Successful Management of a ROS1-Rearranged Pulmonary Pleomorphic Carcinoma Using Serial Tyrosine Kinase Inhibitors

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Abstract: Pulmonary pleomorphic carcinoma (PPC) generally lacks actionable driver mutations such as epidermal growth factor receptor mutations or anaplastic lymphoma kinase or c-ros oncogene 1 (ROS1) rearrangements. The response to crizotinib, ceritinib, brigatinib, and lorlatinib in ROS1-positive advanced non-small cell lung carcinoma is well established; however, there is little mention of their successful administration in pulmonary pleomorphic carcinoma cases. We report a case of a stage II PPC with recurrence after surgical resection and developed multiple distant metastasis. The tumor was refractory to chemotherapy and immunotherapy with progressive disease. EZR-ROS1 fusion was detected by next-generation sequencing and showed a good response to serial ROS1 inhibitors combined with surgery and radiotherapy. Now under lorlatinib, all her lesions responded well during the follow-up with sustained partial remission for more than 18 months. A sustainable treatment effect can be achieved in pulmonary pleomorphic carcinoma with driver mutations with tyrosine kinase inhibitor treatment. Driver mutations should be regularly tested in pulmonary pleomorphic carcinomas.

Keywords: pulmonary pleomorphic carcinoma, lung cancer, ROS1-rearranged, tyrosine kinase inhibitor, lorlatinib

Introduction
Pulmonary pleomorphic carcinoma (PPC) is a rare malignant tumor of the lung, accounting for around 0.1–0.3% of all lung cancers.1 According to the revised 2015 World Health Organization classification, PPC is one of the five subtypes of pulmonary sarcomatoid carcinoma (PSC) and contains at least 10% spindle and/or giant cells. The clinical course and response to systemic therapy are generally poor in PPC, which has a lower incidence of druggable driver mutations than lung adenocarcinoma. Kirsten rat sarcoma virus oncogene homolog (KRAS), MET exon 14 skipping, and epidermal growth factor receptor (EGFR) mutations are the most common driver oncogenes in PSC, whereas anaplastic lymphoma kinase (ALK) and c-ros oncogene 1 (ROS1) fusions are seldom identified.2 In recent years, several newer generation tyrosine kinase inhibitors (TKIs) have emerged and shown good activity in treatment-naïve or heavily treated patients, including alecinib, ceritinib, brigatinib, and lorlatinib. However, in the real-world practice of lung adenocarcinoma harboring druggable driver mutations, not all patients respond well to each type of TKI, making the switch between TKIs a mandatory and reasonable approach to maximize the benefit of
targeted therapy. However, the data regarding TKIs other than crizotinib in PPC with fusion genes are lacking.

Herein, we describe a heavily treated ROS1-rearranged PPC patient successfully managed by serial ROS1 TKIs. Appropriate written informed consent was obtained for publication of this case report and accompanying images.

**Case Report**

Chest computed tomography revealed a 3.5 cm tumor in the right middle lobe in April 2017 in a 45-year-old non-smoking woman. She was diagnosed with stage II pleomorphic carcinoma (pT2aN1M0 according to the American Joint Committee on Cancer handbook 7th edition) and underwent right middle lobe lobectomy in May 2017, followed by four cycles of adjuvant chemotherapy (cisplatin and pemetrexed every 3 weeks, which ended in August 2017). However, tumor recurrence occurred 3 months later, with a new left lower lung tumor, bone (left clavicle and right femur) and brain (left cerebral frontal lobe) metastases (Figure 1). The tests for driver oncogenes, including EGFR, ALK, and ROS1 were negative, using the MassARRAY® system (Agena, CA, USA), VENTANA ALK (D5F3) CDx Assay, and ROS1 (D4D6) rabbit monoclonal antibody immunohistochemical (IHC) assay, respectively.

