CASE REPORT

The efficacy of erlotinib as molecularly targeted maintenance therapy in advanced non-small-cell lung cancer (NSCLC): Case reports in western Iran

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Received: July 1, 2014
Accepted: October 9, 2014
Online Published: October 24, 2014
DOI: 10.5430/jst.v4n3p38
URL: http://dx.doi.org/10.5430/jst.v4n3p38

Abstract
In developing countries, majority of cancer patients have low income. Therefore, we could hardly do maintenance therapy about the number of patients with non-small-cell lung cancer and other cancers. Maintenance therapy for advanced non-small-cell lung cancer (NSCLC) refers to the use of an effective agent in the absence of disease progression following platinum-based combination chemotherapy to improve progression-free survival. In our Clinic, we identified three cancerous patients with metastasis for lung cancer. The Epidermal Growth Factor Receptor mutation on exon 19 started erlotinib therapy for several courses. After a short time, in all patients, lesions decreased and patients became stable.

Key words
Brain blood barrier, EGFR mutation, Maintenance therapy, NSCLC, TTF-1

1 Introduction
Lung cancer is one of the leading causes of death in the world, and non-small cell lung carcinoma accounts for approximately 75%-85% of all lung cancers [1]. Maintenance therapy for advanced non-small-cell lung cancer (NSCLC) refers to the use of an effective agent in the absence of disease progression following platinum-based combination chemotherapy to improve progression-free survival (PFS) and overall survival. This type of intervention is continued for a determined period of time unless there is disease progression or significant toxicities develop [2]. Epidermal growth factor receptor (EGFR) tyrosine kinase inhibitors (TKIs), such as erlotinib, have been extensively investigated in previously treated advanced NSCLC [3], and in our cases it is clear the efficacy of (EGFR-TKI: erlotinib) in a molecularly targeted maintenance therapy in advanced non-small-cell lung cancer (NSCLC) [3].

Vincenten J et al. [4] reported that thyroid transcription factor-1 (TTF-1) associated with the presence of EGFR mutations in NSCLC patients.
In this study, we describe three patients with metastasis of NSCLC who was treated with maintenance therapy by erlotinib. Also, this study may explains the overall situation of NSCLC cancer patients in the western Iran and more importantly define that TTF-1 and EGFR mutation were effective diagnostic factors in metastasis of NSCLC. MTOR inhibitors in combination with chemotherapy have been found to be additive or synergistic. Compared with single agent therapy, the combination of rapamycin with chemotherapy enhances apoptosis in vitro and enhances anti tumor efficacy in vivo [5-7].

Figure 1. Chemical structure of erlotinib.

2 Case reports

Case A: In March 2012, a 58-year-old man referred to Clinic of Neurology with headache, seizure and right facial weakness with a history of smoking. The MRI of his brain showed brain’s lesions (see Figure 2A). Stereotactic surgery and biopsy were performed, too. In his pathological report, poorly differentiated carcinoma had been confirmed. Moreover, Immunohistochemistry (IHC) showed metastatic adenocarcinoma in brain. The results of markers showed CD20 and CD45 were negative, but (epithelial membrane antigen) EMA, cytoskeratin 7(CK7) and TTF-1 were well positive. It was clear that brain metastasis is due to appearance of primary lung cancer. Due to lower safe of brain lesions (<3 lesions), Gamma Knife radiosurgery was performed for him. He was treated with 4 courses of combination chemotherapy. The EGFR mutation with exon 19 deletion was documented for him. Two months after start of the chemotherapy, results were positive, and he was eligible for TK inhibitor (erlotinib) therapy. After 4 months of erlotinib monotherapy, no lesion was observed in brain MRI and all of them had been resolved (see Figure 2B). The patient was then treated for one year with combination of erlotinib (TK inhibitor) + sirolimus (mTOR inhibitor) + temodar (topoisomerase inhibitor). Unfortunately, 16 months after treatment of combination therapy, the patient suffered of dyspnea. In lung CT, apicoposterior segment lesion of left upper lobe was detected. Again, he was treated with secondary single chemotherapy, for 3 months. Respiratory complaint regressed and lung lesion not progressed. In March 2014, he died with sudden cerebrovascular accident (CVA).

