Mediastinal Benign Mature Teratoma in Young Girl, Catastrophic Delayed and Complications of Surgery: A Case Report

Ketut Putu Yasa*, A. A. Christ Tedy Pernmana†, Sri Mahendra Dewi‡

1Cardiothoracic Vascular and Endovascular Division, Department of Surgery, Udayana University, Sanglah Hospital, Denpasar, Bali, Indonesia; †General Surgery Training Program, Department of Surgery, Udayana University, Sanglah Hospital, Denpasar, Bali, Indonesia; ‡Department of Pathology, Udayana University, Sanglah Hospital, Denpasar, Bali, Indonesia

Abstract

BACKGROUND: Teratomas are the most common of germ cell tumor and have different epidemiology, histology, and biology than the others. These tumors occur equally in men and women are generally benign, occur in the mediastinum and compromise approximately 20% of anterior mediastinal masses. All forms of these tumors have a peak incidence in young adults.

CASE REPORT: A huge mediastinal teratoma present in young girl 15 years old, a tumor located in the right chest with mass compression effect to spine, heart, and lung, produce prolong symptoms of dyspnea and chest pain. Delayed clinical presentation influences to post-operative complications such as prolong atelectasis, rupture of tumor mass which spoiled of tumor contents, empyema, and sepsis.

CONCLUSION: Mediastinal mature teratoma with the delayed clinical presentation will give technical difficulties for resection and increase the risk of post-operative complications.

Introduction

Mediastinal teratomas are thought to be the result of pluripotent germ cell that fails to migrate from mediastinum to the gonads during early development. These tumors occur equally in men and women, are generally benign, and consist of at least two of the three embryologic layers: Endoderm, mesoderm, and ectoderm[1]. Teratoma may be mature, well-differentiated, poorly differentiated, immature, or with malignant transformation [2]. A mature teratoma has differentiated into mature somatic structure such as skin, bone, teeth, fat, and epithelial tissue. Most of the symptoms are a result of compression of nearby structures. Treatment of choice for mediastinal teratoma is complete surgical resection[1].

Case Presentation

Young girl 15 years old with chief complain shortness of breath in the past 4 years ago. Shortness of breath felt burdensome in the last 1 month ago; sometimes, the right chest feels painful. Patients feels that her body weight was decreased, and body weight loss was around 5 Kg during last 3 months. Patients with a history of control to cardiothoracic out-patient clinic and already diagnosed with tumor mediastinum 1 year ago, and then plans for surgical removal of the tumor were carried out, but the patient and family refused.

On physical examination, there was bulging of breast on the right chest with no breath sound on the right lung, slight dilatation of chest superficial vein, but no edema on the face and upper extremities (Figure 1). Chest plain X-ray (Figure 2) and computed tomography scan (Figure 3) revealed huge cystic well-defined mass shadow with signs of calcification in the right chest with mass compression effect to spine, heart, and lung. Laboratory findings for tumor marker beta-human chorionic gonadotropin (β HCG) and alpha-fetoprotein (AFP) were normal limit, another laboratory test was normal. Pre-operative clinical diagnosis of suspect mediastinal teratoma was entertained. The patient underwent complete surgical excision of a mediastinal tumor through right posterior lateral thoracotomy.

Intra-operative findings, there was huge solid...
cystic mass fulfill entire of right pleural space, and there was persistent atelectasis, rupture on some area of cystic mass during difficult dissection with spoiled tumor contents material to the operation field (Figure 4a). The complete excised mass consisted of mature squamous epithelium (skin), salivary gland, hair, mature fat tissue, mature cartilage, and mature respiratory epithelium (Figure 4b). Histopathologic examination confirmed the diagnosis of benign mature teratoma (Figure 5a and b). Post-operative the patient has complications such as persistent atelectasis on chest X-ray (Figure 6a), with prolonging intubation, empyema, and surgical site infection as wound dehiscence (Figure 6b).

