Transformation from Small Cell to Squamous Cell Carcinoma in a Thymic Carcinoma Patient with a Durable Response to Anlotinib: A Case Report

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Abstract: The most common pathologic type of thymic carcinoma (TC) is squamous cell carcinoma (SCC). Small cell carcinoma is relatively rare, accounting for approximately 2% to 5% of all thymic tumors. Histologic transformation of TC has not yet been reported. Available treatments for TC patients who progress after first-line therapy are limited, which contributes to their poor prognosis. We reported an extraordinary case of a 66-year-old man who was diagnosed with thymic small cell carcinoma that transformed into SCC after third-line treatment. Surprisingly, the patient had a progression-free survival (PFS) of 25 months and an overall survival (OS) of 10 years on anlotinib as fourth-line therapy. The tolerance was well. Thus, anlotinib may be a safe and promising treatment for TC patients, especially those who undergo histologic transformation.

Keywords: thymic carcinoma, squamous cell carcinoma, small cell carcinoma, histologic transformation, anlotinib

Introduction

Thymic carcinoma (TC) is a relatively rare but highly aggressive disease with a 5-year survival rate of only 36%.1,2 The histopathologic subtypes of TC vary in their degree of differentiation and aggressiveness, and squamous cell carcinoma (SCC) is by far the most common subtype. Small-cell neuroendocrine carcinoma is relatively rare, accounting for approximately 2% to 5% of all thymic tumors.3 The histologic transformation of TC has not yet been reported, and the selection of salvage therapies for them is elusive. About 46.5% of TC patients have advanced disease at diagnosis and are administered platinum-based chemotherapy as standard first-line treatment.4,5 However, there are few treatment options for patients who progress after first-line therapy, and the efficacy of further-line treatments is poor.

Anlotinib is an oral small molecule receptor tyrosine kinase inhibitor (TKI) that exhibits significant anti-angiogenesis and broad-spectrum antitumor activity via highly potent and selective suppression of vascular endothelial growth factor receptor-2 (VEGFR-2).6,7 It was recommended for use in the treatment of multiple cancers, including non-small-cell lung cancer (NSCLC), small cell lung cancer (SCLC) and soft tissue sarcoma.6 However, the effect of anlotinib in patients with TC, especially those who undergo histologic transformation, remains unclear. We reported an uncommon case of thymic small cell carcinoma that underwent histologic transformation to SCC and discussed the patient’s long-term response to anlotinib.

Case Presentation

A 66-year-old man was admitted to the hospital with sternal pain in December 2011. His chest computerized tomography (CT) scan showed anterior mediastinal space occupying lesions with mediastinal lymph node metastases. A biopsy of the
mediastinal lesions showed small cell carcinoma (Figure 1). The patient was diagnosed with thymic small cell carcinoma cT2N1M0 stage IVa. From December 2011 to May 2012, six cycles of first-line chemotherapy with irinotecan (65 mg/m^2 d1, 8) and cisplatin (60 mg/m^2 d1) were administered. During chemotherapy the patient also received 60 Gy/30f local radiotherapy for his mediastinal lesions. Partial response (PR) was achieved as assessed with Response Evaluation Criteria in Solid Tumors (RECIST) version 1.1. Radiation-induced pneumonia occurred, but improved after active treatment.

The patient felt sternal pain again in May 2017, and a positron emission tomography/computed tomography (PET-CT) scan showed progressive disease (PD) with right parasternal tissue enlargement and chest wall metastases (Figure 2). A puncture biopsy of the right parasternal tissue revealed metastatic thymic small cell carcinoma. Second-line chemotherapy with five cycles of irinotecan (65 mg/m^2 d1, 8) and cisplatin (60 mg/m^2 d1) was completed in September 2017. Efficacy was reported as stable disease (SD) after chemotherapy, and the patient’s pain improved.

