EML4-ALK positive lung adenocarcinoma with skeletal muscle metastasis in the right calf which was treatable with lorlatinib after resistance to treatment with alectinib

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SUMMARY
This report concerns a patient with skeletal muscle metastases due to lung adenocarcinoma harbouring an echinoderm microtubule-associated protein-like-4 (EML4)-anaplastic lymphoma kinase (ALK) rearrangement, who was successfully treated with lorlatinib after resistance to alectinib. A right lower lobectomy based on a diagnosis of lung adenocarcinoma was performed on a 77-year-old Japanese woman. After 7 months of surgical resection, a mass in the right calf was observed. A fine-needle aspiration biopsy from the mass was performed and the mass was diagnosed as metastatic adenocarcinoma harbouring EML4-ALK rearrangement. Alectinib was administered for 10 months. Then, administration of lorlatinib, an ALK tyrosine kinase inhibitor classified as third generation, was initiated after resistance to treatment with alectinib. After starting treatment with lorlatinib, the gastrocnemius tumour diminished and has maintained a stable condition. Our case suggests that EML4-ALK positive lung adenocarcinoma is treatable with lorlatinib after resistance to treatment with alectinib.

BACKGROUND
Lung cancer is the major cause of cancer-related death, and it is generally known that lung cancer commonly spreads to other parts of the body, such as the brain, the bone and the liver and rarely metastasises to skeletal muscle.1 2 Skeletal muscle metastases (SMM) were mostly found in the trunk or upper limbs and rarely in the lower limbs.3 It is also said that the presence of SMM represents a higher risk of mortality.4 However, the mechanism of SMM is not clear.

Nowadays tyrosine kinase inhibitors have changed treatment strategies for advanced lung cancer. Echinoderm microtubule-associated protein-like-4 (EML4)-anaplastic lymphoma kinase (ALK) gene rearrangement is one of the driver mutations first reported in 2007.5 ALK inhibitors have demonstrated good activity in patients with EML4-ALK positive non-small cell lung cancer (NSCLC).6 7 Lorlatinib is an ALK and c-ros oncogene 1 tyrosine kinase inhibitor classified as third generation and is expected to overcome ALK resistance mutations that can develop during treatment with ALK inhibitors classified as first or second generation.8

We report on a patient diagnosed as having EML4-ALK positive lung adenocarcinoma with skeletal muscle metastases in the lower leg who responded well to lorlatinib after showing resistance to treatment with alectinib.

CASE PRESENTATION
A 77-year-old woman was referred to our hospital because of an abnormal shadow in the right lower lung. Her medical history included postoperative thyroid papillary carcinoma when she was 71. She...
and had no history of smoking. Chest X-ray revealed a solid tumour in the right middle lung field (figure 1A). Thoracic CT revealed a solid tumour in the lower lobe of the right lung (figure 1B). A right lower lung lobectomy was performed based on the diagnosis of lung adenocarcinoma in the right lower lobe without any distant metastasis (pT1cN1M0, stage IIB seventh edition). After the surgical resection, the patient was observed carefully without adjuvant chemotherapy. About 7 months after the operation, a relapse of the lung cancer, a positron emission tomography (PET)-CT was performed revealing an abnormal uptake in the lung field (figure 1B). Both H&E stains from the gastrocnemius tumour (A) and the lung (C) revealed tumour cells with acinar pattern.

**INVESTIGATIONS**

The patient’s routine complete blood count and chemical laboratory data were all within the normal range. Her carcinoembryonic antigen was normal (3 ng/mL), and her sialyl lewis X-i antigen was slightly elevated (48.8 U/mL). With concern regarding a relapse of the lung cancer, a positron emission tomography (PET)-CT was performed revealing an abnormal uptake in the right lung hilar lymph node, the right popliteal lymph node and the right gastrocnemius (figure 2A,B). Contrast-enhanced MRI of the right leg showed a tumour in the gastrocnemius on the T2 emphasised image, which was of heterogeneously high-signal intensity, with the rim of the tumour enhanced by the gadolinium (figure 2C,D). A fine-needle aspiration biopsy (FNAB) from the gastrocnemius tumour was performed. The specimen obtained revealed moderately differentiated adenocarcinoma (figure 3A). Immunohistochemistry (IHC) studies revealed that diffused expression of cytokeratin 7 (CK7) (figure 3B), thyroid transcription factor-1 (TTF-1) was negative, whereas IHC of the primary lung tumour revealed expression of both of them. The primary lesion in the right leg tumour could not be clarified from the IHC stains. However, in the morphological feature of the primary lung cancer (figure 3C) and the leg tumour, both the H&E stains showed an acinar pattern. Thus, the patient was diagnosed with relapsed lung adenocarcinoma. In addition, using IHC, it was proven that the primary lung cancer was harbouring EML4-ALK rearrangement.

**TREATMENT**

Alectinib (300 mg two times a day) was administered as the initial treatment, and the gastrocnemius tumour shrank in a month. After 10 months of treatment with alectinib, the gastrocnemius tumour increased in size with pain again. And in spite of two cycles of second-line therapy using pemetrexed, the disease could not be controlled. Lorlatinib (100 mg one time a day) was initiated as third-line therapy.

