Case report

Giant teratoma of anterior mediastinum in a 14-year-old girl as an example of potential diagnostic problems and errors

Przemysław Wolak1,2, Renata Skiba1, Wojciech Niedziela2

1Department of Correct and Functional Anatomy, Institute of Nursing and Obstetrics, Faculty of Health Sciences, Jan Kochanowski University, Kielce, Poland
Head of the Department: Prof. JKU Piotr L. Chłosta
2Department of Pediatric Surgery, Urology and Traumatology, Voivodeship Specialist Pediatric Hospital, Kielce, Poland
Head of the Department: Dr. Piotr Stepień
3Operating Theater of the Voivodeship Specialist Pediatric Hospital, Kielce, Poland
Head of the Theater: Dr. Wojciech Wąsacz

Key words: mediastinal teratoma, mediastinal tumor, fever of unknown etiology, exudative pleuritis.

Abstract

Teratomas are tumors originating from the three primary germ layers. The term “teratoma” originates from the ancient Greek word for monster. It was first used by Virchow in 1869 to describe a tumor in the sacrocaudal region [1]. Teratomas are tumors derived from the ectoderm, endoderm and mesoderm, although they may also be formed of a single type of germ layer [2]. The most common locations include the sacrocaudal region (35–60%) and gonads (ovaries and testicles, mediastinum (15%), rarely stomach (1%), retroperitoneal space (5%) or intracranially [1, 2]. Chest locations are rare. In particular, mediastinal teratoma showing no tumor-specific symptoms may be treated as exudative pneumonia. The goal of the article was to present a case encountered in our practice as a showcase of possible diagnostic problems and errors. Despite a thorough medical examination with additional exams (ultrasound scans of pleural cavities, chest X-ray and laboratory analyses), the diagnosis of a thoracic tumor was made only after chest computed tomography scan was performed following ineffective attempts at antibiotic therapy and pleural drainage. Following the diagnosis of mediastinal tumor, the patient was subjected to surgery. A giant teratoma (confirmed in histopathological examination) was removed upon left-sided thoracotomy. Following the procedure, lung expansion and patient recovery were observed. Computed tomography of the chest should be performed routinely upon encountering difficulties in the treatment of exudative pneumonia in children. In every case of pneumonia with pleural effusion in children, inflammatory mask of mediastinal tumors should be ruled out.

Introduction

Teratomas are tumors originating from the three primary germ layers. The term “teratoma” originates from the ancient Greek word for monster. It was first used by Virchow in 1869 to describe a tumor in the sacrocaudal region [1]. Teratomas are tumors derived from the ectoderm, endoderm and mesoderm, although they may also be formed of a single type of germ layer [2]. The most common locations include the sacrocaudal region (35–60%) and gonads (ovaries and testicles, mediastinum (15%), rarely stomach (1%), retroperitoneal space (5%) or intracranially [1, 2]. Chest locations are rare. In particular, mediastinal teratoma showing no tumor-specific symptoms may be treated as exudative pneumonia. Teratomas can be classified as mature or immature (with the structures of yolk sac tumor or ca embryonale, respectively) [1, 3]. In the case of immature teratomas, increase in the levels of tumor markers (α-fetal protein (AFP), human chorionic gonadotropin (β-HCG)) may sometimes be observed [1, 4, 5]. Determination of the levels of these markers facilitates the diagnosis and assessment of the dynamics of the pathological process. In patients under follow-up observation after completion of treatment, it may be the first indicator of cancer relapse. In most cases, teratomas are diagnosed immediately after birth or in infancy [3, 6–8].

The goal of the article was to present a case encountered in our practice as a showcase of possible diagnostic problems and errors. Despite a thorough medical examination with additional exams (ultrasound scans of pleural cavities, chest X-ray and laboratory analyses), the diagnosis of a thoracic tumor was made only after chest computed tomography (CT) scan was performed following ineffective attempts at antibiotic therapy and pleural drainage.

