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Liquid biopsy for therapy monitoring in early-stage non-small cell lung cancer

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Abstract

Liquid biopsy is now considered a valuable diagnostic tool for advanced metastatic non-small cell lung cancer (NSCLC). In NSCLC, circulating tumor DNA (ctDNA) analysis has been shown to increase the chances of identifying the presence of targetable mutations and has been adopted by many clinicians owing to its low risk. Serial monitoring of ctDNA may also help assess the treatment response or for monitoring relapse. As the presence of detectable plasma ctDNA post-surgery likely indicates residual tumor burden, studies have been performed to quantify plasma ctDNA to assess minimal residual disease (MRD) in early-stage resected NSCLC. Most data on utilizing liquid biopsy for monitoring MRD in early-stage NSCLC are from small-scale studies using ctDNA. Here, we review the recent research on liquid biopsy in NSCLC, not limited to ctDNA, and focus on novel methods such as micro RNAs (miRNA) and long non-coding (lncRNA).

Keywords: Liquid biopsy, Circulating nucleic acid, ctDNA, ctRNA, miRNA, lncRNA, Therapy monitoring, minimal residual disease (MRD), Early-stage non-small cell lung cancer (NSCLC)

Introduction

Liquid biopsy has been incorporated in clinical practice as a valuable diagnostic tool for advanced metastatic non-small cell lung cancer (NSCLC). In NSCLC, circulating tumor DNA (ctDNA) analysis is known to increase the chances of identifying targetable mutations and has been postulated clinically as a safer alternative than tissue biopsies.

Circulating cell-free DNA (cfDNA) in the peripheral blood originating from tumors, either in association with nucleosome (80–200 bp) or encapsulated in extracellular vesicles, is called ctDNA. ctDNA makes its way into the circulation during apoptosis and necrosis of the tumor cells [1]. The increased level of ctDNA in cancer patients results from limited infiltration of immune cells at the tumor site [2]. Due to intra-tumoral heterogeneity, detection of ctDNAs in liquid biopsies has some advantages over tumor tissue biopsies. ctDNA may offer a representative view of the genetic variations present in the tumor [3] and may provide a “real-time” assessment of the tumor burden, given the typical short half-life of cfDNA (between 16 min and 2.5 h) [4–6].

Typically, the ctDNA fractions in a total cfDNA bio-fluid samples are very low [7–9], which depends on cancer type and stage, requiring the necessity of highly sensitive detection technologies. Besides quantitative PCR (qPCR) and next-generation sequencing (NGS), recently developed techniques, such as digital PCR (dPCR), droplet digital PCR (ddPCR), and beads emulsion amplification magnets (BEAMing) are in use to analyze ctDNA [1]. Sequencing approaches include CAncer Personalized Profiling by deep sequencing (CAPP-Seq) [10], Tagged AMplicon deep sequencing (TAM-Seq) [11], Safe-sequencing (Safe-Seq) [12], and Duplex sequencing...
The noninvasive nature of obtaining ctDNA is an advantage in serial monitoring. Several studies have suggested the potential utility of ctDNA monitoring to detect the presence of minimal residual disease (MRD) post-resection in early-stage NSCLC [7, 15–18]. While there are emerging data on utilizing liquid biopsy to monitor MRD in early-stage NSCLC, most are small-scale studies using ctDNA. This paper reviews the current research on liquid biopsy in NSCLC, not limited to ctDNA, and focuses on novel methods such as micro RNA (miRNA) and long non-coding (lncRNA).

Main Text
Clinical use of ctDNA

ctDNA and targeted therapy

The most common scenario, where liquid biopsy utilizing ctDNA would be considered in the clinic, is in the context of newly diagnosed advanced metastatic NSCLC. The utilization of ctDNA has been reported to increase the chances of identifying targetable mutations such as EGFR mutations, ALK fusions, ROS1 fusions, BRAF V600E mutation, ERBB2 (HER2) mutations, RET fusions, MET amplification, and MET exon 14 skipping variants [19, 20].

Another common scenario where ctDNA testing could be utilized is perhaps at the time of disease progression in a patient on a tyrosine kinase inhibitor (TKI) treatment. ctDNA can help to identify the mechanism of resistance in such scenarios, and some secondary molecular alterations may be amenable to yet another TKI [21–25].

Indeed, the International Association for the Study of Lung Cancer (IASLC) suggests that liquid biopsy approaches have significant potential to improve patient care, and immediate implementation in the clinic is justified in several therapeutic settings relevant to NSCLC [26].

As the utility of ctDNA testing to guide targeted molecular therapy has been widely assessed through NSCLC patients with EGFR sensitizing mutations, we will use this as an example to reflect on how these tests were incorporated into clinical practice.

The ENSURE study showed that plasma EGFR mutations were useful in detecting patients who would benefit from erlotinib treatment [27]. Data from this study led to the subsequent US Food and Drug Administration (FDA) approval of the Cobas EGFR Mutation Test v2 (Roche Diagnostics, Indianapolis, IN) as the first liquid biopsy test and a companion diagnostic tool to guide treatment choices regarding EGFR TKIs. However, plasma EGFR mutations were detected only in 76.7% of the patients with EGFR mutations, confirmed from tumor tissues. Therefore, plasma-negative patients are recommended to be retested using tissue samples.

Subsequent data from the AURA phase II extension cohort [28] and AURA2 [29] studies led to the approval of the same method to detect the T790M mutation and determine the eligibility of osimertinib post-1st or-2nd generation EGFR TKIs. These studies reported a testing sensitivity of 61.4% using plasma, again suggesting the necessity of tissue testing when plasma-based testing is negative.

In addition to the Cobas EGFR Mutation Test v2, the Therascreen EGFR RGQ PCR Kit (Qiagen, Valencia, CA) has been approved by the European agency to detect EGFR mutations when tumor tissue is insufficient. The Qiagen method demonstrated reasonable mutation status concordance of 94.3% [sensitivity 65.7%, specificity 99.8%, positive predictive value (PPV) 98.6%, negative predictive value (NPV) 93.8%] among 652 matching tissue/cytology and plasma samples [30].

While osimertinib, a third-generation EGFR TKI, is now considered standard first-line therapy in patients with EGFR-mutated NSCLC [31], clinicians in many parts of the world may be bound by their respective health care system. They may still rely on the use of the first or second-generation EGFR TKIs upfront. Upon progression, it is critical to check for the most common resistance mutation on EGFR, T790M. If EGFR-T790M is detected through ctDNA testing, such patients would be eligible for the use of osimertinib as the second-line agent [28, 29]. However, if T790M is not detected through ctDNA, in that case, it is suggested that a tissue-based biopsy should be performed due to the possible false-negative results using a plasma-based test and also to rule out other resistance mechanisms, including small cell transformation [32].