Re-debridement of wound dehiscence and open drainage of empyema was done. The wound was well treated and there is no dehiscence, but unfortunately, the patient was passed away cause of sepsis on day-9 of surgery.
Discussion

Mediastinal mature teratoma is uncommon. However, teratomas are the most common form of germ cell tumor (GCTs) and have different epidemiology, histology, and biology than the others. These tumors occur equally in men and women, are generally benign [1]. GCTs both seminomas and non-seminoma occur in the mediastinum and compromise approximately 20 percent of anterior mediastinal masses. All forms of these tumors have a peak incidence in young adults. In our case, the patient was a young girl 15 years old; similar to the literature non-seminomas include embryonal carcinoma, teratocarcinoma, choriocarcinoma, yolk sac tumors, and benign teratoma. Teratoma of the mediastinum usually grows slowly [3] that is why patients do not feel any significant complaints for a long period of time, the tumor to become very large, filling the entire thoracic cavity with adhesions to the surrounding organs, this makes it difficult to remove the tumor and risks the cyst rupture. Most of the symptoms are related to compression of nearby structures such as chest pain, dyspnea, cough, or pulmonary infection [4]. In our case, there was prolong symptom of dyspnea as long as 4 years, but patients family refused for surgical resection, therefore, delayed clinical presentation can influence to the post-operative complications such as prolong atelectasis, rupture of tumor mass which spoiled of tumor contents, empyema, and sepsis.

Laboratory findings for tumor marker β HCG and AFP were the normal limit, same as other reports. Patients with benign mature mediastinal teratoma, serum β HCG, lactate dehydrogenase (LDH), and AFP levels were within the normal range [5]. Benign teratoma (pure mature teratoma) does not secrete AFP and β HCG. Serum levels of AFP, β HCG, and LDH are well-validated in their utility as noninvasive diagnostic of GCTs. A significant elevation of serum AFP or β HCG indicates significant components of yolk sac tumor or choriocarcinoma, respectively, and rules out pure mature teratoma or seminoma [6]. During the diagnosis and follow-up of mediastinal teratoma, it is necessary to monitor serum tumor markers AFP and β-HCG. Elevated serum AFP or β-HCG level indicates a malignant component to the teratoma such as embryonal carcinoma, endodermal sinus tumor, or choriocarcinoma [7].

Patients with benign mediastinal teratoma have a good prognosis after surgical resection [7]. Complete surgical excision is the treatment of choice because of probable complications such as compression of adjacent structures, rupture, or malignant transformation [8]. Complete excision of mediastinal teratoma usually performed through median sternotomy [9] or thoracotomy [10] or thoracoscopy [11], however, excision may be difficult because of tumor large size and adherent to adjacent and vital intrathoracic structure. The tumors of relative size are not too large, well-defined lobulated round mass, and there was no definite invasion to nearby structures, complete surgical resection giving good results without complications [6] and recurrence [12]. Otherwise, the tumor is very large, filling almost entire of the thorax cavity which has been adhered to the surrounding organs, there is a risk of damage to surrounding organs [12]. In our case, complete surgical excision was successful, but because of huge tumor size and has been adhered to the surrounding organs, there was complication persistent atelectasis, and rupture on some area of cystic mass during difficult dissection with spoiled tumor contents material to the operation field, put a risk on surgical site infection, empyema, sepsis, and death.

Conclusion

Mediastinal mature teratoma is uncommon. Delayed clinical presentation will give technical difficulties for resection and increase the risk of post-operative complication such as prolong atelectasis, tumor rupture, surgical site infection, empyema, sepsis, and death.

Authors’ Contributions

KPY [1] conceived the case report, contributed to collection of clinical details and writing, reviewing and finalization of the manuscript; AACTP [2] prepared the first draft besides contribution to collection of clinical details and illustrations; SMD [3] contributed to reviewing pathological specimen and illustration. All authors reviewed and finally approved the final manuscript.

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