Correlation of circulating tumor DNA EGFR mutation levels with clinical outcomes in patients with advanced lung adenocarcinoma

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Abstract
Background: Circulating tumor DNA (ctDNA) is a promising biomarker for non-invasive epidermal growth factor receptor mutations (EGFRm) detection in lung cancer patients, but existing methods have limitations in sensitivity and availability. In this study, we used the ΔCt value (mutant cycle threshold [Ct] value–internal control Ct value) generated during the polymerase chain reaction (PCR) assay to convert super-amplification-refractory mutation system (superARMS) from a qualitative method to a semi-quantitative method named R-superARMS (R-superARMS), and evaluated its performance in detecting EGFRm in plasma ctDNA in patients with advanced lung adenocarcinoma.

Methods: A total of 41 pairs of tissues and plasma samples were obtained from lung adenocarcinoma patients who had known EGFRm in tumor tissue and were previously untreated. EGFRm in ctDNA was identified by using superARMS. Through making use of ΔCt value generated during the detection process of superARMS, we indirectly transform this qualitative detection method into a semi-quantitative PCR detection method, named R-superARMS. Both qualitative and quantitative analyses of the data were performed. Kaplan–Meier analysis was performed to estimate the progression-free survival (PFS) and overall survival (OS). Fisher exact test was used for categorical variables.

Results: The concordance rate of EGFRm in tumor tissues and matched plasma samples was 68.3% (28/41). At baseline, EGFRm-positive patients were divided into two groups according to the cut-off ΔCt value of EGFRm set at 8.11. A significant difference in the median OS (mOS) between the two groups was observed (EGFRm ΔCt ≤8.11 vs. >8.11: not reached vs. 11.0 months; log-rank P = 0.024). Patients were divided into mutation clearance (MC) group and mutation incomplete clearance (MIC) group according to whether the ΔCt value of EGFRm test turned negative after 1 month of treatment. We found that there was also a significant difference in mOS (not reached vs. 10.4 months; log-rank P = 0.021) between MC group and MIC group. Although there was no significant difference in PFS between the two groups, the two curves were separated and the PFS of MC group tended to be higher than the MIC group (not reached vs. 27.5 months; log-rank P = 0.088). Furthermore, EGFRm-positive patients were divided into two groups according to the cut-off of the changes in ΔCt value of EGFRm after 1 month of treatment, which was set at 4.89. A significant difference in the mOS between the two groups was observed (change value of ΔCt >4.89 vs. ≤4.89: not reached vs. 11.0 months; log-rank P = 0.014).

Conclusions: Detecting EGFRm in ctDNA using R-superARMS can identify patients who are more likely sensitive to targeted therapy, reflect the molecular load of patients, and predict the therapeutic efficacy and clinical outcomes of patients.

Keywords: Lung adenocarcinoma; Non-small cell lung adenocarcinoma; Liquid biopsy; Super-amplification-refractory mutation system; EGFR mutation

Introduction
The guidelines recommend non-small cell lung cancer (NSCLC) patients with epidermal growth factor receptor mutations (EGFRm) can be treated with tyrosine kinase inhibitors (TKIs), but in clinical practice, the effective rate of targeted therapy in patients with EGFR positive mutation was only about 70%, and most patients develop resistance within 9 to 12 months. Therefore, detection of EGFRm and monitoring the efficacy of treatment are of great significance in such patients. The majority of EGFR genotyping is assessed by conventional tumor biopsy, which however, has several shortcomings: the tissue specimen showed heterogeneity of time and space; it is hard to obtain biopsy tissue after recurrence or metastasis; and intrathoracic biopsy is easy to cause complications. In addition, the clinical evaluation of...
the therapeutic effect of solid tumor is currently based on
alterations in imaging; however, patients with the same
degree of imaging response have different treatment
outcomes, supposing that there exist different degrees of
tumor molecular response to guide treatment options and
to predict prognosis. Recently, with the development of
technology, liquid biopsy, especially circulating tumor
dNA (ctDNA) detection, can bring us enough informa-
tion, present information earlier than changes in imaging,
and also overcome the limitations of tissue biopsy.[10-12]