Moreover, whether be it the first- or second-line therapy, patients inevitably progress on osimertinib. Acquired resistance, such as EGFR-C797S [21] and various additional alterations, including mutations (BRAF-V600 and KRAS) and amplification (HER2 and MET), have been reported [22, 23]. Detection of such alterations utilizing ctDNA may allow subsequent targeted therapy upon progression on osimertinib [24]. Similarly, plasma genotyping has also been used in NSCLC with other molecular alterations [25].

cTDNA for monitoring MRD after definitive therapy

Although patients with early-stage NSCLC are treated with curative intent, recurrence remains a challenge. Based on the Lung Adjuvant Cisplatin Evaluation (LACE) meta-analysis, the 5-year-survival of patients treated with adjuvant chemotherapy was only 5.4% with a hazard ratio (HR) of death at 0.89 (95% CI 0.82–0.96, p = 0.005) [33]. Consolidation immunotherapy with durvalumab after chemoradiation has shown improved overall survival (OS) and progression-free survival (PFS)
in unresectable stage III NSCLC. In exchange, over 30% of patients receiving durvalumab experienced at least one grade 3 or 4 adverse events (AEs) [34]. To truly optimize therapy in the curative setting, it is critical to identify those at high risk of recurrence and provide them with effective treatment.

Previous reports have demonstrated a rapid decline in ctDNA levels post-resection in early-stage lung cancer, which suggests the potential role of ctDNA as a measure of detecting recurrence, possibly before overt disease imaging. By identifying six somatic driver mutations of EGFR, KRAS, TP53, BRAF, PIK3CA, and ERBB2 through targeted sequencing utilizing the SV-CA50-ctDNA panel (San Valley Biotech Inc., Beijing, China), Guo et al. reported the changes in tumor and blood ctDNA levels in 41 NSCLC patients before and after surgery. Overall, 19 plasma samples (19/41; 46.3%) were positive for ctDNA before surgery, and among these 19 samples a total of 24 mutations were found including EGFR (17/24; 70.8%), KRAS (2/24; 8.3%), TP53 (3/24; 12.5%), BRAF (1/24; 4.2%), and PIK3CA (1/24; 4.2%). The average plasma ctDNA mutation frequency prior to surgery was 8.88% (range: 0.1–81.06%), whereas postsurgery it drastically reduced to 0.28% (range: 0.00–3.01%) [15]. A similar study by Chen et al. utilized a 50 gene-panel target sequencing to identify somatic mutations in matched tumor tissue DNA and plasma ctDNA samples. They showed that the average post-operation plasma ctDNA mutation frequency was reduced to 0.28% +/- 0.32%, from 1.21% pre-surgery (p < .0001) [16]. Detection of ctDNA post-surgical resection may guide clinicians to consider adjuvant therapy, whereas “non-detection” or reduced detection of ctDNA may be utilized to monitor disease recurrence.

If we know that ctDNA is reduced post-resection, could we then utilize it to monitor disease recurrence? How would we know that we are not detecting random DNA and are monitoring for primary disease recurrence?

Newman et al. developed the CAPP-Seq approach, which combines optimized library preparation methods for low DNA input with multiphase bioinformatics, to design a ‘selector’ composed of biotinylated DNA oligonucleotides that target recurrently mutated regions in cancers. The selector is applied to tumor DNA to identify the patient’s cancer-specific genetic aberrations and then directly to circulating DNA to quantify them over time. The authors give a few examples in NSCLC; a patient with stage IB disease who had surgery and another stage IB patient post-SBRT. In both patients, ctDNA was not detectable after definitive therapy, and both had long-term survival. In contrast, another patient who had chemoradiation for stage IIIB disease had a recurrence following an increase in ctDNA concentration post-therapy, although initial imaging showed a near-complete response [10].

Chaudhuri et al. also reported their use of CAPP-Seq to detect ctDNA. In a study of 40 patients treated with curative intent for stage I–III lung cancer, ctDNA was detectable in the first post-treatment blood sample in 94% of the patients experiencing recurrence [17].

In the TRACERx study, pre and post-surgical plasma ctDNA profiling was performed blinded to relapse status in a sub-group of 24 patients. In this study, multiplex-PCR assay panels were synthesized for each patient, targeting clonal and sub clonal single-nucleotide variants (SNVs) selected to track phylogenetic tumor branches in plasma. The multiplex-PCR NGS platform was validated to have a sensitivity of above 99% for detecting SNVs at frequencies above 0.1%, and the specificity of detecting a single SNV was 99.6%. At least two SNVs were detected in ctDNA from early-stage NSCLCs that had been analyzed in their discovery cohort data [35], suggesting biological sensitivity of a two-SNV threshold for ctDNA detection. Therefore, a threshold of two detected SNVs for calling a sample ctDNA-positive was selected. In this study’s validation cohort, at least two SNVs were seen in 13 out of 14 (93%) patients with confirmed NSCLC relapse before or during clinical relapse. In contrast, only 1 out of 10 (10%) patients, with no clinical evidence of NSCLC relapse, had at least two SNVs detected [18].

CAPP-Seq is an economical and ultrasensitive method for quantifying ctDNA. By providing personalized profiling through deep sequencing, the clinical application of CAPP-Seq to monitor and detect disease recurrence may become a vital surveillance tool in the future.

cDNA in treatment monitoring for metastatic NSCLC
cDNA testing may help monitor the therapeutic response in the metastatic setting, especially when there is a known driver, such as the sensitizing EGFR mutation. Several studies have evaluated the quantitative changes in serial ctDNA testing and have found a correlation between tumor response and survival.

One of the earlier studies in this area was reported by Mok et al., where ctDNA level was monitored in EGFR-mutated patients treated with gemcitabine followed by erlotinib or placebo. In erlotinib-treated patients, the objective response rate (ORR) of those with undetectable ctDNA at 12 weeks was more remarkable than those with detectable ctDNA levels (82.5% vs. 66.7%). Also, the lack of detectability of ctDNA at 12 weeks was associated with a significant increase in PFS and OS (16.6 vs. 7.8 months and 32.4 vs. 17.7 months, respectively) [36].