**OUTCOME AND FOLLOW-UP**

After initiating lorlatinib, the gastrocnemius tumour diminished within a month and has maintained a stable condition (figure 4). In 2 months of taking lorlatinib, dyslipidaemia (low density lipoprotein cholesterol 265 ng/mL, triglyceridaemia 474 mg/dL) developed as adverse events, treatable with rosuvastatin and bezafibrate. No other severe side effects have been revealed so far.

**DISCUSSION**

It is generally known that lung cancer commonly metastasises to other parts of the body. About 20%–50% of patients with NSCLC have metastases when lung cancer is diagnosed.9 The common sites of distant metastases are the brain (10%), bone (7%) and the liver (5%).2 Besides these organs, the skeletal muscle is one of the sites where NSCLC metastasises. SMM is a rare condition, estimated at 2.6% of NSCLC at the time of initial diagnoses.1 The presence of SMM is considered as a progressive condition. Pop et al reported that the median survival time of patients with lung cancer with SMM was 6 months.4 The reason for the rarity of SMM is still unclear, but there are several hypotheses, one of which suggests that muscle resists metastatic disease, such as mechanical, metabolic or immunological. None of these can explain the whole mechanism but in combination theorising may be possible.4

Most of the instances of SMM were accidentally diagnosed during routine CT, because a large proportion of patients with SMM are asymptomatic or neglected.1 There are several CT imaging patterns. It was reported that clinical symptoms such as severe pain or restriction of local movement tend to arise when SMM revealed a rim-enhanced tumour using enhanced CT.10 MRI is also a helpful technique for characterising soft tissue regions. Li et al reported that soft tissue sarcoma, single haematomata, haemangioma or malignant fibrous histiocytoma can mimic SMM in MRI.11 Nevertheless, CT and MRI are sometimes not sufficient for screening all soft tissue tumours. PET-CT could provide additional diagnostic information to a greater extent than other staging examinations.12 Although the metastatic tumour was in the lower leg in this case, the trunk and upper limbs are common metastatic sites according to a systematic review relating muscle metastases.3

**Figure 3** Histopathology slides obtained from the gastrocnemius tumour and the lung. H&E stain exam (×20) from the gastrocnemius tumour (A) and the lung (C). Immunohistochemistry studies (×20) from the gastrocnemius tumour revealing diffused expression of cytokeratin 7 (B). Both H&E stains from the gastrocnemius tumour (A) and the lung (C) revealing tumour cells with acinar pattern.

**Figure 4** Enhanced CT showing the skeletal muscle metastases in the right leg before (A) and after (B) 2 months of taking lorlatinib. The tumour exhibited a response with lorlatinib.
Biopsy is the most reliable test to confirm SMM. Trocar biopsy and FNAB are valid in order to distinguish carcinoma from other diseases. In the present literature, the most common histological diagnosis of SMM confirmed by FNAB was adenocarcinoma. Panel markers such as CK7 and CK20, and TTF-1 are necessary to detect the origin of a metastatic tumour. However, TTF-1 positivity is observed in 75%–80% of lung adenocarcinomas with sensitivity decreasing with poor differentiation as well as in mucinous adenocarcinoma.

There are some treatment options for SMM such as radiotherapy, surgical excision and medication. SMM is sometimes painful and excision of the painful mass may help the patients. Local treatments of the mass might affect the rate of recurrence and survival time. Tuoheti et al reported in a case series that a patient with lung cancer who underwent the excision of the muscle metastatic lesion was disease-free for 92 months and patients with some other cancer survived for an average of 18.5 months after radiation therapy.

EML4-ALK gene rearrangement occurs in 3%–5% of NSCLC in East Asian countries. Mostly, ALK inhibitors are the preferred choice for advanced EML4-ALK positive NSCLC. Crizotinib and ceritinib have improved progression-free survival (PFS) compared with chemotherapy. Then alectinib showed longer PFS than crizotinib (34.8 months with alectinib vs 10.9 months with crizotinib). Considering these facts, ALK inhibitors can be the appropriate choice for patients with EML4-ALK positive lung cancer with SMM.

Resistance mechanisms to ALK inhibitors depend on secondary mutation in the kinase domain or the variant type of ALK fusion. These mechanisms could be related to resistance to treatment with alectinib in our case. Furthermore, the shorter duration of response for alectinib might be connected to the aggressive state of the disease, such as the presence of SMM.

Lorlatinib is an ALK tyrosine kinase inhibitor classified as third generation with a multi-coverage of ALK kinase domain inhibitor. In the global phase 2 study, lorlatinib proved efficacious in patients with EML4-ALK positive lung cancer who had been treated with former generation ALK inhibitors such as alectinib or crizotinib. It was reported that 47% of patients previously treated with at least one ALK inhibitor improved clinically and their median PFS was 6.9 months. The median time for first tumour response was 1.4 months. Such a rapid response time and high response rate might be helpful for a patient with painful SMM.

Finally, SMM from lung adenocarcinoma is a rare but aggressive condition. Lorlatinib is an effective choice for advanced EML4-ALK positive lung cancer after the failure of alectinib treatment with good tolerance. Furthermore, appropriate assessments of pathological and genetic diagnosis are important steps to provide better treatments.

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