ADENOCARCINOMA OF THE LUNG WITH RARE INSERTION MUTATION IN EGFR EXON 19 THAT HAD PARTIAL RESPONSE TO GEFITINIB: A CASE REPORT

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Aim: Classic activating mutations L858R and deletions in exon 19 (19del) in the gene for epidermal growth factor receptor (EGFR) are associated with sensitivity of the non-small cell lung cancer (NSCLC) to therapy with tyrosine kinase inhibitors (TKI). Insertions in EGFR exon 19 (19ins) are rare mutations in NSCLC; response of cases with 19ins to TKI is not well studied. Here we report a case of NSCLC with 19ins in a Russian patient who was treated with gefitinib. We also overview cases of 19ins reported in the literature. Case description: A 48 years old female Russian patient was diagnosed with adenocarcinoma of the lung (T3N2M1, stage IV). Mutation 19ins was detected in the tumor biopsy by fragment analysis and genotyped by Sanger sequencing as p.1744_K745insKIPVAI. Treatment with gefitinib (250 mg/day) resulted in clinical and radiological improvements scored as partial response that lasted 12 months. Conclusion: Treatment with gefitinib of lung adenocarcinoma that carries mutation EGFR 19ins can result in durable response.

Key Words: lung adenocarcinoma, EGFR, exon 19 insertion, gefitinib.

Some cases of non-small cell lung cancer (NSCLC) carry activating mutations in the epidermal growth factor receptor (EGFR) gene which are associated with sensitivity of the tumor to tyrosine kinase inhibitors (TKI). Frequency of the mutations varies from 2 to 40% in different groups of patients and positively correlates with adenocarcinoma histology, female gender, non-smoker status and Asian ethnicity [1]. Most of activating mutations in EGFR gene (85–90%) are small in-frame deletions in exon 19 (19del), or missense mutation L858R. Insertions in EGFR exon 19 (19ins) are rare mutations which comprise about 1% among cases of NSCLC with mutant EGFR [2]. Due to the small number of patients response to TKI of cases with 19ins is not well studied. Here we describe a case of NSCLC with 19ins in a Russian patient that was treated with gefitinib.

CASE REPORT

Female Caucasian patient, 48 years old, non-smoker, was diagnosed with lung cancer (T3N2M1, stage IV). Major clinical symptoms were pronounced dyspnea and cough with large amount of sputum (up to 500 ml/day). Computed tomography (CT) scans showed peripheral tumor of the right lung (Fig. 1, a). Thoracoscopic biopsy of the right lung was performed to verify morphological diagnosis. Biopsy of the tumor had histology of adenocarcinoma.

Molecular tests of the DNA from FFPE tumor biopsy did not find common mutations EGFR L858R or 19del. The assays we used are based on allele-specific real-time polymerase chain reaction (PCR) for mutation L858R and wild-type blocking real-time PCR for 19del (BioLink, Russia). Some rare deletions as well as insertions in exon 19 are not included in the real-time PCR assay, therefore we test for these mutations by fragment analysis of the DNA after PCR with forward primer 5’-dGGTGAGAAAGTTAAAATTCCCGTCGC and reverse primer 5’-dTCAGAGATTTCCCGTCTGGC and sequence AAAATCCGTCTCGC (c.2215_2232dup) that resulted in mutation p.1744_K745insKIPVAI (Fig. 2, b). Fragment analysis of the DNA detected insertion in EGFR exon 19 (Fig. 2, a, case # 692).

During the first month after diagnosis the patient had 1 course of chemotherapy (carboplatin AUC5, etoposide 120 mg/m²). After detection of the EGFR mutation patient was switched to gefitinib (250 mg/day). One month after treatment with gefitinib there was pronounced improvement in clinical symptoms (minimal dyspnea at physical exercise, rare cough with small amount of sputum). Due to diffused infiltration of the lung parenchyma and pleuritis it was not possible to select target on CT scans for evaluation of response according to RECIST 1.1. Case was scored as partial response based on clinical improvements in patient performance from ECOG 2 to ECOG 0-1 and radiological improvements (diminished ground-glass opacities and decreased amount of fluid in pleural cavity on CT scans) after 2 months of treatment with gefitinib (Fig. 1, b). CT scans were performed every 2 months. Radiological and clinical disease progression was revealed after 12 months of gefitinib treatment. After that patient was switched to paclitaxel (175 mg/m²) and due to lack of clinical response after 3 courses of paclitaxel the best available supportive care was provided. Patient died 18 months after initiation of treatment.
Published data of response of lung cancers with 19ins to TKI (gefitinib, erlotinib or afatinib) includes 22 cases that were reviewed recently [5]. By searching PubMed in addition to the cases reviewed by Lin et al. [5] we found a report of Arabic female lung cancer patient with 19ins responding to erlotinib [6]. Together with our study total number of patients with 19ins treated with TKI is limited to 24 cases would-wide. Patients were predominantly female non-smokers and most cases were sensitive to TKI. We observed a case of lung adenocarcinoma (stage IV) with mutation p.I744_K745insKIPVAI that had partial response to gefitinib that lasted 12 months. Combined with data from Iyevleva et al. [3] time to progression of Russian patients (n = 4) with 19ins on gefitinib was on average 9.3 months (range 5–12 months). For comparison, Caucasian patients who had lung cancer with mutant EGFR and were treated with gefitinib in the first-line had progression free survival 9.7 months [7]. It appears that sensitivity of tumors with 19ins and duration of response to TKI is similar to classic mutations in EGFR gene, however data on larger number of patients is needed.

In conclusion, our report contributes to small number of cases in other studies indicating sensitivity of the lung adenocarcinoma with insertions in exon 19 to therapy with TKI. The study also shows clinical usefulness of fragment analysis of EGFR exon 19 in addition to real-time PCR assays that do not include mutation 19ins.

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DISCLOSURE

The authors declare no conflict of interest.

REFERENCES