Similarly, Taus et al. reported 221 plasma ctDNA correlated samples in response to therapy in 33 EGFR-mutated NSCLC patients treated with TKI or chemotherapy. In 13 of 14 evaluable cases (93%), ctDNA
decrease predicted radiological response. Contrasting, in 17 of 19 evaluable cases (89%), ctDNA increase predicted radiological progression. In patients with circulating EGFR mutation becoming undetectable during follow-up had a significantly better PFS than those with detectable ctDNA, with a median of 295 vs. 55 days (HR 17.1; p < 0.001), respectively [37].

Monitoring the treatment effects of immunotherapy on NSCLC patients may be uniquely challenging due to an exacerbated inflammatory response that may mimic cancer progression (pseudo-progression). Goldberg et al. compared longitudinal changes in ctDNA levels of 28 NSCLC patients receiving immunotherapy. In this study, a ctDNA response correlated with superior PFS (HR 0.29; 95% CI, 0.09–0.89; p = 0.03) and OS (HR 0.17; 95% CI, 0.05–0.62; p = 0.007) [38]. Interestingly, Lee et al. evaluated the association between ctDNA and pseudo-progression in patients with metastatic melanoma treated with immunotherapy. In this study of 125 patients, the number of ctDNA copies was reduced by more than ten folds within 12 weeks of treatment in those who responded to therapy and also accurately identified all nine patients (7%) who had pseudo-progression based on imaging [39]. Future studies to evaluate the combination of ctDNA and imaging may further advance the field by measuring treatment response with greater accuracy.

**Tumor mutational burden (TMB) in treatment monitoring for metastatic NSCLC**

Another promising biomarker for patients receiving immunotherapy is TMB. Traditionally, TMB is measured using whole-exome sequencing (WES) or an NGS cancer gene panel from the tumor tissue. A high TMB correlates with response to immunotherapy in lung cancer [40, 41]. Gandara et al. reported that high blood TMB (bTMB), demonstrated by the FDA-approved FoundationOne (F1) CDx NGS assay [42], and was associated with increased PFS with atezolizumab over docetaxel in the second-line treatment for NSCLC. In patients with bTMB ≥16, a significant increase in PFS with atezolizumab (versus docetaxel) was shown [43]. The B-FIRST study was one of the first prospective trials to evaluate bTMB as a biomarker to predict the benefit of atezolizumab, used in the first-line setting for metastatic NSCLC. Utilizing the same cut-off values of bTMB ≥16, median PFS was 5.0 vs. 3.5 months, and median OS was 23.9 vs. 13.4 months in the bTMB ≥16 vs. < 16 patients, respectively [44]. Wang et al. recently showed the concordance of bTMB with matching tumor TMB calculated using WES. In their validation cohort, the investigators reported that a bTMB ≥6 was associated with superior PFS [45]. Georgiadis et al. developed a novel liquid biopsy method utilizing a hybrid-capture-based 98-kb pan-cancer gene panel with multifactorial error correction and peak-finding algorithm. This approach identified TMB and microsatellite instability (MSI) from plasma cfDNA with a specificity of > 99% (N = 163), and sensitivities of 67% (N = 15) and 78% (N = 23), respectively [46]. While testing for TMB and MSI using liquid biopsy appears promising for real-time evaluation and global measurement of a tumor, testing methods need further optimization and validation of cut-off values to be clinically meaningful.

**What is circulating non-coding RNA (ncRNA)?**

Circulating genomic materials in the bloodstream encompass nucleotide fragments of DNA and RNA molecules [1]. The RNAs that originate from tumor cells are called circulating tumor RNAs (ctRNAs) [47], which includes both coding and non-coding RNAs (ncRNAs). The ncRNAs comprise a large group of RNAs without the translational ability, involving micro RNAs (miRNAs), transfer RNAs (tRNAs), long non-coding RNAs (lncRNAs), circular RNAs (circRNAs), small nuclear RNAs (snRNAs), small nucleolar RNAs ( snoRNAs), and piwi interacting RNAs (piRNAs) [48, 49].

**Potential role of ncRNA utilization in monitoring early-stage NSCLC**

The detection of ncRNA is a noninvasive, innovative approach for diagnosis [1]. As potential biomarkers, ncRNAs were assessed for their ability to provide qualitative and quantitative information (e.g., expression levels), detect the mutational pattern in the transcript, and identify cancer-associated fusion transcripts or alternative splice variants [48]. Among ncRNAs, miRNAs, lncRNAs, and recently characterized circRNAs are more suitable as biomarkers [48]. Cutting-edge technologies, such as ddPCR and comprehensive characterization of RNA using RNASeq, dramatically enhance the probability of ncRNAs as biomarkers. These are capable of detecting new transcripts and can precisely quantify ncRNAs with higher sensitivity [48].

It has been shown that crizotinib-associated clinical outcomes correlated with EML4-ALK fusion transcripts in NSCLC [50]. Nilsson et al. demonstrated that ctRNA extracted from platelets can detect EML4-ALK rearrangements with 65% sensitivity and 100% specificity [50]. In an index patient, EML4-ALK fusion transcripts reappeared as ctRNA 2 months before radiographic confirmation of crizotinib resistance. Such detection of fusion RNA transcript in the circulation can help monitor and predict targeted therapies’ outcomes [50].

**miRNAs as biomarkers**

In 1993, after the discovery of miRNA in Caenorhabditis elegans as an LIN-4 gene transcript [51, 52], the miRNA field advanced immensely with an in-depth understanding...
of its role in human pathobiology [53]. miRNAs are classified as the main category of small ncRNAs. miRNAs consist of 18–25 nucleotides (nts) [54]. miRNAs represent about 3% of the human genome with 700 members, responsible for the regulation of 20–30% of coding genes [55, 56]. Circulating miRNAs have been investigated as biomarker candidates in various types of cancers [57]. The origin of cancer-associated circulating miRNAs is unclear [58]. They may be secreted by circulating tumor cells, tumor cells from the primary site or by cells that have metastasized. The infiltrating immune cells at the tumor microenvironment may also be considered as a source. However, irrespective of their origin, miRNA can be useful biomarkers for tumor progression [59, 60]. Both radio and chemotherapy can influence the circulating miRNA as evident from several studies [61]. Many patients with early-stage NSCLC require adjuvant chemotherapy. miRNAs may serve as a biomarker to assess such therapy’s sensitivity quite early, which playing an important role in patient management decisions and reducing unnecessary therapeutic toxicities [57]. Table 1 summarizes various ctRNAs assessed as a biomarker in NSCLC.

