Alectinib as a treatment option following recovery from crizotinib-induced interstitial lung disease in patients with anaplastic lymphoma kinase-positive advanced non-small-cell lung cancer

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Received November 3, 2015; Accepted March 21, 2016

DOI: 10.3892/mco.2016.838

Abstract. Crizotinib is a tyrosine kinase inhibitor that displays antitumor activity against anaplastic lymphoma kinase (ALK)-positive advanced non-small-cell lung cancer. However, crizotinib-associated interstitial lung disease (ILD) has been reported as an infrequent, but potentially fatal complication. We herein describe the case of a 63-year-old male patient with ALK-rearranged advanced lung adenocarcinoma. Chest computed tomography (CT) revealed extensive bilateral ground-glass opacity and airspace consolidation with traction bronchiectasis on day 27 of crizotinib therapy. No signs of infection or left heart failure were identified and we considered the lesions to be consistent with crizotinib-induced ILD. Following corticosteroid treatment and discontinuation of crizotinib, CT revealed improvement of ILD, but also showed regrowth of the tumor. Alectinib, a novel alternative ALK inhibitor, was initiated, and has been successfully continued, with neither disease progression nor recurrence of ILD. The present case indicates that alectinib may be considered as an alternative agent in cases of crizotinib-induced ILD, irrespective of the pattern of ILD, i.e., a diffuse alveolar damage (DAD) or non-DAD pattern, with careful observation.

Introduction

Crizotinib is a tyrosine kinase inhibitor that has antitumor activity against anaplastic lymphoma kinase (ALK)-positive advanced non-small-cell lung cancer (NSCLC) (1). However, crizotinib-associated interstitial lung disease (ILD) has been reported as an infrequent but potentially fatal complication (2,3).

Alectinib is a novel oral ALK inhibitor with high potency and selectivity for ALK, which displays promising antitumor activity in NSCLC. Alectinib has shown promising activity in patients with crizotinib-resistant disease and has been generally well-tolerated in clinical trials (4,5). The safety of alectinib for patients who develop crizotinib-induced ILD has not been determined. We herein describe a case of ALK-positive NSCLC successfully treated with alectinib after developing crizotinib-induced ILD.

Case report

A 63-year-old Japanese man with a 25 pack-year smoking history presented with a 4-month history of cough. The patient was diagnosed with advanced lung adenocarcinoma (cT4N3M1b), with bone, brain and multiple lymph node metastases (Fig. 1A). Fluorescence in situ hybridization analysis revealed the presence of ALK gene rearrangement. Crizotinib (250 mg twice daily) was administered as first-line chemotherapy. On day 27, the patient developed high-grade fever and exertional dyspnea. Chest computed tomography (CT) revealed a decrease in tumor size; however, there was new extensive bilateral ground-glass opacity (GGO) and airspace consolidation with traction bronchiectasis (Fig. 1B). With the patient on oxygen at a flow rate of 5 l/min via a mask, the arterial blood gas analysis revealed a PaO2 of 48.5 mmHg. Although the chest CT revealed bilateral pleural effusions, there was no evidence of left heart failure. There were also no signs of infection. Based on these findings, crizotinib-induced ILD was suspected. Crizotinib was discontinued immediately, and methylprednisolone pulse therapy (1 g/day for 3 days) followed by oral prednisolone (PSL) at a dose of 65 mg/day was initiated. Given that the patient’s symptoms and radiological findings improved, the PSL dose was gradually tapered. Ten weeks after crizotinib was discontinued, the PSL dose was decreased to 25 mg/day. The chest CT showed no recurrence of crizotinib-associated interstitial lung disease.
of ILD, but the primary lesion had regrown (Fig. 1C). After discussing the risk of ILD exacerbation and obtaining the patient’s informed consent, alectinib (300 mg twice daily) was initiated. After 2 weeks, the primary lesion had decreased in size, without any exacerbation of the ILD. The patient received continuous alectinib treatment, with neither disease progression nor recurrence of ILD for more than 4 months, without the need for steroid therapy (Fig. 1D).

Discussion

Treatment with the first-generation ALK inhibitor crizotinib is associated with a risk of ILD development. ILD, which may be severe and occasionally fatal (2,3), develops in 2% of the cases. Crizotinib-induced ILD may present with at least two types of radiological findings, namely as the ‘diffuse alveolar damage (DAD) pattern’ and the ‘hypersensitivity pneumonia (HP) pattern’ (6). The DAD pattern is severe and fatal in most cases. Chest CT shows a rapid, bilateral and widespread development of GGO. The HP pattern is less severe and may not be associated with a definitive need for crizotinib withdrawal. Chest CT shows a predominant GGO pattern, which is localized and faint. Some cases of successful crizotinib retreatment after crizotinib-induced ILD have been reported, but they all had the non-DAD pattern (6,7). Retreatment with crizotinib should be avoided in patients with the DAD pattern, as the mortality rate of such patients is very high.

While crizotinib is a multitargeted inhibitor of the ALK, MET, and ROS1 receptor tyrosine kinases (8,9), alectinib is highly selective for ALK. The high selectivity for ALK may contribute to its reduced toxicity compared with crizotinib (4). We were able to successfully treat our patient, who had developed a crizotinib-induced DAD pattern of ILD, with alectinib. Therefore, alectinib may be considered as an alternative agent in cases of crizotinib-induced ILD, irrespective of the pattern of ILD, i.e., a DAD or non-DAD pattern, with careful observation.
References