At present, several molecular detection methods have been
developed, such as amplification-refractory mutation
system (ARMS), droplet digital polymerase chain reaction
(ddPCR), and next generation sequencing (NGS).[13-15] The
ARMS has been approved by the European Union and
National Medical Products Administration (NMPA) for
clinical testing of EGFRm using tissue or blood samples.
However, it is only a qualitative detection technique and the
sensitivity (48.2–67.4%) needs to be improved compared to
the matched tumor tissue biopsy.[16,17] ddPCR and NGS
have relatively high sensitivity and absolute quantification.
But the development of ddPCR kit is not easy, and
the construction of NGS database is complicated and
the standardization of NGS detection platform needs to be
improved, which makes these two methods more suitable
for translational medical research[18,19] rather than clinical
practice. Most recently, a newly developed method based on
ARMS, super-amplification-refractory mutation system
(superARMS), has been available for detecting EGFRm
status in the plasma ctDNA with a qualitative polymerase
chain reaction (PCR) method claimed to be highly selective,
sensitive, and economic, and has been approved by (NMPA)
for detecting the status of ctDNA EGFRm in patients with
advanced NSCLC and conventionally applied to clinical
practice.[20,21] However, it is still not a quantitative
detection method, which limits its application in further
clinical research.

In this study, all patients were routinely applied the
superARMS method for plasma ctDNA detection. We
proposed a reformed superARMS method named re-
formed-superARMS (R-superARMS). Through making
use of ΔCt value generated during the detection process, we
indirectly transform this qualitative detection method into
a semi-quantitative PCR detection method. This research
aimed to study the effects of different baseline ctDNA
EGFRm levels and changes of ctDNA EGFRm with R-
superARMS on the outcomes and prognosis of patients
receiving targeted therapy.

Methods

Ethical approval

This study was approved by the Institutional Review Board
of the First Hospital of Jilin University (No. NCT2017-
444). Informed consent was obtained from all patients.

Patients and treatment

This study was conducted in the Cancer Center, the First
Hospital of Jilin University in Changchun, China. A total
of 41 Patients were enrolled from January 2017 to June
2018. Inclusion criteria are described as follows: (1)
pathologically confirmed lung adenocarcinoma; (2) ad-
vanced clinical stage (stage IIIb or IV); (3) newly diagnosed
or recurrence after surgery; (4) a subsequent treatment did
not begin at the time of sample collection; and (5) tumor
tissue and blood specimens were available and the interval
between tissue collection and blood collection was less
than 14 days.

All enrolled patients received first-generation TKIs mono-
therapy (erlotinib, gefitinib, or cetinib) or combination
with antiangiogenic drugs (bevacizumab or apatinib) as
first-line treatment. Antitumor drugs were used according
to the routine clinical practice (National Comprehensive
Cancer Network Clinical Practice Guidelines in Oncology:
NSCLC). Patient data were collected from medical records
and radiological images were used to assess baseline
information, tumor response, and progression-free surviv-
(al (PFS). After tumor progression, patients can withdraw
from the study using other treatment options. Tumor
assessment was performed using the response evaluation
criteria in solid tumors (RECIST) Version 1.1, and
EGFRm in plasma was assessed at baseline and 1 month
after treatment. A schematic representation of the study is
shown in Figure 1. The primary endpoint was PFS. Overall
survival (OS) was evaluated as a secondary endpoint.

DNA extraction

Plasma was separated from the whole blood within 2 h of
collection. ctDNA was extracted from the plasma within
2 h by AmoyDx circulating DNA kit (Amoy Diagnostics,
Xiamen, China) and stored at –80°C before use. The
optical density of the extracted DNA samples was
measured using a NanoDrop 2000 spectrophotometer
(Thermofisher Scientific, Carlsbad, CA, USA). The A260/
A280 value of all samples was 1.8 to 2.0. The extracted
DNA was used for EGFR molecular testing.