The levels of miR-1246 and miR-1290 in serum were associated with the initiation, progression, and metastasis of NSCLC [62]. miRNA levels correlated with the tumor stage and clinical response and increased by multiple folds compared to healthy controls, suggesting the utility of miR-1246 and miR-1290 as clinical biomarkers [62]. Mutational changes in miRNAs may also predict the outcome of malignancies. Single nucleotide polymorphisms (SNPs) at let-7 complementary sites (LCS) are associated with an increased risk of NSCLC. In a study, 74 NSCLC tissue samples were sequenced to detect mutations at let-7 complementary sites (LCS) in the KRAS 3′ untranslated region. Specific LCS6 allelic variant is speculated to be a potential risk factor for NSCLC [63]. Detection of mutated miRNAs in the body fluids of NSCLC patients can be a noninvasive approach to determine the patients at risk.

Hu et al. investigated serum miRNA levels in long-term (mean survival 49.54 months) and short-term (mean survival 9.54 months) survivors of stage I-IIIA NSCLC [64]. After validation of differentially expressed serum miRNAs in 243 patients, the study suggested a panel of miRNAs, involving miR-486, miR-30d, miR-1, and miR-499, that were linked to survival [64].

In an NSCLC study, patients treated with cisplatin and/or other chemotherapeutic regimens had increased circulating miR-125b. Elevated serum level of miR-125b was indeed associated with inferior OS (P = 0.0012) [65]. Upregulation of miR-125b has been reported to inhibit apoptosis caused by cisplatin and associated with cisplatin resistance in breast and ovarian cancers [66, 67]. The level of miR-125b, therefore, may help in deciding platinum-based chemotherapy response in NSCLC patients. Another study identified a set of 14 circulating miRNA including miR-134, miR-200b, miR-574, miR-858, and let-7c to predict grade 3+ radiation-induced cardiac toxicity (RICT) in NSCLC [68].

Among other miRNAs, the diagnostic and prognostic potential of serum miR-141 was demonstrated in a prospective cohort study involving 108 NSCLC patients and 54 matched healthy controls [69]. Receiver operating characteristic (ROC) curve analysis showed significant elevation of miR-141 expression in NSCLC patients (P < 0.001). Moreover, the expression of miR-141 could distinguish both lymphatic (P = 0.015) and distant (P = 0.025) metastasis. Subtype analysis revealed an association between miR-141 expression and OS of lung adenocarcinoma but not squamous cell carcinoma [69]. Using multivariate Cox regression analyses, Xu et al. assessed elevated plasma levels of miR-18a, miR-20a, and miR-92a as prognostic biomarkers for NSCLC [70]. In an approach to combine miRNAs for the diagnosis of NSCLC, Abdollahi et al. suggested a miR-panel (miR-21, miR-148, miR-152 and miR-638) of 96.4% sensitivity and 86.67% specificity [71]. Recently, exosomal plasma miRNA of NSCLC patients namely miR-23b-3p, miR-10b-5p and miR-21-5p significantly correlated with poor OS (hazard ratio between 2.12–2.42). Several miRNAs in sputum, including miR-21, miR-143, miR-155, miR-210, and miR-372, have also been reported to detect NSCLC earlier [72, 73].

Long non-coding RNAs (lncRNAs) as biomarkers

Aside from miRNAs, another encouraging source of ncRNA-based biomarkers is the lncRNA. lncRNAs are composed of 200 nts or more without any open reading frame but contribute to various cellular activities and diseases, including cancer [74]. There are approximately 60,000 ncRNAs in the human genome; > 70% are lncRNAs [74]. Only ~ 20% of lncRNAs are annotated and shown to regulate diverse cellular functions [75]. The regulation of gene expression by lncRNAs occurs through cis and trans mechanisms [76, 77]. Additionally, lncRNAs can regulate mRNA splicing and subcellular localization of proteins [78]. Therefore, it is not surprising that abnormal lncRNA expression contributes to cancer development and progression [79].

Recent research on lncRNA has revealed several potential biomarkers for the detection and prognosis of NSCLC. lncRNAs XIST and HIF1A-AS1 were significantly elevated in serum samples of patients with NSCLC [80]. Notably, these were significantly reduced after surgery. Patients with NSCLC and the control group were strongly separated in the ROC curve analysis (area under the curve (AUC) of XIST and HIF1A-AS1 were 0.834 and 0.876, respectively). The combination of XIST
| ctRNAs                   | Expression | Function            | Bio-fluids          | Patients types | References (PMID)                          | Notes                                      |
|-------------------------|------------|---------------------|---------------------|----------------|--------------------------------------------|--------------------------------------------|
| lncRNA MALAT1           | Down       | Diagnostic          | Whole blood         | NSCLC          | 26,137,228; 24,313,945                     | Patients vs healthy controls               |
| lncRNA NR-026689        | Up         | Diagnostic          | Whole blood         | NSCLC          | 26,908,441                                 | Patients vs healthy controls               |
| lncRNA XIST             | Up         | Diagnostic          | Serum               | NSCLC          | 26,339,353                                 | Patients vs healthy controls               |
| lncRNA HIF1A-AS1        | Up         | Diagnostic          | Serum               | NSCLC          | 26,339,353                                 | Patients vs healthy controls               |
| lncRNA SOX2OT           |            |                     | Serum               | NSCLC          | 29,504,701                                 | Patients vs healthy controls               |
| lncRNA ANRIL            |            |                     | Serum               | NSCLC          | 29,504,701                                 | Patients vs healthy controls               |
| lncRNA16 (ENST00000539303) | Up       | Diagnostic          | Plasma              | NSCLC          | 27,999,202                                 | Patients vs healthy controls               |
| lncRNA GAS5             | Down       | Diagnostic          | Plasma              | NSCLC          | 27,631,209                                 | Patients vs healthy controls               |
| lncRNA UCA1             | Up         | Diagnostic          | Plasma              | NSCLC          | 26,380,024                                 | Patients vs healthy controls               |
| lncRNA SPRY4-IT1        | Up         | Diagnostic          | Plasma              | NSCLC          | 26,453,113                                 | Patients vs healthy controls               |
| lncRNA ANRIL            | Up         | Diagnostic          | Plasma              | NSCLC          | 26,453,113                                 | Patients vs healthy controls               |
| lncRNA NEAT1            | Up         | Diagnostic          | Plasma              | NSCLC          | 26,453,113                                 | Patients vs healthy controls               |
| lncRNA RP11-397D12.4    | Up         | Prognostic          | Plasma              | NSCLC          | 26,393,913                                 | Patients vs healthy controls               |
| lncRNA AC007403.1       | Up         | Prognostic          | Plasma              | NSCLC          | 26,393,913                                 | Patients vs healthy controls               |
| lncRNA ERICH1-AS1       | Up         | Prognostic          | Plasma              | NSCLC          | 26,393,913                                 | Patients vs healthy controls               |
| lncRNA AFAP1-AS1        |            | Diagnostic          | Serum               | NSCLC          | 29,080,690                                 | EGFR-TKI therapy                           |
| lncRNA DLX6-AS1         |            | Diagnostic          | Plasma              | NSCLC          | 31,612,030                                 | EGFR-TKI therapy                           |
| lncRNA NR-026689        | Up         | Diagnostic          | Cell lines (human)  | NSCLC          | 26,908,441                                 | EGFR-TKI therapy monitoring               |
| lncRNA SCARNA7          |            | Diagnostic          | Pleural effusions   | NSCLC          | 31,691,525                                 | EGFR-TKI therapy                           |
| lncRNA MALAT1           |            | Diagnostic          | Pleural effusions   | NSCLC          | 31,691,525                                 | EGFR-TKI therapy                           |
| ctRNAs | Expression | Function | Bio-fluids | Patients types | References (PMID) | Notes |
|--------|------------|----------|------------|----------------|------------------|-------|
| IncRNA NONHSAT017369 | | Diagnostic or prognostic | Pleural effusions | NSCLC | 31,691,525 | EGFR-TKI therapy monitoring |
| IncRNA HNF1A-AS1 | Up | Lung cancer including NSCLC | 25,863,539 |
| IncRNA CRNDE | | Lung cancer including NSCLC | 28,940,804; 30,554,121; 30,230,527; 31,275,444 |
| Combined IncRNA GASS and IncRNA SOX2OT | | Diagnostic and prognostic | Plasma | NSCLC | 31,077,615 |
| IncRNA SOX2OT, ANRIL with tumor markers CEA, CYFRA21-1 and SCCA | | Diagnostic | Serum | NSCLC | 29,504,701 |

