

CASE REPORT

Great Efficacy of Afatinib in a Patient with Lung Adenocarcinoma Harboring EGFR L833V/H835L Mutations: A Case Report

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Keywords: non-small-cell lung cancer, epidermal growth factor receptor, afatinib, EGFR compound mutation, L833V/H835L

Introduction

Epidermal growth factor receptor (EGFR) is the most common driver gene in non-small cell lung cancer (NSCLC) patients. The mutations in exons 18 to 21, especially exon 19 deletion and exon 21 L858R account for a large proportion of all EGFR mutations. In recent years, with the application of the next generation sequencing (NGS), more and more uncommon EGFR mutations have been detected, which may potentially benefit from EGFR TKIs. However, more evidence should be provided for the clinical significance of rare EGFR mutations, especially complex mutations. Here, we present a patient with NSCLC harboring EGFR exon 21 L833V/H835L complex mutation, who had clinical response to first-line afatinib therapy.

Case Report

A 65-year-old never-smoking male patient presented to our hospital in September, 2019 due to 1-month history of cough and a few days of thoracalgia. According to the subsequently high resolution computed tomography (HRCT) scanning and pathological examination, the patient was diagnosed with stage IV lung adenocarcinoma. For genomic analysis, capture-based ultra-deep targeted sequencing (depth:

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10,000×, LOD: 0.2%) was performed on circulating tumor DNA (ctDNA) extracted from pleural effusion and plasma by a large panel consisting of all exons and critical introns of 520 cancer-related genes (Oncosreen PlusTM, Burning Rock Biotech. Guangzhou, China). The results showed that the patient only had a complex EGFR mutation: L833V/H835L in exon 21 (VAF: 6.72%. Figure 1). The tumor mutation burden (TMB) was 2.4 mutations/Mb and the expression of PD-L1 was negative. After literature research and multiple disciplinary team (MDT) discussion, daily oral of afatinib (40mg, bid.) was administered starting at the middle of September 2019. No grade III or IV

adverse event (AE) was observed. Surprisingly, after one month of treatment, HRCT scan revealed a significant shrinkage in the nodules on the middle right lobe of the lung and decrease in number of nodules throughout the lung as compared to the baseline, indicating a partial response to afatinib (Figure 2A and B). In addition, no mutations were detected in plasma DNA using the previous NGS panel, also suggesting a good efficacy of afatinib. In December 2019, no apparent abnormalities were found by color ultrasound of neck and abdomen, and the level of circulating carcinoembryonic antigen (CEA) was significantly decreased compared with the

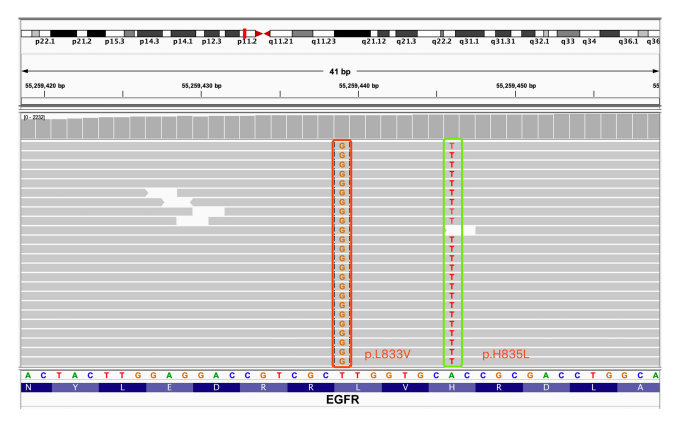


Figure 1 Identification of L833V and H835L in EGFR exon 21 from liquid biopsy.

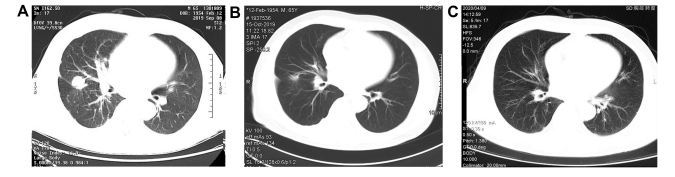


Figure 2 Medical imaging. CT scan with a right middle lobe lung mass before (A) and after (B) I month and 7 month (C) of afatinib treatment.

