An operated case of locally advanced thymic atypical carcinoid in anterior mediastinum: a case report

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Abstract: Primary thymic atypical carcinoid (TAC) is a kind of neuroendocrine tumors of the mediastinum, which has a poor prognosis due to its invasive behavior, metastasis and postoperative recurrence. We present a relatively rare case who came to hospital because of chest pain and tightness from a large mediastinal mass of 115 mm × 95 mm compressing left brachiocephalic veins, pericardium and upper-lobe of left lung. Although the operation was rather challenging, we performed complete resection including local lymph node dissection by open median sternotomy. The pathology of combining hematoxylin/eosin staining and immunohistochemical was confirmed to be primary TAC, grade 2 according to 2015 WHO Classification of Tumors of the Thymus. After radical surgery, the case underwent 6 cycles of platinum-based adjuvant chemotherapy. To date, the man has survived over 11 months without recurrence and distant metastasis. In conclusion, open surgery is a safe and effective method for locally advanced TAC and radical resection combination with adjuvant chemotherapy may lead to a long-term survival.

Keywords: Thymic atypical carcinoid (TAC); mediastinum; female; resection; case report

Case presentation

On January 15th in 2019, a 42-year-old male was referred to our hospital, complaining of chest pain and tightness for over 1 month, and the symptoms were getting worse gradually in the last week. He had no history of malignancy, and tumor family history, as well as genetic history, was not found. The patient reported no fever, cough, hemoptysis, difficulty swallowing, dyspnea, hoarseness and
limb weakness. No positive signs were detected in routine physical examination and tumor markers were normal, such as carcino-embryonic antigen (CEA), ferritin (FER). Chest CT revealed a large left-anterior mediastinum mass. Further evaluation demonstrated a 115 mm × 95 mm mass with patchy necrosis, heterogeneous enhancement and small blood vessels. Moreover, the large mass compressed left brachiocephalic veins, pericardium and upper-lobe of left lung (Figure 1). Based on clinical and CT appearances, the patient was suspected of invasive thymoma or thymic carcinoma pre-operatively. Due to the estimated great-difficulty by video-assisted thoracoscopic surgery (VATS), median sternotomy was performed to remove the tumor in January 30th, 2019. During the surgery, we found that the tumor was located in the left-anterior mediastinum with incomplete capsule, and that part of upper-lobe of left lung was compressed by the large mass. Complete tumor resection with combined regional mediastinal lymph node dissection were performed. Hematoxylin-eosin (HE) staining showed that the tumor cells were arranged in a nested, or ribbon pattern, with mild atypia, local hemorrhage, foci of necrosis and abundant interstitial blood vessels (Figure 2). Immunohistochemistry demonstrated positivity for CD56, Syn, CgA, CK (Figure 3A,B,C,D), and staining indices for Ki-67 was 15% (Figure 3E). No metastasis was found in all the 4 resected mediastinal lymph nodes. Postoperative pathology turned out to be primary TAC, Masaoka stage II (7). Postoperative chest CT (Figure 4A) showed that the tumor was resected completely and no recurrence occurred in postoperative follow-up (Figure 4B). From March 3rd on, a total of 6 cycles of postoperative adjuvant chemotherapy was given per 3 weeks successfully, and the chemotherapy regimen was gemcitabine (1,000 mg/m² on 1st day) plus cisplatin (80 mg/m² on 1st day). No 3/4 grade adverse effects (AEs) occurred during the perioperative period. There was no tumor recurrence or metastasis during the postoperative follow-up. A time lime showed the whole medical procedure of the special case (Figure 5).

**Discussion**

NETs have been reported in pancreas (8), gastrointestinal tract (9), lung (10), thymus, pituitary (11), thyroid (12),
parathyroid, breast, skin, adrenal gland, paraganglia and genito-urinary system (13). In 1972, thymic carcinoid was first described by Rosai and Higa et al. (14). So far, almost 220 TAC cases have been reported since then. Among all the NETs, TAC is relatively rare and often found by accident. It mainly occurs in men (15,16) and nearly one-third of patients with this malignancy are asymptomatic (17). Szolkowska et al. summarized that the percentage of thymic carcinoid in primary mediastinal neoplasms was nearly 1% (18). Thymic NETs originate from neuroendocrine cells, which are diffusely distributed in different tissues and organs. Primary TAC only represents a minority of NETs. However, prognosis of locally advanced primary TAC is not optimistic due to the high risk of recurrence and metastasis. And there are no accepted guidelines for us to deal with the rare disease. In this report, we presented a rare case

![Figure 3](image1.png)  ![Figure 4](image2.png)

**Figure 3** Immunohistochemistry demonstrated positivity for CD56, Syn, CgA, CK (A-D), and staining indices for Ki-67 was 15% (E) (×100).

**Figure 4** Neither residual tumor nor recurrence was observed upon a repeat CT scan in 1 month (A) and 9 months (B) after surgery, respectively.
of TAC located in left-anterior mediastinum in a middle-aged male. To our delight, the patient is still alive without local recurrence or metastasis after timely radical operation and post-operative adjuvant chemotherapy. However, no population-based data are available to build up guidelines for diagnosis, treatment, and follow-up.