**micro RNA (miRNA /miR)**

| miR-21 | Up | Lung cancer including NSCLC | 18,719,201 |
|        |    |                            |            |
|        |    | combination of miR-145 and miR-451 | Diagnostic or serum | Lung cancer including NSCLC | 28,060,535; 23,301,032 |
| let-7 | SNP | Diagnostic | NSCLC | 18,922,928 |
| miR-142-3p | Up | Prognostic | Serum | NSCLC | 22,983,388; 23,794,259 | Cisplatin and pemetrexed-based chemotherapy |
| miR-125 | Up | Prognostic | Serum | NSCLC | 22,983,388; 23,794,259 | Cisplatin and pemetrexed-based chemotherapy |
| miR-22 | Up | Prognostic | Serum | NSCLC | 22,983,388; 23,794,259 | Cisplatin and pemetrexed-based chemotherapy |
| miR-1290 | Prognostic | Serum | Lung cancer including NSCLC | 27,325,363 | Metastasis |
| miR-1246 | Prognostic | Serum | Lung cancer including NSCLC | 27,325,363 | Metastasis |
| miR-590-5p | Prognostic | Plasma | NSCLC | 31,520,555 |
| miR-141 | | Diagnostic and prognostic | Serum | NSCLC | 31,938,373 |
| miR-18a | Up | Prognostic | Plasma | NSCLC | 29,266,846 |
| miR-20a | Up | Prognostic | Plasma | NSCLC | 29,266,846 |
| miR-92a | Up | Prognostic | Plasma | NSCLC | 29,266,846 |
| miR-21, miR-148, miR-152 and miR-638 | Diagnostic | NSCLC | 31,236,600 |
| miR-375 | Diagnostic and prognostic | Serum | NSCLC (Indian patients cohort) | 32,007,320 |
| miR-320a | Diagnostic and prognostic | Serum | NSCLC (Indian patients cohort) | 32,007,320 |
and HIF1A-AS1 significantly improved the AUC value (0.931) as a predictive biomarker for NSCLC [80]. An increase in exosomal (plasma) lncRNA Distal-less homeobox six antisense RNA 1 (DLX6-AS1) levels were also determined as a potentially promising diagnostic biomarker. The study analyzed 72 serum samples of NSCLC patients and 64 healthy donors and observed an association with advanced disease, lymph node metastasis and poor tumor differentiation. The sensitivity and specificity were 77.5 and 85.9% respectively. Notably, the expression of DLX6-AS1 in the serum decreased after surgical resection [81].

Recently, an association between lncRNAs and EGFR mutation was demonstrated. Using the Clarion D Human chip technology, in the plasma samples collected from pleural effusions, three lncRNAs (SCARNA7, MALAT1, and NONHSAT017369) were found to be significantly associated with EGFR mutation and thus could be considered as biomarkers of therapy monitoring [82]. These biomarkers could help predict EGFR mutation status and monitor EGFR-TKI efficacy in patients with EGFR-positive NSCLC.

It is evident from studies that the combination of biomarkers significantly increases the diagnostic and prognostic sensitivity and specificity. Plasma levels of GAS5 and SOX20T lncRNAs together showed superior sensitivity (83.8%) and specificity (81.4%) for the diagnosis and prognosis of NSCLC [64]. Circulating GAS5 in combination with other biomarkers can be helpful in monitoring after surgical treatment for NSCLC patients [83]. In a cohort study, serum lncRNA biomarkers SOX20T and ANRIL, along with tumor markers CEA, CYFRA21-1, and SCCA, demonstrated higher sensitivity (77.1%) and specificity (79.2%) for NSCLC diagnosis [84].

