Introduction

EGFR-TKI resistance and mechanisms

Epidermal growth factor receptor tyrosine kinase inhibitors (EGFR-TKIs) are current treatments for advanced non-small cell lung cancer (NSCLC) harboring activating EGFR gene mutations. Although studies show an increased progression free survival (PFS) with use of EGFR TKIs in the first-line setting, most patients will develop resistance to therapy after the first about 10 months. Undoubtedly it is critical to choose an optimal clinical strategy for patients with EGFR sensitive mutation undergoing EGFR-TKI resistance. The second biopsy should be applied under condition permission to verify specific resistance mechanism. Here we discussed the mechanism of drug resistance and the choice of therapeutic regimen, also compared the superior and inferior of each treatment plan, which proposed a novel perspective for NSCLC target therapy.
for SCLC transformation; associated pathway inhibitors combined with TKI treatment can be applied for genes abnormal activation and amplification. Currently, all the MET, HER2, HER, IGF1R, AXL, MEK, AKT, PI3K, HSP90 inhibitors are in the development of clinical research, of which the outcomes are worth looking forward to[9,10].

T790M, the second mutation of EGFR gene, is considered to be the main causes of primary EGFR-TKI resistance, accompany with secondary resistance of EGFR-TKI[11,12]. Studies have shown that T790M changes the spatial configuration of EGFR tyrosine kinase domain, which hinders the binding of TKI to EGFR receptor and weakens the effective competitive ability between TKI and ATP, eventually leading to resistance. Clinically, T790M mutations are commonly associated with sensitive mutations[4,5]. In recent years, according to newly reports about T790M mutation from China, Japan and United States, using various detection methods including arrays, direct sequencing, PCR-RFLP, qPCR respectively, investigators found the presence of T790M mutation may be more extensive than we thought, even up to 60-80%, fully illustrated that T790M mutation is the most common cause of EGFR-TKI resistance[4,6,13]. In addition Kuang, Y., et al discovered T790M genes can only be detected from 54% patients whose plasma DNA were collected from CR patients in EGFR-TKI targeted therapy and there was no significant difference compared with tumor tissue sequencing data. However, examining the plasma of patients with known EGFR T790M mutation showed 71% (5/7) of the patients whose plasma DNA was able to detect T790M EGFR, declaring EGFR T790M can be detected from plasma DNA of EGFR-TKI resistant patients and this non invasive method is also useful for monitoring drug resistance in the future, providing supplementary basis for guideline of subsequent drug use[14]. The ultimate aim of studying T790M mutation was to find an effective drug that can overcome TKI resistance associated with mutations. AZD9291 is the third generation oral drug and irreversible selective EGFR mutated inhibitor, which possessed strong anti-tumor efficacy and high selectivity for both EGFR-TKI sensitive mutations (EGFR+) and drug resistance mutations (T790M). The overall response rate and clinical benefit rate for T790M positive patients were 64% and 90%, both of which were extremely higher than negative ones (23% and 64%), and PFS was also up to 13.5 months[15]. What’s more, Rocaletinib (CO-1686), as another third generation EGFR-TKI, whose overall response rate and clinical benefit rate reached to 64% and 90% respectively and PFS got up to 8-10 months was also achieved good therapeutic effect[16]. In conclusion, the third generation of EGFR-TKI will undoubtedly prove to be the essential substances to overcome drug resistance caused by T790M. Currently which is being studied for the third generation of EGFR-TKI includes HM61713, EG816, ASP8273, besides domestic is also developing third generation EGFR-TKI, such as Avitinib and BPI-15086. According to the T790M mutation status, the current prevailing perception for NSCLC patients with EGFR sensitive mutation is, all the patients with resistance to the first generation or the second generation EGFR-TKI can be divided into two classes: Patients with second mutations of T790M will achieve good therapeutic effect adopted by the third generation of EGFR-TKI, however for the resistant patients with T790M negative status, clinical studies on the mechanisms of drug resistance may be the best choice[17].

Currently how to deal with NSCLC patients with resistance to the first generation of EGFR-TKI in clinical, especially before the third generation of EGFR-TKI listed?

Clinical retrospective study of Faehling, M., et al found patients with previously effective erlotinib treatment who developed disease progression applying continuous EGFR-TKI therapy even can acquire an average survival time of 14.5 months or more, overall survival of nearly 50 months since the diagnosis. However patients who discontinued TKI treatment obtained an average survival time of only 2 months after progression, and overall survival of 28 months since the diagnosis, which was significantly lower than that of patients who continued to use TKI therapy after disease progression[18]. The Japanese scholar Hosomi Y., reported a multicenter, prospective observational study in ASCO 2014 on assessment of PD. Clinical practice treatment mode and curative effect of NSCLC patients with EGFR mutation positive receiving the first-line TKI therapy based on RECIST criteria and the observation showed PFS (R-PDFS) was 264 days while the clinical judgment of disease progression (C-PDFS) was almost 461 days. Patients developed without obvious clinical progression can acquire clinical stability for 6 months if performing continuous TKI treatment after disease progression. What’s more, the prognosis of patients with initial TKI therapy appeared clinical progression was significantly better than that of patients with discontinued TKI treatment[19]. 2014 ESMO results from ASPIRATION, a prospective, randomized controlled study investigated by Park K et al suggested, continued erlotinib treatment after the RECIST PD can prolong PFS for 3.1 months (from 11.0 to 14.1 months) with no new adverse events[20]. 2015 NCCN guidelines recommended of patients should apply erlotinib continual therapy when progress appeared in imaging diagnosis but no clinical symptoms were found. Retrospective investigation published by Nishie K., et al displayed NSCLC patients with EGFR sensitive mutation applied ongoing Gefitinib treatment followed with disease progression by TKI can acquire the median survival time for 32 months, whereas only 23 months were appeared among patients who received replacement for chemotherapy[21]. Therefore, from a point of view, for patients with initial effective treatment of TKI, disease progress in imaging diagnosis but no significant symptoms, both of disable TKI and replacement with chemotherapy is inferior to continual use for TKI to benefit patients. Undoubtedly, it is necessary to use chemotherapy for patients with resistance, rapid progression of disease or obvious clinical symptoms, even cannot continue acquired benefit.