Detection of ctDNA EGFRm

The EGFRm in plasma ctDNA was detected by super-
ARMS EGFRm detection kit (Amoy Diagnostics, Xiamen,
China), which can identify 41 types of the most common
somatic EGFR mutations in exons 18 to 21. The assay
was performed using a real-time PCR according to the
manufacturer’s instructions, while PCRs were performed
using Stratagene Mx3000P quantitative PCR system
(Agilent, Santa Clara, CA, USA). All reactions were
carried out in a volume of 70 μL DNA sample (or negative/
positive controls), plasma epidermal growth factor recep-
tor (P-EGFR) enzyme mix, and P-EGFR reaction mix. PCR
amplification conditions were performed with 10 min incubation
at 95°C, followed by 15 cycles at 95°C for 40 s, 64°C for
40 s, and 72°C for 30 s, and then 28 cycles of 93°C for 40 s,
60°C for 45 s, and 72°C for 30 s, respectively. ΔCt value
(mutant cycle threshold [Ct] value-internal control Ct
value) was calculated to identify the presence of EGFRm,
which was automatically calculated from PCR amplifica-
tion plots of fluorescence and its corresponding num-
ber of cycles. The ΔCt cut-off value was defined as 11 (19Δd/
L858R/20ins), 8 (T790M), or 12 (G719X/L861Q/S768L).
When the ΔCt value was less than the cut-off value, the sample was considered as mutant. Mutation clearance (MC) group refers to that the ΔCt value of original EGFRm completely changed to the negative mutation value set above, and mutation incomplete clearance (MIC) group refers to that although the ΔCt value of EGFRm changed, it was still positive and did not completely turn negative.

**Statistical analysis**

The concordance rate was assessed by comparing the match between primary tissue and ctDNA sequencing results. The Kaplan–Meier analysis was performed to estimate survival probabilities. A receiver operating characteristic (ROC) curve was applied to calculate the cutoff values for the parameters (ΔCt value of EGFRm or the changes in ΔCt value of EGFRm) and area under the curve (AUC). The cutoff point was determined using Youden index. Fisher exact test was used for analysis of categorical variables. The hazard ratio for each cutoff point was calculated using the results from each of the corresponding survival analyses. The PFS was defined as the time from the start of the TKIs treatment to the first documentation of tumor progression or the last follow-up. OS was defined as the time from the start of follow-up to the death of patients or the end of follow-up. Throughout the analyses, all P values were based on a two-sided hypothesis, and those less than 0.05 were considered statistically significant. All statistical analyses were performed using GraphPad Prism version 6.0 for Mac and SPSS 22.0 (IBM Corp., Armonk, NY, USA).

**Results**

**Characteristics of patients**

A total of 41 lung adenocarcinoma patients met the criteria and were enrolled into the study. The patients’ clinical characteristics are listed in Table 1. The mean age was 63 years with a range of 39 to 75 years. Most patients were female and non-smokers. About 51.2% (21/41) had extrathoracic metastatic diseases (M1b; mainly for bone metastasis and brain metastasis) and 48.8% (20/41) only had intrathoracic metastatic diseases. About 68.3% (28/41) received first-generation epidermal growth factor receptor tyrosine kinase inhibitors (EGFR-TKIs) mono-therapy, and others received first-generation EGFR-TKIs combined with antiangiogenic drugs.

**Detection of ctDNA EGFRm using superARMS and R-superARMS**

The EGFRm status in 41 pairs of tissue and matched plasma samples collected before the administration of TKIs treatment was tested by superARMS, and the concordance rate was 68.3% (28/41). Individual patient results are reported in Table 1. EGFR 19del (31.7%, 13/41) and L858R mutations (29.3%, 12/41) were the two most common type of mutations. Other rare mutation types included G719X and T790M. About 31.7% (13/41) were negative for EGFRm detected by superARMS in plasma samples but positive in the tissues.