### Table 1 ctRNAs as a biomarkers in NSCLC (Continued)

| ctRNAs                        | Expression | Function          | Bio-fluids | Patients types | References (PMID) |
|-------------------------------|------------|-------------------|------------|----------------|-------------------|
| miR-21                        |            | Diagnostic or prognostic | Sputum     | NSCLC          | 30,165,346        |
| miR-143                       |            | Diagnostic or prognostic | Sputum     | NSCLC          | 30,165,346        |
| miR-155                       |            | Diagnostic or prognostic | Sputum     | NSCLC          | 30,165,346        |
| miR-210                       |            | Diagnostic or prognostic | Sputum     | NSCLC          | 30,165,346        |
| miR-372                       |            | Diagnostic or prognostic | Sputum     | NSCLC          | 30,165,346        |
| circular RNA (circRNA)        |            | Prognostic         | Plasma     | NSCLC          | 29,588,350        |
| circ-0005962 (up), circ-0003958 (up), circ-0086414 (down), circ-0001936 (down) |            |                   |             |                 |                   |
| circFARSA                     |            |                   | Plasma     | NSCLC          | 29,722,168        |
| others                        |            |                   |            |                |                   |
| mRNA CCL5                     |            | Prognostic         | NSCLC      |                | 31,215,484        |
| mRNA CLU                      |            | Prognostic         | NSCLC      |                | 31,215,484        |
| mRNA SPARC                   |            | Prognostic         | NSCLC      |                | 31,215,484        |
| mRNA SRGN                     |            | Prognostic         | NSCLC      |                | 31,215,484        |
| mRNA with KRAS mutation       |            | Diagnostic and prognostic | Lung cancer including NSCLC |                | 12,459,728 |
| snoRD33/66/76                 |            | Diagnostic         | Plasma     | NSCLC          | 20,663,213        |
| snoRDs-66                    |            |                   | Sputum     | NSCLC          | 27,176,474        |
| snoRDs-78                    |            |                   | Sputum     | NSCLC          | 27,176,474        |
| snRNA U2 (RNU2–1)             |            | Diagnostic         | Lung cancer including NSCLC |                | 23,527,303        |

Nagasaka et al. Molecular Cancer (2021) 20:82
circRNAs as biomarkers

circRNAs were described first in the early 1970s as a non-functional by-product of RNA splicing [85]. They are cell lineage-specific, enriched in exosomes, and observed in different body fluids [86]. Due to its enhanced stability compared to other ncRNAs, they are more suitable as biomarkers for cancer diagnosis, monitoring development, and progression [87]. circRNAs are a large unique family of ncRNAs that have joined covalent ends. They intricate in the complex regulation of normal processes via interacting with proteins, nucleic acids, and other molecules [88–90]. Recently, circFARSA has been detected and confirmed in the plasma of patients with NSCLC, with high confidence ($p < 0.001$) [91], which warrants future studies. The average plasma half-life of circRNAs is more than 48 h, which is much longer than mRNAs (~10 h) [92]. The hidden 3’ ends of circRNAs protect them against nuclease-mediated degradations compared to other ncRNAs [90, 93]. It is assumed that circRNAs overexpressed in lung cancer cells have the potential to be released in the circulation. Such notion has been validated, and indeed a differential expression of circRNAs has been observed in the plasma of early and late-stage NSCLC patients [94]. The study further identified a panel of circRNAs with prognostic significance in NSCLC, which includes two overexpressed (circ-0005962 and circ-0003958) and underexpressed (has-circ-0086414 and has-circ-0001936) circRNAs [94].

snRNA/snoRNAs play essential roles in the early steps of gene expression [95]. Recently, a panel of two snoRNAs, including snoRD-66 and snoRD-78, have been identified in sputum samples of patients with NSCLC. The panel detected NSCLC with 74.6% sensitivity and 83.6% specificity, which increased further to 89% sensitivity and 89% specificity when combined with miRNAs (miR-21, miR-31, and miR-210) [96].

Potential roles and challenges of ctDNA analysis

While multiple studies have proven the utility of ctDNA MRD measurement in post-therapy NSCLC patients and may allow personalized MRD-based adjuvant treatment options, they are still challenged by many limitations.

Concordance between the tissue and liquid

When utilizing ctDNA to detect MRD, it is vital to confirm the concordance rates between plasma samples and the original tumor. A systemic review and meta-analysis of 32 studies with 4527 advanced NSCLC patients reported a pooled sensitivity and specificity of 0.70 and 0.98 respectively for ctDNA EGFR mutation status in plasma samples compared to tumor tissues [97]. The discordance between tumor tissue and plasma DNA likely arises from the limitations of ctDNA detection.

Detection limitation

Data from the TRACERx study demonstrated that tumor volume correlates with the mean plasma VAF of clonal SNVs in ctDNA-positive NSCLC. A primary NSCLC tumor volume of 10 cm$^3$ predicts a ctDNA plasma VAF of 0.1%. The tumor size may constrain the sensitivity of clonal-SNV ctDNA. It may not also be helpful for the identification of very early disease recurrence or as an initial approach to screen early-stage NSCLC. Lung nodules smaller than computed tomography resolution could release ctDNA with allele frequencies as low as 0.00018%. This would challenge even the most efficient ctDNA technologies, which have reported detection thresholds of 0.00025% under optimal conditions. While ctDNA-based methods would need improvements in sensitivity thresholds for identifying microscopic diseases, there might still be a challenge of missing very minimal residual disease. Practically, the combination of such technologies with clinical correlation would be critical in evaluating patients.

False positives

The specificity of ctDNA testing could also be a confronting issue, as not all mutations detected in plasma are related to a cancerous process. For example, BRAF V600E mutation, which is known as the driver mutation in melanoma and NSCLC, is found more frequently in tissue specimens of benign nevi [98]. Indeed, Di Giorgi et al. reported a case where BRAF V600E mutated DNA was detected in both plasma and formalin-fixed tissue specimens from a man with benign nevi [99].

Furthermore, cfDNA primarily originates from hematopoietic cells, and evidence of somatic mutations in these cells are common with aging (10% in healthy adults > 70 years), defined as clonal hematopoiesis of indeterminate potential (CHIP) [100]. The presence of CHIP does not necessarily imply the presence of cancer and can confound ctDNA testing results. Hu et al. reported that most JAK2 mutations, some TP53, and rare KRAS mutations detected in cfDNAs were from clonal hematopoiesis and not tumors. They suggested using paired peripheral blood cells genotyping so that such clonal hematopoiesis-derived mutations would not be misdiagnosed as an occult malignancy [101]. Li et al. reported a novel approach using cfDNA and matching white blood cells with a hybrid capture panel, covering 37 lung cancer-related gene sequencing up to 50,000x raw target coverage, filtering somatic mutations attributable to clonal hematopoiesis. Plasma NGS sensitivity for known oncogenic drivers was 75% ($N = 68/91$), and the specificity was 100% ($N = 19/19$) [102].

In contrast, germline mutations can also complicate ctDNA testing as their mere presence does not necessarily imply a cancerous process. However, a high allele
fraction (typically between 30 and 70%) is suspicious for a germline variant than a somatic hit [103].

On a different note, caution must be taken when performing liquid biopsy testing in pregnant women. The fact that fetal cfDNA are shed into the maternal bloodstream [104] may obscure the interpretation.