Which is a better choice comparing chemotherapy alone and TKI combined with chemotherapy?

A retrospective analysis on Comparing with the effects of single drug chemotherapy and erlotinib with chemotherapy combination in patients with acquired resistance which implanted by Goldberg SB, at Massachusetts General Hospital found Whatever adopting single drug chemotherapy or combined with platinum chemotherapy, even erlotinib with chemotherapy combination, the PFS are not superior than that of single chemotherapy[22]. 2014 ESMO Tony Mok reported the outcomes of IMPRESS study Which is a randomized, double-blinded, placebo-controlled phase III global multi center clinical trial, aimed to compare the efficacy and
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safety of ongoing Gefitinib associated with chemotherapy with single chemotherapy in advanced Gifitinib resistance NSCLC pa-
tients. Research results show that the two main endpoint was PFS was 5.4 months, Orr and DCR without significant difference,
consequently it is not recommended adopting the second-line chemotherapy which contained two drugs based on platinum with
Gefitinib combination for patients with disease progression[23].

**How to choose the optimal therapy strategy at the end of the second line chemotherapy?**

Namba Y, in a retrospective research discovered the median survival time can reached 955 days for rechallenge TKI ther-
apy; whereas only 615 days of median survival time can be reached if abandoning TKI rechallengement, suggesting the survival time
was obviously higher for patients who received TKI rechallenge therapy than not (p = 0.0194). Nevertheless, T790M disappearance
may occur in some part of patients undergoing chemotherapy, also with the second mutations of T790M, which may be explained
as a new effective process for patients re-sensitive and re-challenged to TKI.

In addition, the survival time of patients who received multiple line therapies followed with TKI treatment was significant-
ly longer than those who only received first-line or less line therapy[24]. Therefore it is necessary for patients to administrate with
EGFR-TKI therapy as long as possible be under the premise of drug tolerance, however, if the patients cannot acquire evidential
benefits of clinical symptoms, the second line chemotherapy should be performed, and TKI treatment should not be administrated
until no more use of chemotherapy for patients. What’s more, it is possible for NSCLC patients to be able to achieve benefits from
TKI rechallenge therapy or multi line treatment at the end of the second line chemotherapy, and tumor resistance rebiopsy or partic-
ipation in clinical research are also recommended for those who have failed in target therapy.

**How to perform the further cure facing the acquired resistance of the third generation TKI?**

Obviously patients with T790M (+) should be given the third generation TKI treatment[25]. A study published in the Nat
Med 2015 discovered cell growth conditions using BA/F3 cells with L858R or T790M mutation which transected with C797S mu-
nants or not in different concentrations of AZD9291 treatments. The data displayed the susceptibility of L858R or T790M mutated
Ba/F3 cells transfected with C797S mutant to AZD9291 was startling attenuated, which was also occurred in CO-1686,as the second
generation of selective mutation EGFR-TKI targeted drug, exerting the good function to T790M positive NSCLC patients. In gen-
eral, C797S mutation may be a mediator of AZD9291 acquired resistance in lung cancer patients, consequently, it is suggested that
effective targeting therapy for C797S mutations need to be managed in patients with AZD9291 acquired resistance[26]. Nowadays,
the new use of old drugs is also emerging. 2014 Clin Cancer Res reported that the combined treatment with metformin and TKI may
reverse the efficacy of EGFR mutation NSCLC patients and prolong the overall survival time, which proposed a novel perspective
for NSCLC target therapy[27].

**Conclusion**

In summary, NSCLC patients undergoing EGFR-TKI acquired resistance are not a homogeneous population, but a col-
lection of different types of patients which cannot be treated with the same method. We should deal with each case on its merits in
clinical practice: The third generation of EGFR-TKI was the best choice for patients with T790M acquired resistance; applying the
second biopsy under the condition permit to verify the specific mechanism of drug resistance; Distinguish various progresses for
patients, such as slow, asymptomatic or local progression, and the continual EGFR-TKI (or combined with local therapy) therapy
should not suspended until the patient can no longer acquired benefit from the TKI treatment. The effective second line therapy
such as cisplatin or pemetrexed can relieve if symptomatic, rapid and extensive progress were appear, whereas if it is on the basis of
combination chemotherapy, the application of EGFR-TKI should not be continued. Undoubtedly, which option on EGFR-TKI treat-
ment undergo disease progress to take in the future is still needed to further distinguish the development pattern and the resistance
mechanism, consequently adopting individualized therapy strategy.

**References**

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