**Baseline levels of ctDNA EGFRm affect the outcomes of patients**

The median PFS of all patients was 14.9 months. During a median follow-up period of 2.5 years, disease progression and death occurred in 36.6% (15/41) and 24.4% (10/41) of patients, respectively. All adverse events were tolerable and no treatment-related deaths occurred. Considering the variability in the effective rate of targeted therapy in patients with EGFR positive mutation, the semi-quantitative R-superARMS method was used in our study to detect the different values of EGFRm in EGFRm-positive patients. We set the cut-off ΔCt value of baseline EGFRm at 8.11 using ROC curve (AUC = 0.579; Youden index = 0.236).
Before survival analysis, we analyzed the equilibrium of clinical characteristics (including gender, age, treatment, and tumor stage) between different groups with baseline $\Delta$Ct values of $EGFR_m \leq 8.11$ and $>8.11$, and found $P > 0.05$ for all comparisons, demonstrating that the balance was good [Table 2].

Firstly, all patients were divided into $EGFR_m$-negative and $EGFR_m$-positive groups by detecting baseline ctDNA $EGFR_m$ status in plasma samples using superARMS, and found that there was no significant difference in the median progression-free survival (mPFS) (22.2 vs. 27.5 months; log-rank $P = 0.914$) and median OS (mOS) (22.2 months vs. not reached; log-rank $P = 0.683$) between the two groups. Then the OS was compared between $\Delta$Ct value $>8.11$ group and $\leq 8.11$ group. As shown in Figure 2, the patients with $\Delta$Ct $\leq 8.11$ had better OS (not reached vs. 11.0 months; log-rank $P = 0.024$).

### Changes of ctDNA $EGFR_m$ during treatment affect the outcomes of patients

Next, we detected the changes of ctDNA $EGFR_m$ during treatment by qualitative and quantitative analysis with R-superARMS, and observed the effects of $EGFR_m$ changes on patients’ outcomes. According to the MC status after one month of treatment, 28 patients with positive $EGFR_m$ were divided into MC group (67.9%, 19/28) and MIC group (32.1%, 9/28). No significant difference was detected between the MC and MIC groups in clinical characteristics (all $P > 0.05$) [Table 2]. The results showed that mOS between two groups (not reached vs. 10.4 months; log-rank $P = 0.021$) had significant difference. Although there was no statistical difference in PFS between the two groups, the two curves were separated and the PFS of MC group tended to be higher than the MIC group (not reached vs. 27.5 months; log-rank $P = 0.088$) [Figure 3]. Moreover, quantitative analysis of the data was performed to further clarify whether changes in the amount of mutation affect the outcomes of the patients. The cut-off value of the changes in $\Delta$Ct value of $EGFR_m$ after one month of treatment derived from the ROC curve was set at 4.89 (AUC $= 0.737$; Youden index $= 0.600$). There was no significant differences between the two groups in clinical