The optimal timing of ctDNA testing also needs to be further evaluated in prospective studies. While Chen et al. suggested utilizing ctDNA detection on the third day after R0 resection as the post-operative baseline value [105], Szpechcinski et al. reported a drastic increase in plasma DNA levels a week after primary tumor resection, likely due to the surgical trauma [106]. In the metastatic setting, peak concentration of ctDNA early after the start of therapy was reported [107], which is also likely related to the rapid release of tumor DNA by treatment-induced tumor cell death. Distinguishing these and actual residual/resistant disease must be examined further.

In addition to the timing, the definition of a “positive” MRD, as well as standardized criteria to define “response” and “progression” or “relapse” during longitudinal monitoring, are critical to minimize variations among studies and provide reliable practices.

**Challenges and the prospect of ncRNAs as biomarkers**

The unstable nature and very short half-life of naked ncRNAs in plasma compared to ctDNA is a challenge for ncRNAs to be considered as biomarkers [108]. The stability of ctRNA is also dependent on the complexes they form. For example, miRNA in microvesicles are resistant to RNase-mediated degradation compared to protein-bound miRNA [109]. It is indeed difficult for an individual ctRNA to qualify as a candidate biomarker because of its overlapping levels in patients and healthy controls, limiting its sensitivity and specificity [110]. However, there are some advantages of ncRNAs over ctDNAs. The ctDNAs levels in the circulation are very low (between 0.01–1% of total ctDNA), whereas, ncRNAs are far more abundant than ctDNAs. Dysregulation of ncRNAs in different biofluids is frequent, even in the earliest stages of cancer [4, 111]. A combination of various ncRNAs and other established circulation nucleic acids should be considered to enhance ncRNAs’ potential as a non-invasive biomarker.

**Cost**

Collaborators of Ontario Health conducted a systematic review. They determined that utilizing liquid biopsy as a triage test before performing a tissue biopsy was more cost-effective in the setting of detecting EGFR T790M resistance mutation post 1st or 2nd generation EGFR TKIs, as the estimated cost was $700 for a liquid biopsy vs. $2500 for a tissue biopsy [112]. Furthermore, upfront NGS testing in patients with metastatic NSCLC can have substantial cost reductions compared to single-gene testing for both Centers for Medicare & Medicaid Services and US commercial payers [113]. The cost-effectiveness of molecular testing in early-stage NSCLC needs further evaluation. It is estimated that the cost of a personalized ctDNA profiling per patient for sequencing of a single tumor region, synthesis of a patient-specific assay panel, and profiling of five plasma samples is US $1750 [17].

**Future directions**

Detecting and targeting ctDNA, particularly those with sensitizing molecular mutations, such as EGFR, would be especially beneficial in NSCLC with the availability of specific targeted inhibitors. In the recently reported ADAURA study, adjuvant osimertinib demonstrated a 79% reduction in the disease recurrence risk or death in resected stage IB-IIIA EGFR-mutated NSCLC patients (DFS HR 0.21, 95% CI 0.16–0.28, p < 0.0001) [114]. In addition to erlotinib, the ongoing ALCHEMIST study is evaluating the utility of crizotinib (ALK inhibitor) and nivolumab (PD1 inhibitor) in completely resected NSCLC post standard care adjuvant therapy. Many other studies are assessing the role of chemotherapy with or without additional treatments, such as immunotherapy in the adjuvant setting. Table 2 shows the ongoing studies utilizing ctDNA monitoring in early-stage NSCLC.

While detectable plasma ctDNA post-surgery appears promising to utilize as a biomarker for MRD, the interpretation of a negative ctDNA test could be a challenge; as there are limitations to the detection sensitivity and there always will remain a chance of false-negative testing. To minimize the deleterious consequences of diagnosing a patient with a false-negative ctDNA result as “cancer-free”, further long-term prospective studies utilizing serial plasma ctDNA monitoring post-resection with endpoints in OS would be of benefit.

At this time, the most robust evidence in liquid biopsies is reported from NSCLC studies, but recently, the role of liquid biopsies are being expanded into multiple tumor types.

Although the National Comprehensive Cancer Network (NCCN) guidelines recommend RAS, BRAF, and MSI or mismatch repair deficiency (dMMR) testing for all patients with metastatic colon cancer [115], a retrospective study showed that only 40% of these patients received guideline-recommended testing [116]. Direct comparison of tumor tissue-based approach vs. plasma-based ctDNA testing has shown concordance in detecting APC, TP53, KRAS, NRAS, and BRAF mutations in treating naïve and non-anti-EGFR-treated cohorts of colorectal cancer [117]. With limited tissue availability in some patients, utilizing ctDNA for detecting these markers could increase the number of patients being
tested and increase the chance of receiving appropriate targeted therapies. Besides, ctDNA could be used to predict resistance to anti-EGFR therapy. Diaz et al. found that 40% of the patients with metastatic colon cancer, who were initially KRAS wild-type, developed detectable KRAS mutations in their serum using ctDNA after 5–6 months of anti-EGFR therapy with panitumumab [118].

In stage III of colon cancer, Tie et al. reported a significant difference in the 3-year recurrence-free interval in patients with detectable vs. undetectable levels of ctDNA after surgery (47% vs. 76%, HR 3.8; \( p < 0.001 \)) and after completion of chemotherapy (30% vs. 77%, HR 0.75; \( p < 0.001 \)) [119]. This suggests that post-surgical and post-chemotherapy ctDNA analyses may identify patients with stage III colon cancer with a high risk of recurrence. Prospective randomized trials (NCT04068103, NCT03803553) are underway to answer these critical questions.

In resectable pancreatic cancer, Groot et al. monitored ctDNA in 59 patients using ddPCR to detect the most commonly seen somatic KRAS mutations (G12D, G12V, G12R, and Q61H) [120]. ctDNA was detected in about 50% of the patients before resection, and this was a strong predictor for worse PFS and OS. Those who received neoadjuvant chemotherapy were less likely to have detectable preoperative ctDNA (21% vs. 70%; \( p < 0.001 \)). Detecting ctDNA post-resection was associated with worse OS.

Late recurrence may also be detected by ctDNA in specific settings. A secondary analysis with ctDNA was done in hormone receptor-positive HER2-negative operable breast cancer patients from a randomized clinical study. Presence of ctDNA approximately 5 years after the initial diagnosis positively correlated with late disease recurrence [121].