Because of the chemoresistant nature of PPC, the patient received an aggressive multidisciplinary approach including lung tumor resection (S6 segmentectomy) in February 2018, radiation to the bone metastases (35 Gy in five fractions to the left clavicle and right femur) in February 2018, and brain tumor resection in March 2018 followed by radiation to the brain tumor bed (18 Gy in a single fraction) (Figure 1). The pathologic findings of the lung (Figure 2A) and brain tumors showed pleomorphic carcinoma (Figure 2B) arising in an adenocarcinoma, which was positive for thyroid transcription factor 1 (Figure 2C) and vimentin (Figure 2D) and negative for CK7. PD-L1 (Dako PD-L1 IHC 22C3 pharmDx assay) was expressed in 80% and 45% of tumor cells in the lung and brain tumors, respectively. Unfortunately, new lung, and renal metastases developed during treatment; therefore, a combination of docetaxel (35 mg/m² on days 1 and 8 every 3 weeks) and pembrolizumab (100 mg every 3 weeks) was administered as salvage treatment for the new metastases. However, progressive disease with new bone metastasis (left rib, right ischium, right femur) was identified by positron emission tomography (PET) after two cycles of chemotherapy-immune checkpoint inhibitor combination therapy.

![Figure 1](image-url)  
**Figure 1** Summary of the treatment course in this case report. Arrows indicate the target lesions in the imaging studies.  
**Abbreviations:** PPC, pulmonary pleomorphic carcinoma; Cis, cisplatin; Pem, pemetrexed; Doc, docetaxel; PEM, pembrolizumab; TrAE, treatment-related adverse event; RT, radiotherapy; RML, right middle lung; LLL, left lower lung; RUL, right upper lung.
Owing to the refractory nature of her disease, we performed next-generation sequencing (NGS) using the previous surgical specimens (Foundation One assay, hybrid-capture-based NGS to detect 324 genes, Foundation Medicine, Inc.) to find possible driver mutations in tumor of young non-smoker female patient. Unexpectedly, an EZR-ROS1 fusion was identified. Using a more sensitive ROS-1 antibody (SP384), we disclosed both resected lung tumor and brain tumor had ROS-1 protein expression (Figure 2F). Therefore, crizotinib treatment was started in April 2018, and she achieved a partial response 2 months later (according to the Response Evaluation Criteria in Solid Tumors, version 1.1), with shrinkage of the lung and renal tumors. Despite the remarkable efficacy of TKIs, grade 4 treatment-related hepatoxicity was observed during crizotinib therapy. The peak levels of aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were 865 U/L and 2451 U/L, respectively; total bilirubin and alkaline phosphatase were within the normal limits. Her liver enzymes returned to normal levels at day 39 after crizotinib discontinuation (AST 19 U/L, ALT 20 U/L). We changed the treatment to ceritinib in July 2018, and the partial response to TKIs persisted until November 2018, although new lung tumors emerged in the right upper and lower lobes (Figure 1). Therefore, we changed the treatment to the third-generation ROS1 TKI lorlatinib (100 mg daily) in combination with radiation therapy to the lung tumors (5250 cGy in 15 fractions). The lung tumor, bone, and renal metastases responded well during follow-up (last image study in June 2020 as showed in Figure 1), except a pseudoprogression of the left frontal lobe brain tumor occurring in March 2019, which was confirmed as radiation necrosis after surgical resection (Figure 3). Recurrence of radiation necrosis was noted 3 months after surgery, and it later spontaneously resolved. The patient is still under lorlatinib therapy since November 2018 with a sustained systemic partial remission. The total treatment course is summarized in Figure 1.

**Discussion**

Herein, we describe a patient with heavily treated ROS1-rearranged PPC successfully managed by serial ROS1 TKIs. The patient developed severe liver toxicity after a partial response to crizotinib and was successfully switched to ceritinib and lorlatinib uneventfully. The disease was well-controlled for years. Our report demonstrates the efficacy of newer generation ROS1 TKIs even in PPC, a tumor type with a lower incidence of driver mutations.