The patient’s sternal pain worsened again in March 2018, and a CT scan showed an abnormal density in his right anterior chest wall and destruction of his ribs and sternum. Four cycles of third-line paclitaxel (175mg/m^2 d1) and cisplatin (60 mg/m^2 d1) chemotherapy were completed in June 2018. During the follow-up period, the patient’s response to chemotherapy was rated as SD (Figure 3A).

The patient reported a progressively enlarging right testicular mass and worsening sternal pain again in March 2019. An enhanced CT was performed which showed right clavicle and neck lymph node metastases, multiple new bone

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**Figure 1** The pathology of a mediastinal biopsy in 2011 showed thymic small cell carcinoma. (A and B) Hematoxylin and eosin (HE), (magnification 200×, 400×); (C–F) Immunohistochemistry: CD56+, CgA+, Syn+, Ki67 40%, (magnification 100×).
Figure 2. PET-CT scans in May 2017 showed soft tissue enlargement and chest wall metastasis. (A) Anterior mediastinal residual adjacent soft tissue density without increased FDG metabolism, representing local tumor biologic activity loss. (B) Bony destruction of the right 3rd and 4th anterior ribs and the adjacent sternum as well as the right 6th to 9th posterior ribs. (C) Corresponding parts of the right front chest wall soft tissue thickening.

Figure 3. Enhanced CT scans showed PD in March 2019 compared with prior imaging from June 2018. (A) Chest-enhanced CT scans showed multiple bone metastases with surrounding soft tissue masses in June 2018. (B–F) Enhanced CT scans in March 2019. (B) Right clavicle and neck lymph node metastases. (C) Soft tissue density shadow in the anterior mediastinum. (D) Enlarged soft tissue in the right chest wall. (E) Soft tissue image of the right spermatic cord. (F) Right testicular mass.
metastases with surrounding soft tissue masses, and right testicle metastases (Figure 3B–F). A biopsy was performed again on account of the patient’s extensive metastases. Interestingly, the pathology of the right parasternal mass biopsy showed SCC rather than small cell carcinoma (Figure 4). Due to the size of the right testicular mass, a right orchiectomy was performed, and postoperative pathology also confirmed SCC from the thymus gland (Figure 5).

The patient’s Eastern Cooperative Oncology Group performance status (ECOG PS) score was 3. He had a history of hypertension and heart disease. His pro-B-type natriuretic peptide (pro-BNP) reached 2055 pg/mL and his left ventricular ejection fraction was 53%. Expectant and supportive treatment was administered to improve the patient’s condition. His ECOG PS score improved to 2, his pro-BNP fell to 421.5 pg/mL and his left ventricular ejection fraction improved to 61%. Even so, the patient was not likely to tolerate further chemotherapy. We therefore chose to pursue palliative radiotherapy and anti-angiogenic therapy for optimal safety and efficacy. Palliative radiotherapy of 54 Gy/27f was administered to the parasternal mass due to sternal pain, with treatments ending May 2019. A CT after the completion radiotherapy showed that the size of the right chest wall metastasis was smaller, and the patient’s chest wall pain was significantly improved. The patient started to take anlotinib orally in June 2019 at a dose of 8 mg/day, once daily on days 1–14 of a 21-day cycle (2-weeks on, 1-week off) while taking medicine to lower his blood pressure and improve the condition of his heart. The patient developed grade II hand-foot skin reactions while taking anlotinib, which improved to grade I after supportive treatment. Over a routine observation period the patient’s response to anlotinib was evaluated as SD. In July 2021, the patient died of respiratory failure because he caught a cold that developed into a pulmonary infection. The patient therefore achieved a PFS of 25 months during anlotinib treatment, and an OS of up to 10 years from diagnosis (Figure 6). The patient’s treatment timeline is shown in Figure 7.

**Discussion**

Here we discuss a patient with a rare case of a thymic small cell carcinoma that transformed into SCC, who had a long-term response to anlotinib as fourth-line treatment.