Interestingly, there is data to support the use of ctDNA testing even in patients with cancers of unknown origin (CUP). Kato et al. reported the genomic landscape of ctDNA testing in 442 patients with CUP. Overall, 80% of patients (\( N = 353/442 \)) had detectable ctDNA and

### Table 2: Ongoing studies utilizing ctDNA monitoring in early-stage NSCLC

| NCT#        | Type          | Scope                                                                 | Cohorts/Arms                                                                 | Primary outcome measures                          | Number to be enrolled |
|-------------|---------------|----------------------------------------------------------------------|----------------------------------------------------------------------------|---------------------------------------------------|-----------------------|
| NCT04367311 | Phase 2       | Adjuvant chemo + atezolizumab in resected NSCLC and clearance of ctDNA | Non-squam: cisplatin, pemetrexed, atezo Squam: cis, docetaxel, atezo       | Percentage of pts. with undetectable ctDNA post adjuvant treatment | 100                   |
| NCT04267237 | Phase 2       | Atezolizumab +/- RO7198457 (personalized cancer vaccine) following adjuvant chemo in ctDNA positive pts. post resection | Atezo, atezo + RO7198457                                                  | DFS                                              | 332                   |
| NCT01629498 | Phase 1/2     | Intensity modulated photon (IMRT) or proton radiation in stage II-IIIB NSCLC | IMRT or proton radiation therapy                                         | MTD of IMRT and proton therapy                    | 100                   |
| NCT03521154 | Phase 3       | Osimertinib following chemoradiation in stage III NSCLC (LAURA)      | Osimertinib, placebo                                                      | PFS                                              | 200                   |
| NCT04385368 | Phase 3       | Durvalumab with chemo in resected stage II-III NSCLC (MERMAID-1)     | Chemotherapy + durvalumab/placebo                                        | DFS in MRO positive analysis set                  | 590                   |
| NCT03774758 | Observational | Biomarkers for risk stratification in lung cancer                     | Benign nodule on screening, incidental benign nodule, presumed lung cancer, suspicious nodule, suspicious incidental nodule, post-treatment lung cancer | Sensitivity and specificity of ct-DNA LUNAR assay, |
| NCT0355350  | Observational | Role of ctDNA from liquid biopsy in resected lung tumor (LIBERTI)     | Completely resected early stage NSCLC                                     | Correlation between ctDNA after surgery with DFS  | 500                   |
| NCT0351732  | Observational | ctDNA exposure in peripheral blood                                    | Healthy volunteer, those with stage 0-IV cancer (i.e. lung, colon, pancreatic) | ctDNA exposure in peripheral blood using a novel process, a feasibility study | 10,000                 |
| NCT04354064 | Observational | ctDNA for early treatment response assessment of solid tumors         | Healthy volunteer, those with stage I-IV solid tumors (i.e. lung, breast, colon) | Freedom from progression                         | 3362                  |
| NCT03838588 | Observational | Tracking molecular evolution for NSCLC (T-MENC)                       | Monitoring of ctDNA in stage IB, II, III NSCLC post resection            | Clonal evolution assessed with liquid biopsy, concordance of ctDNA with PFS and OS | 200                   |

Nagasaka et al. Molecular Cancer (2021) 20:82
66% (N = 290/442) had at least one characterized alteration, and most (N = 289/290) had an alteration that would be hypothetically targetable [122]. Combining different ctRNAs is one strategy to enhance the sensitivity and specificity as a biomarker for assessing disease detection and progression with increased precision. Simultaneous detection of two mRNAs and a lncRNA in serum exosomes was shown to have more excellent diagnostic value in colorectal cancer [123]. The lack of consensus on an optimal method [108] needs to be overcome for less variable recovery rates for the processing of samples. Notably, the advent of a lab-on-a-chip microdevice technology may allow rapid processing of samples [124]. The microfluidic system called integrated comprehensive droplet digital detection (IC3D) can detect a very low level of mRNAs in plasma within few hours [125]. With the promising potential as a biomarker in different cancer progression stages, the recently identified ctRNAs warrant further validation studies.

Conclusions

Liquid biopsy approaches utilizing ctDNA, miRNA, and lncRNA have a significant potential to improve patient care. Further prospective studies are warranted to optimize their use in early-stage NSCLC.

Abbreviations

AE: Adverse events; ALK: Anaplastic lymphoma kinase; APC: Adenomatous polyposis coli; AUC: Area under the curve; BEAMing: Beads emulsion amplification magnets; Braf: V-Raf murine sarcoma viral oncogene homolog B1; CAPP-Seq: Cancer personalized profiling by deep sequencing; CI: Confidence interval; ctDNA: Cell-free DNA; circRNA: Circular RNA; ctRNA: Circulating tumor RNA; ddPCR: Droplet digital PCR; dNMR: Mismatch repair deficiency; dPCR: Digital PCR; EGFR: Epidermal growth factor receptor; BRBB2: Enzymeblastic oncogene B; FDA: Food and drug administration; HER2: Human epidermal growth factor receptor 2; HR: Hazard ratio; iASLC: International association for the study of lung cancer; IDES: Integrated digital error suppression; KRAS: Kirsten rat sarcoma viral oncogene homolog; LCS: Let-7 complementary sites; LncRNA: Long non-coding RNA; MET: Mesenchymal epithelial transition factor; miRNA: Micro RNA; MDD: Minimal residual disease; MSI: Microsatellite instability; ncRNA: Non-coding RNA; NCCN: National comprehensive cancer network; NGS: Next generation sequencing; NTRAS: Neuroblastoma ras viral oncogene homolog; NSCLC: Non-small cell lung cancer; ORR: Objective response rate; OS: Overall survival; Pik3Ca: Phosphatidylinositol-4,5- bisphosphate 3-kinase catalytic subunit alpha; pIRNA: PAXI interacting RNA; PFS: Progression-free survival; pPCR: Quantitative PCR; PET: Rearranged during transfection; ROC: Receiver operating characteristics; Safe-Seq: Safe-sequencing; SNP: Single nucleotide polymorphisms; snRNA: Small nuclear RNA; snORNA: Small nuclear RNA; TAM-Seq: Tagged amplicon deep sequencing; Tki: Tyrosine kinase inhibitor; TMB: Tumor mutational burden; TP53: Tumor protein 53; tRNA: Transfer RNA; WES: Whole-exome sequencing

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Authors’ contributions

Dr. Nagasaka and Dr. Uddin developed the first draft of the manuscript. All authors contributed to the planning, organization, data collection, and writing of the manuscript. All reviewed the manuscript and provided critical edits. The final version of the manuscript was approved by all authors.

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NA

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