Thymic carcinoids are more likely to have a large mass, irregular contour, heterogeneous intensity, heterogeneous enhancement and local invasion on CT or MRI (19,20). A necrotic or cystic component is often seen in atypical carcinoid. Small biopsy often cannot distinguish AC from TC, so surgical specimens are required to figure out precise classifications. Atypical carcinoid tumor is featured by a high degree of malignancy and invasiveness, and patients often complain of respiratory or local chest symptoms, such as cough, dyspnea, and chest pain. Thymic NETs are frequently associated with autoimmune disease or endocrine disorders, and the most common is Cushing syndrome, characterized by ectopic production of adrenocorticotropic hormone (ACTH) (21,22). The case in our study presented to hospital due to local chest symptoms, which may attribute to compression by the large mediastinal mass. Because of asymptomatic status at the early stage, a majority of patients were initially diagnosed at an advanced stage. Walts et al. (21) found that thymic NETs were usually large at high Masaoka stage at initial diagnosis and were associated with poor prognosis in spite of aggressive treatment.

It is generally accepted that strong and diffuse expression of more than one of four neuroendocrine markers (chromogranin A, synaptophysin, CD56 and NSE) in >50% of tumor cells has been maintained to diagnosed with NETs. Histopathologic characteristics of TAC include carcinoid morphology, 2–10 mitosis/2 mm², or foci of necrosis. In addition, TAC is well differentiated and it is suggested as intermediate grade neuroendocrine tumor (23). Moreover, thymic NETs genetics differ from each other (24), so precision genetics analysis may provide new avenues for diagnosis and treatment for TAC in future. Treatment strategies for TAC mainly include surgery, chemotherapy, and radiotherapy. Due to its aggressive behaviour and lack of effective chemotherapy or radiotherapy, surgical approach may be the best treatment to deal with TAC. Complete resection is deemed to be a favorable prognostic factor for patients with thymic NETs (15-17), but the rate of postoperative recurrence is high. A retrospective study in Japan (25) found that distant metastasis in bone and lung tissues was more common than local recurrence. Araki et al. (26) revealed that the rate of recurrence or metastasis in thymic NETs patients was 79%, including intra-thoracic metastasis (thoracic lymph nodes, pleura, lung, pericardium, postoperative mediastinum) and extra-thoracic metastasis (bone, abdominal lymph nodes, liver, pancreas, kidney, adrenal gland, spleen, brain). This article concluded that metastasis, as well as recurrence, was frequent, mainly involving thoracic lymph nodes, but extra-thoracic metastasis may also happen. Wu et al. (27) reported a case presented with neck pain and right upper limb numbness, and the case was diagnosed with cervical vertebra metastasis resulting from TAC. Studies have revealed that postoperative adjuvant chemotherapy or radiotherapy can prolong survival time. In our article, the case underwent complete resection and have survived to the present day without recurrence or metastasis. We chose open thymectomy rather than VATs because open surgery may remove the tumor to the maximum possible extent. Also, postoperative adjuvant chemotherapy was regularly

Figure 5 A timeline showing the medical procedure of the case.
performed to reduce the rate of recurrence or metastasis.

A population-based study by Jennifer suggested that the median overall survival time was 73 months and the 5-year survival rate was 56% in 254 thymic NETs cases, and patients who underwent surgery had a significantly longer survival time than those without surgery (28). Tumor size, proliferation index, histologic type, surgical resection, the Masaoka staging and postoperative radiotherapy have an impact on prognosis (16,17,29). Han et al. (30) found that the number of mitosis was an independent prognostic factor for recurrence and death in patients with atypical carcinoid tumor of the lung and thymus. In the above study, all seven TAC patients underwent total thymectomy with neck lymph node dissection, and six of seven received adjuvant chemotherapy, radiotherapy or chemoradiotherapy, but recurrence/metastasis appeared only in one patient, whose recurrence sites appeared on the mediastinum, pleura, brain, and bone. To the best of our knowledge, radical surgery is very necessary to perform for those locally advanced TAC. However, the management of TAC is full of challenge and requires a multi-discipline team for the final diagnosis and treatment.

We suggest that all NETs need to have a comprehensive assessment after clear diagnosis. In the case of our report, combination of radical surgery and adjuvant chemotherapy provide an excellent alternative for locally advanced TAC patients. Due to the tumor's large size, open thymectomy was performed instead of mini-invasive thoracoscopy, which is helpful to increase the R0 resection rate. Literature review has indicated that a male with a large anterior mediastinal mass should consider the possibility of thymic TAC. These patients might have concomitant diseases, such as Cushing syndrome or MEN type I syndrome. Radical resection is still the first choice for those early or locally advanced thymic TAC patients. The favorable prognosis can be achieved after radical surgery, and postoperative adjuvant therapies should be taken into consideration seriously.

However, there were several limitations in the whole course of treatment. First, neo-adjuvant chemotherapy or radiochemotherapy was not performed, which may induce tumor decreased and easier to remove. Second, the follow-up time is relatively short and a long-term follow-up is very necessary to evaluate the therapeutic efficacy.

In conclusion, for those who have a large mediastinal mass with compression syndrome, open surgery is a safe and effective method for locally advanced TAC and radical resection combination with adjuvant chemotherapy may lead to long-term survival.

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Footnote
Conflicts of Interest: 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. Written informed consent was obtained from the patient for publication of this manuscript and any accompanying images.

References