Case B: A 68-year-old woman referred to Clinic of Oncology with cough without a history of smoking. Her image was evaluated. Whole body isotopic bone scan were positive for metastasis. In CT spiral scan of whole body, several randomly distributed pulmonary parenchymal nodules were detected (see Figure 3A). Following CT guided core biopsy; a poorly differentiated/metastatic adenocarcinoma in left lung was detected. In IHC evaluation, TTF-1 was positive. In EGFR mutation analysis, “exon19 deletion” was detected. Regarding this diagnosis, she was eligible to EGFR-TK (erlotinib) therapy. Three months after beginning of erlotinib therapy, lung nodules significantly decreased (see Figure 3B). At now, after 24 months she is stable and all of lung lesions have disappeared and also in bone scan, lesions were sclerosed (see Figure 3B, arrows).

Case C: A 47-year-old woman referred to Clinic of Oncology with complaint of progression dyspnea and weight loss with generalized bone pain without a history of smoking. In evaluation of imagine of her chest CT scan, moderate degree of right pleural effusion with small thickness in anterior portion of pleural, and small lesion at level of RML-lower lobe of bronchi were detected (Figure 4A). In abdominal CT scan a hypodense (19×22 mm) lesion, with extension to
periampullary area of pancreas, was detected. Pleural cytology and biochemical analysis reported inflammatory reaction with lymphocyte predominant. Analyses of her pleural biopsy showed that she suffer from metastatic adenocarcinoma with papillary features. In whole body, isotopic bone metastatic was detected although genitourinary, breasts and thyroid were normal.

In IHC assessments, TTF-1 and CK7 were positive while CK2, CK5/6, CK20, calretinin, ER, PR were negative. In final report/diagnosis, metastatic adenocarcinoma with lung origin was recommended. The paraffin block was evaluated for any changes in exons 18 to 21, and insertional change in exon 19 (as heterozygous) and EGFR mutation was detected. Then, based on this pathology analysis, she was eligible for therapy with EGFR kinase inhibitor. She was treated with erlotinib (150 mg/day), and after 3 months of treatment all complaints of patient reduced and in imaging evaluation, all of lesions decreased more than 50% (see Figure 4B). At now, she is stable, and this policy will be continued for her.

Figure 2. MRI of brain (A) before (B) after erlotinib therapy. A March 2012, B January 2013. Arrows show the tumor (lesion) which has proved to be metastatic adenocarcinoma.

Figure 3. CT scan of Lung (A) before (B) after erlotinib therapy. A July 2012, B February 2013. Arrows show lung lesions improved compared with previous lung CT of patient.
3 Discussion

Our department of oncology is in Kermanshah province, which is one of western provinces in IRAN, and neighbor with Iraq country. In 2009, a study [8] showed that health care payments in western Iran, compared with other developing countries in Asia, are very high. Majority of peoples this province and surrounding provinces are rural and living in (IRAN-IRAQ) war zone. In 2012, Rostaei S et al. [9] reported that 43% of people in Kermanshah city are below the poverty line. Therefore, majority of cancer patients may have low income ($35 - $150 per month). In recent years, due to sanctions on Iran, and since majority of patient samples should be sent to Tehran (capital of Iran) for further evaluation, diagnosis, treatment and drug costs have had a huge increase. Despite financial limitations for majority of patients, because of appropriate clinical facilities in Kermanshah province compared to the neighboring provinces, the number of patients referred to our clinic is very high, and over a few past years we could do maintenance therapy for the number of wealthy cancer patients. In this way, so we could stable lung cancer for three patients (top case reports).