### Table 2: Comparison of clinical characteristics between groups with different ctDNA $EGFR_m$ $\Delta$Ct values.

| Items                   | Baseline ctDNA $EGFR_m$ $\Delta$Ct values | Qualitative changes of ctDNA $EGFR_m$ 1 month after treatment | Quantitative changes of ctDNA $EGFR_m$ $\Delta$Ct values 1 month after treatment |
|-------------------------|------------------------------------------|---------------------------------------------------------------|--------------------------------------------------------------------------|
|                         | $\leq 8.11$ ($n = 23$) | $>8.11$ ($n = 5$) | MC group ($n = 19$) | MIC group ($n = 9$) | $\leq 4.89$ ($n = 12$) | $>4.89$ ($n = 16$) | $P$ |
| Age (years)             |                                         |                                                               |                                                                         |                                                                          |                                                                         |                                                                         |
| $<65$                   | 12 (52.2)                               | 4 (80.0)                                                     | 0.36                                                                    | 12 (63.2)                                                              | 4 (44.4)                                                               | 0.43                                                                    | 8 (66.7)                                                               | 8 (50.0) | 0.46    |
| $\geq 65$               | 11 (47.8)                               | 1 (20.0)                                                     | 1.00                                                                    | 7 (36.8)                                                              | 5 (55.6)                                                               | 0.68                                                                    | 5 (41.7)                                                              | 5 (31.2) | 0.70    |
| Gender                  |                                         |                                                               |                                                                         |                                                                          |                                                                         |                                                                         |                                                                          |                                                                  |
| Male                    | 8 (34.8)                                | 2 (40.0)                                                     | 1.00                                                                    | 6 (31.6)                                                              | 4 (44.4)                                                               | 0.68                                                                    | 5 (41.7)                                                              | 5 (31.2) | 0.70    |
| Female                  | 15 (65.2)                               | 3 (60.0)                                                     | 1.00                                                                    | 13 (68.4)                                                             | 5 (55.6)                                                               | 0.68                                                                    | 7 (58.3)                                                              | 11 (68.8) | 0.70    |
| Tumor stage             |                                         |                                                               |                                                                         |                                                                          |                                                                         |                                                                         |                                                                          |                                                                  |
| M1a                     | 10 (43.5)                               | 2 (40.0)                                                     | 1.00                                                                    | 8 (42.1)                                                              | 4 (44.4)                                                               | 1.00                                                                    | 7 (58.3)                                                              | 5 (31.2) | 0.25    |
| M1b                     | 13 (56.5)                               | 3 (60.0)                                                     | 1.00                                                                    | 11 (57.9)                                                             | 5 (55.6)                                                               | 1.00                                                                    | 9 (75.0)                                                              | 7 (43.8) | 0.14    |
| Treatment               |                                         |                                                               |                                                                         |                                                                          |                                                                         |                                                                         |                                                                          |                                                                  |
| First-generation EGFR-TKIs | 12 (52.2)                               | 4 (80.0)                                                     | 0.36                                                                    | 11 (57.9)                                                             | 5 (55.6)                                                               | 1.00                                                                    | 9 (75.0)                                                              | 7 (43.8) | 0.14    |
| First-generation EGFR-TKIs + anti-angiogenic drugs | 11 (47.8) | 1 (20.0) | 0.36 | 11 (57.9) | 5 (55.6) | 1.00 | 9 (75.0) | 7 (43.8) | 0.14 |

Data were presented as $n$ (%). ctDNA: Circulating tumor DNA; $EGFR$: Epidermal growth factor receptor; $EGFR_m$: Epidermal growth factor receptor mutation; MC: Mutation clearance; MIC: Mutation incomplete clearance; TKIs: Tyrosine kinase inhibitors.
characteristics (all $P>0.05$) [Table 2]. In the group with the change value of $\Delta Ct >4.89$, mOS was not reached, while in the group with the change value of $\Delta Ct \leq 4.89$, mOS was 11.0 months, showing a significant difference (log-rank $P = 0.014$). However, there was no significant difference in mPFS (27.5 vs. 11.0 months; log-rank $P = 0.199$) [Figure 4].

In addition, we also analyzed the relationship between the change in the amount of mutations and the degree of response based on imaging in EGFRm-positive patients. It was found that patients in the MC group showed higher proportions of stable disease (SD) (4/12, 33.3%) and partial response (PR) (5/12, 41.7%) upon RECIST standard (about 2 months after treatment) than those of patients in the MIC group (SD: 1/5, 20.0%; PR: 1/5, 20.0%), with limited available data. This phenomenon was also seen when the group with changed value of $\Delta Ct >4.89$ (SD: 3/12, 25.0%; PR: 6/12, 50.0%) was compared with the group with changed value of $\Delta Ct \leq 4.89$ (SD: 1/5, 20.0%; PR: 1/5, 20.0%).

