perfusion and for the prevention of any anastomotic leak [24-26].

Similar to other surgical situations, in selected cases, one can use a linear stapler in order to resect a pedunculated esophageal mass [25]. In our case, enucleation was possible due to a relatively simple dissection between the tumor and the esophageal muscularis. The mucosa excess was also removed along with the tumor and esophageal suturing was done in safe conditions. Moreover, in terms of preoperative planning, in large esophageal masses – the accurate evaluation of the position related to the thoracic aperture is an important matter, in order to decide between thoracotomy, cervical approach or both. The accuracy of the surgical procedure ensures a quick recovery, reducing the hospital stay and the associated medical expenses [26].

The differential diagnosis of the esophageal polypoid in childhood may include squamous hamartomas fibrovascular papilloma, giant polyps, lipomas, inflammatory fibroid polyps, glomangiomas, pyogenic granulomas [3] or teratomas [27-29]. Other common causes of benign esophageal masses in children are leiomyomas, papillomas, hemangiomas, lymphangiomas or neurofibromas. More rarely, malignant causes are also reported, the most common being squamous cell carcinomas and adenocarcinomas [2]. However, the most frequent site of chondromatous hamartoma is the lung [9,30], accounting for 3% of all lung tumors in the adult population, its possibility of malignant degeneration toward carcinoma or sarcoma being well-known [31].

Highlights

- ✓ Esophageal tumors in the pediatric population are extremely rare entities with a low incidence of occurrence in the clinical practice of a pediatrician.
- ✓ The diagnosis is delayed by nonspecific symptoms and radiologic findings.
- ✓ The wide range of pathological entities similar to it require extensive planning and interdisciplinary collaboration between the radiologist, oncologist, pediatric surgeon and endoscopist. The surgical treatment should preserve the native esophagus and prevent long term complications.

Conclusions

Esophageal tumors are uncommon in the pediatric population. The diagnosis can be delayed due to their insidious onset of symptoms and their unspecific clinical signs, therefore the patient's compliance to follow-up and broad, careful evaluation are mandatory. The respiratory symptoms and the history or clinical signs of nutritional difficulties should lead to a more careful evaluation of the esophagus.

The preoperative evaluation may be difficult because of the size of the tumors. Because of their rarity along with the virtual space defined by the esophageal lumen, when facing large tumors of the posterior mediastinum in the pediatric population, the surgeon should never hesitate to use preoperative imaging tools for a precise definition of the tumor's anatomical relations.

Conflict of interest disclosure

There are no known conflicts of interest in the publication of this article. The manuscript was read and approved by all authors.

Compliance with ethical standards

Any aspect of the work covered in this manuscript has been conducted with the ethical approval of all relevant bodies and that such approvals are acknowledged within the manuscript.

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