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first two times. After 7 months of afatinib treatment, the nodule in the middle lobe of right lung was almost disappeared (Figure 2A and C). No metastasis was found and the mutation in ctDNA remained negative. The patient is still benefit from afatinib treatment at the last follow-up time (July 20, 2020). The PFS of afatinib was over 10 months.

Discussion

In the present case, we showed the clinical efficacy of afatinib in a patient with EGFR L833V/H835L mutations. The patient is still under treatment with afatinib, and the PFS is over 10 months. To the best of our knowledge, this is the first case of first-line treatment with afatinib in Asian NSCLC patient harboring EGFR L388V/H835L mutation.

Complex EGFR mutation is a small group of uncommon mutations, which comprised of two or more EGFR point mutations. Among them, the most frequent type is an EGFR sensitive mutation in combination with a nonclassical mutation, which is called compound mutation.^{5–} ⁷ Another frequent type is EGFR sensitive mutation plus resistant mutation, such as L858R/T790M, 19del/T790M and T790M/C797S.8 Two or more synchronous rare EGFR mutations like L833V/H835L mutations are extremely rare in lung cancer. A previous study which compared the testing methods and tissue types of EGFR detection have found 3 L833V/H835L mutations in 157 NSCLC patients. On the other hand, only a few studies have detailed the clinical significance of this mutation. An Asian patient harboring L833V/H835L mutations were reported to benefit from gefitinib treatment lasting 34 weeks. 10 In addition, a recent cohort study about in cis EGFR mutations in NSCLC has reported a case carrying L833V/H835L mutations. The patient was treated with gefitinib and the PFS was over 15 months. 11 Qin et al have reported a Chinese patient with advanced lung adenocarcinoma harboring a rare triple EGFR mutation (L833V/H835L/R670W) was resistant to gefitinib but sensitive to afatinib, suggesting that R670W may contribute to resistance to first-generation EGFR TKIs.12 Frega et al presented an Italian patient with advanced NSCLC harboring synchronous EGFR L833V/H835L/E709K mutations who achieved partial response after 2 months of afatinib treatment.¹³ However, whether E709K will impact the sensitivity of EGFR L833V/H835L to first-generation EGFR-TKIs was unclear. It has been reported that the efficacy of afatinib is superior to first-generation EGFR-TKIs in NSCLC harboring uncommon EGFR mutations,

such as G719X and L861Q.^{14,15} Recently, some clinical data showed that afatinib may be effective for other rare mutations, such as exon 18 delE709_T710insD and L858M/L861Q compound mutation.^{16,17} In this study, we found that afatinib could be a first-line treatment option in NSCLC patients with L833V/H835L mutations.

It is worth mentioning that the application of NGS provided an opportunity of targeted therapy for the present patient because the traditional ARMS-PCR would not normally detect the L833V/H835L or other rare mutations. In addition, the L833V/H835L mutation reported in previous studies was mostly detected in tumor tissue. 10,13,18 In this study, the mutation was successfully detected using the circulating ctDNA extracted from pleural effusion and plasma, indicating that liquid biopsy could be an alternative mediator for mutation analysis in patients who are difficult to get tumor specimens. It has been reported that circulating cell free DNA (ccfDNA) is a prognostic marker in advanced NSCLC. 19 Under treatment, ctDNA clearance at first evaluation is associated with objective response and longer survival.²⁰ In this case, the ctDNA was undetectable after 1 month and 7 months of afatinib treatment accompanied with the imaging changes, suggesting a good efficacy of afatinib.

In conclusion, this case report suggests that EGFR L833V/H835L are rare but potentially afatinib responsive mutations in NSCLC. Although it is not realistic to design a clinical trial due to the small number of patients, our case provides more evidence for the clinical application of afatinib in this mutation. Inevitably, a few limitations are associated with this study. How the afatinib take effect on this mutation is not known. A well designed in vitro and in silico studies would provide better understanding of its concrete mechanisms.

Ethical Approval

Institutional approval was not required to publish the case details.

Patient Informed Consent

Written informed consent was obtained from the patient for the publication of his case details and images.

Acknowledgment

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Disclosure of Interest

Tian Qin is an employee of Burning Rock Biotech. The authors declare no other potential conflicts of interest in this work.

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