**Discussion**

In this study, we indirectly make use of $\Delta Ct$ value generated during the detection process and transform superARMS from a qualitative into a semi-quantitative PCR detection method, and subsequently apply it to detect any change in ctDNA EGFRm. We have named it R-superARMS, and demonstrated its applicability for detecting EGFRm status in plasma as a simple, fast, and semi-quantitative PCR method that is claimed to be highly selective and sensitive. In our study, the concordance rate of EGFRm in tissue and matched plasma samples using superARMS was 68.3%, which was similar to previous research results.[19,22] However, given that there is a certain time interval between tissue collection and blood
collection, our results may be biased. We also explored the effects of different baseline levels of $\text{EGFR}^{\text{m}}$ in plasma on the outcomes in patients receiving targeted drugs, and found that the baseline status of ctDNA $\text{EGFR}^{\text{m}}$ has an impact on the survival of all patients. Those with positive mutations detected in blood tests were more suitable for targeted therapy. Considering the effective rate of targeted therapy in patients with positive $\text{EGFR}^{\text{m}}$ is only 70%,[1-3] we supposed it is possibly related to the amount of initial $\text{EGFR}^{\text{m}}$. We set the cut-off $\Delta \text{Ct}$ value at 8.11 and found survival curves of the two groups were significantly different, which might partly explain why not all $\text{EGFR}^{\text{m}}$-positive patients were responsive to TKIs. When the $\Delta \text{Ct}$ value of baseline $\text{EGFR}^{\text{m}}$ is less than 8.11, the clinical outcome of patients receiving targeted therapy may be better, while for patients with positive $\text{EGFR}^{\text{m}}$ but $\Delta \text{Ct}$ value greater than 8.11, we consider that the application of targeted therapy may be not effective. Therefore, quantitative detection methods for $\text{EGFR}^{\text{m}}$ may be a good strategy for screening and identifying patients who are more sensitive to targeted therapy in clinical practice. However, our study has a shortcoming that patients excluded from the study due to disease progression may receive other treatment regimens, which had an impact on the overall evaluation of OS. Therefore, the relationship between indicators, such as changes in ctDNA and OS, should be interpreted with caution. More large-scale clinical trials are needed for further confirmation, while our study only provides an idea and direction for future prospective large-scale studies. Previously, Park et al.[23] reported that the corrected delta cycle threshold value, which refers to $\text{EGFR}$ quantification of small biopsy tissue or cytology specimens by peptide nucleic acid-mediated clamping PCR, can predict better objective response and clinical benefit rates of $\text{EGFR}$-TKIs therapy. This is also an important study of $\text{EGFR}^{\text{m}}$ quantitative detection method in identifying the efficacy of targeted therapy, but unlike our quantitative detection of liquid biopsy. This is because microdissecting the tiny tissue obtained by bronchoscopy or needle biopsy is not easy, while the study by Park et al.[23] reflected the clinical limitations of tissue biopsy compared with liquid biopsy.

**Figure 4:** Survival analysis (PFS [A] and OS [B]) of the groups with changes in $\Delta \text{Ct}$ values of ctDNA. $\text{EGFR}^{\text{m}} > 4.89$ and $\leq 4.89$. ctDNA: Circulating tumor DNA; $\text{EGFR}^{\text{m}}$: Epidermal growth factor receptor mutations; OS: Overall survival; PFS: Progression-free survival.