![Figure 4](https://doi.org/10.2147/CMAR.S362858)

**Figure 4** Biopsy of the right parasternal mass showed thymic SCC. (A) Hematoxylin and eosin (HE), (magnification 100×); (B) HE, (magnification 200×); (C and D) Immunohistochemistry: CK5/6+, CKpan+; (magnification 100×).
The initial pathologic type of this TC was small cell carcinoma, as determined at the time of diagnosis via needle biopsy. Transformation into SCC after third-line therapy was confirmed via needle and excisional biopsy. It is possible that the patient’s primary cancer was mixed small cell and SCC because of intratumoral heterogeneity or sampling artifacts. Guo et al reported a case of a recurrent breast cancer whose pathological type changed from invasive ductal carcinoma to metaplastic SCC. The authors’ hypothesis was that the tumor’s transformation into breast SCC was likely to be the result of the malignant transformation of breast cancer stem/progenitor cells. Histologic transformation from adenocarcinoma to small cell carcinoma and SCC in epidermal growth factor receptor (EGFR)-mutant and anaplastic lymphoma kinase (ALK)-rearranged lung cancer patients has been identified as a potential mechanism of resistance to EGFR/ALK inhibition. Although the related mechanism by which small cell carcinoma transforms into SCC has not yet been reported, possible mechanisms for such a histologic transformation include pluripotent cancer stem cell differentiation, intratumor heterogeneity, and early clonal evolution. Further studies are therefore crucial to fully identify the mechanism of tumor pathologic transformation.

The patient had completed third-line treatment when his pathologic transformation occurred. A recent study reported that the median PFS of patients with metastatic or unresectable TC who were treated with first-line chemotherapy was 8.4 months. Significantly, in our case, the patient still achieved a PFS of 2 years after fourth-line anlotinib treatment despite

Figure 5 Postoperative pathology of the right orchiectomy showed thymic SCC. (A) Hematoxylin and eosin (HE), (magnification 200×); (B-F) Immunohistochemistry: CD117+; CK5/6+; CKpan +; SALL-4−; P40+, (magnification 100×).
the tumor’s pathologic transformation. Anlotinib not only effectively inhibited the migration and proliferation of capillary-like tubes adjusted by VEGFR in endothelial cells, but also inhibited tumor cell proliferation.\textsuperscript{19} In clinical trials, anlotinib was more effective than a placebo as salvage therapy for patients with advanced NSCLC, SCLC and soft tissue sarcoma.\textsuperscript{20,21} It has been reported that VEGFR-1 and VEGFR-2 are overexpressed in TC, which suggests that angiogenesis inhibitors might be a valid treatment strategy.\textsuperscript{22} However, no study of anlotinib in TC has been published. Clinical trials are necessary to confirm the efficacy of anlotinib in the treatment of TC.
Conclusion
In summary, this is the first report of a histologic transformation of TC from small cell carcinoma to SCC. The patient achieved exceptional PFS and OS with anlotinib as fourth-line treatment, which suggests that anlotinib may be a promising strategy for treating advanced TC with a transformed pathologic type. This potential benefit should be evaluated in future prospective studies.

Abbreviations
TC, Thymic carcinoma; SCC, squamous cell carcinoma; PFS, progression-free survival; OS, overall survival; TKI, tyrosine kinase inhibitor; NSCLC, non-small-cell lung cancer; SCLC, small cell lung cancer; VEGFR, vascular endothelial growth factor receptor; SD, stable disease; CT, computerized tomography; PR, partial response; PD, progress disease; ECOG PS, Eastern Cooperative Oncology Group performance status; pro-BNP, pro-B-type natriuretic peptide.

Ethics Statement
Written informed consent was obtained from the son of patient for publication of this case report and any accompanying images. The study is exempt from ethics committee approval since treatment decisions were based on clinical reasoning.

Data Sharing Statement
The data are available from the corresponding authors upon reasonable request.

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Disclosure
The authors have no conflicts of interest to declare in this work.

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