Case report

A 14-year-old girl was unsuccessfully treated for 7 days in an outpatient setting with an antibiotic (amoxicillin) for dyspnea, dry cough, fever reaching 39°C, and pain in the chest and left subcostal region. Physical examination upon admission revealed dull percussion above the left lung and lack of audible respiratory murmur in the left chest. A classical X-ray of chest was taken in the P-A plane, revealing shading of the lower 2/3 of the chest with relocation of the heart and mediastinum towards the right (Figure 1).
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The ultrasound scan of the pleura performed upon admission revealed a large amount of fluid (more than 1000 ml) in the left pleural cavity; fluid compartments were visible with part of the fluid being encysted; and the heart was relocated towards the right. Supportive examinations revealed no pathologies other than a positive C-reactive protein (CRP) result and elevated lactate dehydrogenase (LDH) level. Diagnosis of left-sided pneumonia was made and antibiotic treatment (cefuroxime) was initiated. The child was qualified for decompression of the left pleural cavity. The left pleural cavity was punctured to afford 750 ml of xanthochromic fluid collected for general examination and cultures. The patient’s health improved slightly as she experienced no fever and reported no dyspnea. The culture of the fluid collected from the chest revealed no growth of aerobic bacteria; general examination revealed no tumor cells. No tests were performed to screen for tuberculosis. In the follow-up ultrasound scan, the quantity of fluid within the left pleural cavity was measurable and amounted to 875 ml. The patient was transferred to the Department of Pediatric Surgery, Urology and Traumatology. Suction drainage was installed in the left pleural cavity, yielding 750 ml of xanthochromic fluid. Follow-up ultrasound scan and X-ray examination revealed no improvement – a suggestion was made to expand the diagnostic methods as the picture might correspond to cavernous necrosis in the course of pneumonia, infected cystic lung disease or hypertrophic lesion. Amikacin and fluconazole were added to the cefuroxime regime. A CT scan of the chest was performed on the 4th day of hospitalization, revealing a cystiform, multicompartmental mass with calcifications and adipose tissue, causing relocation of mediastinal structures towards the right. The size of the lesion was ca. 170 mm × 150 mm × 105 mm (Figure 2). The left pleural cavity contained a fluid layer of 28-mm thickness; the drain was visualized. The image most likely corresponded to a hypertrophic lesion. The serum levels of tumor markers (AFP and β-HCG) were in the normal range. Bronchoscopic examination was performed (the left upper bronchium was compressed with the lumen reduced to ca. 30% compared to the contralateral structure). The patient was qualified for surgical treatment. Surgery was performed on the 10th day after admission. Left-sided anterolateral thoracotomy was performed in the 5th intercostal space, revealing a very large solid/cystic tumor, as well as a sunken and relocated left lung. The adhesions were removed, tumor volume was reduced by evacuation of fluid from the cystic space, the tumor was dissected from the diaphragm, pericardial sac and left lung; the lung was dissected using a linear stapler. The left lung was decompressed, filling the pleural cavity. The pleural cavity was evacuated using 2 drainage tubes extending through the 6th intercostal space. The chest was closed in layers. The patient was transferred to the Department of Anesthesiology and Intensive Care. The post-operative recovery proceeded well. On the 3rd day after the procedure, bronchoscopy was required with simultaneous aspiration of secretion from the left bronchium and its branches. Antibiotic therapy was continued. On the 5th day after the surgery, the drainage tube was removed from the pleural cavity and the patient was transferred to the Department of Pediatric Surgery. The patient required respiratory rehabilitation. Macroscopic histopathological examination revealed a tumor sized 17 cm × 15 cm × 10 cm, with tooth buds, adipose tissue and hair visible in the cross-sections. The microscopic image corresponds to mature teratoma (teratoma maturum partim cysticum) (Figure 3). Tissues from three

Figure 1. Preoperative X-ray of chest was taken in the P-A plane

Figure 2. CT scan of the chest with tumor
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germ layers were identified within the tumor structure. The patient was discharged from the Department of Pediatric Surgery on the 10th day after the surgical procedure, cardiovascularly and respiratorily stable, with the chest wound properly healed. The treatment was continued at the Department of Pneumology and Allergology, from which the patient was discharged home on the 15th day after the surgery. The patient was under periodic observation at the Outpatient Clinic of Pediatric Surgery and the Outpatient Clinic of Pneumology, showing no complications of the surgical treatment for thoracic teratoma and no subsequent respiratory infection (Figure 4).

Discussion

In case of mediastinal location, teratoma may exert symptoms of respiratory inefficiency in neonates or infants, directly threatening the child’s life [3, 6–8]. According to American authors, 42% of tumors are diagnosed prenatally [3]. Mediastinal teratomas are operated on mainly immediately after birth or during the first months of life [3, 6–9]. However, late diagnosis of thoracic teratomas may occur. A case similar to that of our patient was reported by Golash in a 20-year-old female patient [10]. The giant mass of the tumor (mature teratoma) located within the mediastinum, sized 27 cm × 20 cm × 11 cm and weighing 14 kg, led to symptoms of respiratory disorders and difficulties swallowing due to the compression of the neighboring structures [10]. Another case was reported in Australia and pertained to an 18-year-old female patient with an immature teratoma sized 23 cm × 17 cm × 9 cm, weighing 2005 g and located within the left pleural cavity [11]. Our case was associated with marked diagnostic difficulties. The patient was admitted for exudative pleuritis without any reservations from radiologists, pediatricians and surgeons with relation to the presence of pleural fluid that should have been drained. However, the lack of clinical improve-

ment, xanthochromia of the fluid that was expected to be purulent effusion and, most of all, lack of clear changes in ultrasound scans and X-ray examinations of the chest following evacuation of nearly 2 L of fluid from what we suspected to be the left pleural cavity, suggested otherwise. Expanding the diagnostic range by inclusion of the chest CT scan allowed us to make the correct diagnosis and qualify the patient for surgical treatment. Therefore, we propose the expansion of the diagnostic methods used in dubious cases by computed tomography or magnetic resonance scans, as also highlighted by other authors [2]. Classic X-rays may not visualize fragments of bones or teeth within the tumor outlines; in most cases, the picture is uncharacteristic [2]. Mature teratomas may grow over years, with patients adapting to the anatomical conditions changed by the growth of the tumor. The presence of the tumor may be disclosed only by accompanying infection or trauma with tumor bleeding. Surgical procedures usually allow for complete recovery, as was observed in our case as well.

Conclusions

Computed tomography of the chest should be performed routinely upon encountering difficulties in the diagnosis and treatment of exudative pneumonia in children. In every case of pneumonia with pleural effusion in children, inflammatory mask of chest tumors, including mediastinal tumors, should be ruled out.

References


Figure 3. The macroscopic image of tumor

Figure 4. Postoperative X-ray of chest

Address for correspondence:

Przemysław Wolak
Institute of Nursing and Obstetrics
Department of Health Sciences
Jan Kochanowski University
al. IX Wieków Kielc 19, 25-317 Kielce, Poland
Phone: +48 501 525 549
E-mail: przemyslaw.wolak@ujk.edu.pl

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