RECIST v1.1[24] is commonly used in clinical practice to evaluate the therapeutic efficacy in patients with solid tumor; however, it is well-known that imaging changes in tumor patients lag behind the occurrence of tumor molecular events.[13,15,18,25] The idea of using molecular response to evaluate the efficacy in patients is also reflected in the previously reported case of imatinib treatment in chronic myeloid leukemia,[26] which confirmed the rationale of our research. We detected ctDNA $\text{EGFR}^{\text{m}}$ by qualitative and quantitative methods using R-superARMS. In qualitative analysis, the TKI-treatment outcomes of patients in the MC group tended to be better than those in the MIC group, with OS between two groups being significantly different. Our study showed that ctDNA concentration of patients, who responded to treatment, decreased significantly since the initial treatment, with some decreasing to even undetectable levels. In the phase III AURA3 trial,[27] clearance of plasma $\text{EGFR}^{\text{m}}$ after 3/6 weeks of first-line treatment with TKIs was associated with a numerical improvement in PFS. Quantitative analysis of the data was performed using R-superARMS, and we found that when 4.89 was set as the cut-off of the changes in $\Delta \text{Ct}$ value of $\text{EGFR}^{\text{m}}$ in $\text{EGFR}^{\text{m}}$-positive patients. After 1-month treatment, PFS analysis showed that the results tended to be better in the group with changes in $\Delta \text{Ct}$ value of $\text{EGFR}^{\text{m}} > 4.89$, suggesting that the changes of $\text{EGFR}^{\text{m}}$ may have a potential to predict disease progression. However, significant differences were not observed which may be due to the limited number of patients in this study. In the OS analysis, a statistically significant difference was found between the two groups, suggesting that quantitative detection of the changed value in ctDNA $\text{EGFR}^{\text{m}}$ before and after treatment may predict patient survival, although this may be affected by subsequent treatment. Using the above two analyses, we found the changes of mutations in plasma ctDNA, representing the degree of molecular responses, to be related to the degree of responses based on imaging. Patients in the MC group and in the group with changed value of $\Delta \text{Ct} > 4.89$ showed higher proportion of disease control (SD and PR) upon imaging assessment. Thus, compared to evaluation with imaging, the method with molecular responses has
significant superiority in evaluating the efficacy and prognosis in patients. This approach may help identify the patients with better PFS or OS after treatment, providing insights for clinicians to tailor treatment strategies for specific patients.

Furthermore, conventional staging of malignant tumors follows the tumor-node-metastasis (TNM) notation system. Recently, routine analysis of ctDNA from liquid biopsy has been considered to possibly make cancer diagnosis, treatment, and prognosis more accurate for individual patients. Thus, researchers put forward the concept of TNMB (TNM staging by liquid biopsy [B]) tumor classification, opening a new horizon for precision medicine. In our study, we demonstrated that the baseline status of ctDNA EGFRm has an impact on the survival of patients, confirming that it is valuable to propose the concept of TNMB staging for predicting clinical outcomes and prognosis. Additionally, we identified the clinical features of patients who could be more sensitive to drugs at an early stage, and based on the baseline ΔCt value, we could dynamically monitor the efficacy of TKIs. When changes in ΔCt value of ctDNA EGFRm did not reach cut-off level, we could recommend that clinicians change the medication or consider combination therapy. By using R-superARMS to detect the changes in EGFR mutations after 1 month treatment, we could predict the treatment response of patients. Then an NGS test can be further recommended for the patients with poor response, instead of performing an expensive and complex NGS testing at the time of initial diagnosis. At the same time, this method of measurement provides an opportunity to switch to NGS testing, rather than waiting for the disease progression. Based on the above considerations, ctDNA testing seems to provide us with a good and early detection method.

However, till date, we have rarely found a study like ours using quantitative methods to analyze EGFR changes. Although we obtained meaningful results, a limitation of this study was the small cohort of patients. Additionally, considering the effect of subsequent treatment after the progression of patients on the evaluation of OS, our study still has some shortcomings, and the relationship between changes in ctDNA and OS should be carefully interpreted. Nevertheless, our research as an initial exploration can provide some directions and ideas for larger, more rationally designed clinical studies in the future, including the effect of baseline EGFRm levels and different changes of ctDNA EGFRm values on patient outcomes, the impact of different drug treatments on the degree of molecular response, and the function of different molecular response intensities in predicting patient outcomes. We have proposed a semi-quantitative EGFR blood testing with R-superARMS to identify the people who respond to treatment more accurately and sensitively. As a supplement to the RECIST standard, R-superARMS has offered a good option to address the yet unmet clinical needs, such as early molecular responses, early detection of micrometastasis, prediction of tumor treatment efficacy and prognosis, etc. We look forward to conducting a large randomized controlled trial in the future to confirm the findings of our present study.

In conclusion, R-superARMS is likely to be a simple, fast, economic, and semi-quantitative PCR method, that is, claimed to be highly selective and sensitive for detecting EGFRm levels in plasma. Detecting ctDNA EGFRm using this method can identify patients who are more likely sensitive to targeted therapy, reflect the molecular load of patients, and predict the therapeutic efficacy and clinical outcomes of patients.

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Conflicts of interest

None.

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