Fortunately during last recent five years Erlotinib registered and available, that facilitate patient treatment policy. EGFR is a member of the type I growth factor receptor family, and The EGFR gene is located in chromosome 7 in the short arm TP12-14 district [10]. The epidermal growth factor receptor (EGFR) is a transmembrane glycoprotein that constitutes one of four members of the erbB family of tyrosine kinase receptors [11]. In the vast majority of cases, EGFR mutations are non-overlapping with other oncogenic mutations found in NSCLC (e.g., KRAS mutations, ALK rearrangements, etc.). More than 90% of EGFR mutations are located in the exons 18 to 21, and the most common mutations lie in exons 19 and 21 [12]. In EGFR gene, exons 18-21 are analyzed in tumor-derived DNA [13].

Exon 19 deletion mutations occur in a protein strand (called the β3 strand) adjacent to the C-helix. Similarly, the exon 19 insertion mutations all lead to substitution of residue L747 [14]. In three listed cases, EGFR mutation happened in exon 19, and erlotinib consumption improved metastasis in all of them. So may be patients with exon 19 (deletion/insertion) respond more positively to erlotinib compared with 18 or 21. In future researches is better that this point be considered.

This study showed that erlotinib was effective in the elderly patients in this case series. Although this report includes a small number of patients, but it demonstrates that erlotinib can be a suitable treatment for lung cancer patients in conditions that patients can’t take expensive drugs.

One series found PIK3/CA mutations in 5% of cases of acquired resistance [15], and several trials combining EGFR TKIs with PI3K inhibitors are underway [16].

Figure 4. CT scan of Lung (A) before (B) after erlotinib therapy. A December 2013, B April 2014. Arrows show the tumor (lesion) which has proved to be metastatic adenocarcinoma with lung origin.
A retrospective study has also show higher response rates in setting when chemotherapy is given with a TKI\cite{17}.

In a randomized study conducted in the first-line setting, erlotinib plus chemotherapy was found to be somewhat more toxic than chemotherapy, but no antagonism was seen; efficacy was equal in patients with untreated EGFR-mutant lung cancer\cite{18}.

Thyroid transcription factor-1 (TTF-1) is a 38-kDa nuclear protein initially identified as a mediator of thyroid-specific gene transcription\cite{19}. TTF-1 is a member of the NKX2-1 family; it is expressed in the thyroid, forebrain, lung and other organs, and is a sensitive marker for pulmonary and thyroid adenocarcinomas\cite{20}. Results of Lee JS et al\cite{21} showed that TTF-1 amplification might be a predictive marker of poor response to EGFR-TKI therapy in patients with recurrent tumor after surgical resection. Vallee A et al.\cite{22} reported that EGFR mutation rate was much higher in tumors expressing the TTF-1 antigen than in TTF-1 negative tumors. Also, a study\cite{23} reported that erlotinib may to cross the blood-brain barrier, and other study\cite{24} on patients who underwent whole-brain radiotherapy demonstrated that this procedure may increase the permeability of the blood-brain barrier and eventually lead to a change in the erlotinib concentration in the cerebrospinal fluid. Togashi Y et al.\cite{25} suggested that 150 mg daily erlotinib can increase concentration of erlotinib in cerebrospinal fluid, so erlotinib could be effective for the treatment of central nervous system (CNS) metastases of NSCLC. Of three listed patients, one patient had lung cancer with metastasis of brain. Consumption of erlotinib in patients led to improved of metastasis. It is probably that erlotinib could cross of BBB. Confirmation of this hypothesis needs to more comprehensive studies.

Results of the study suggested that first of all, these case reports show that maintenance therapy with erlotinib according to molecular markers (TTF-1 and kind of EGFR mutation) can help survival rate in patients. Second, in this report illustrates erlotinib may to cross the blood-brain barrier.

**Acknowledgement**

The authors thank to company of AqVida GmbH, Hamburg (Hamburg, Germany) for their financial support.

**References**