PPC is known for its resistance to chemotherapy and low responsiveness to radiotherapy. Therefore, different
treatment strategies have been assessed in recent years. Surgery is the treatment of choice in early-stage PPC with a high recurrence rate; these patients may also benefit from perioperative chemotherapy. In advanced PPC, response to first-line chemotherapy is poor, with an overall survival of 5 months. Multimodality treatment with surgery, radiotherapy, and immunotherapy has been attempted but rarely achieves good response. A high frequency of PD-L1 expression has been observed in PPC patients, but variable response to immune checkpoint inhibitors is reported. Our patient had a poor response to pembrolizumab combined with docetaxel despite a high expression of PD-L1 in the primary and metastatic tumors.

Druggable driver mutation genes are detected by comprehensive genomic profiling in 30% of PSC patients, with MET exon 14 skipping being the most common mutation, adding an additional option for treatment. ROSI rearrangement is rare in PPC and the specificity of ROSI in IHC was low hence ROSI status should be confirmed with fluorescence in situ hybridization (FISH) or NGS. The reason for the discrepancy of results of ROSI IHC stains in our case might be caused by different sensitivity of antibodies. As previous studies reported, SP384 is more sensitive than D4D6 in FISH detected ROSI gene arrangement lung cancer. The discordance between SP384 and D4D6, though rare, was also shown in previous studies.

Crizotinib and ceritinib are ROSI inhibitors with overall response rates of 72% and 62% in NSCLC patients, respectively, and median progression-free survivals of 19.2 and 19.3 months, respectively. Lorlatinib also showed promising results in a Phase 1–2 trial. However, only one PSC case with ROSI alteration has been reported; the patient showed a sustained response to crizotinib for 46 months. Our case describes the first successful treatment with serial ROSI inhibitors. In our patient, crizotinib showed a good effect; however, hepatoxicity occurred, and the treatment was shifted to ceritinib. New lesions developed after 4 months. The treatment was changed to lorlatinib in combination with local radiation treatment. The resistance to ceritinib may be related to kinase domain mutation as opposed to activation of other pathways because the patient had a sustained response to lorlatinib.

Radiation necrosis presenting as pseudoprogression during alectinib treatment of previously radiated brain metastases has been reported by Ou et al. No other similar presentation of ALK inhibitors has been reported. Our patient developed pseudoprogression 11 months after Cyberknife surgery and about 3 months after starting lorlatinib, which is similar to what was observed in two previously reported cases. However, the causal relationship and pathophysiology remain unclear.

**Conclusion**

To our knowledge, this is the first report of ROSI-rearranged lung PPC successfully treated by serial TKIs. Our case highlights the importance of molecular testing even in PPC, which has a low prevalence of driver mutations, sequential use of TKIs, the delicate management of
treatment-related adverse events, multidisciplinary treatment, and the careful differential diagnosis of radiation necrosis rather than true progression, which may contribute to extended patient survival.

**Abbreviations**

*ROSI*, c-ros oncogene 1; *PPC*, pulmonary pleomorphic carcinoma; *PSC*, pulmonary sarcomatoid carcinoma; *KRAS*, Kirsten rat sarcoma virus oncogene homolog; *EGFR*, epidermal growth factor receptor; *ALK*, anaplastic lymphoma kinase; *TKI*, tyrosine kinase inhibitor; *PD-L1*, programmed death-ligand 1; *NGS*, next-generation sequencing; *AST*, aspartate aminotransferase; *ALT*, alanine aminotransferase.

**Consent for Publication**

The patient signed the written informed consent to use and publish the patient’s data, case details, and images for study. No institutional approval was required to publish the case details.

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Chang-Wei Wu, Ching-Yao Yang, and Yih-Leong Chang report no conflicts of interest for this work.

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**References**


