Altered editing level of microRNAs is a potential biomarker in lung adenocarcinoma

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Adenosine-to-inosine (A-to-I) microRNA editing is associated with tumor phenotypes in various cancer types. Recent analyses of The Cancer Genome Atlas (TCGA) dataset have shown several microRNAs that undergo A-to-I editing in human cancers, some of which have been reported to be associated with prognosis. Herein, we examined published small RNA deep sequencing data of 74 cases of lung adenocarcinoma (AD) and the corresponding normal counterpart (NC) specimen in silico in order to identify A-to-I microRNA editing events. Editing levels of miR-379-5p, miR-99a-5p, and miR-497-5p were lower in AD than in NC and, in a large number of cases, the editing level of miR-200b-3p was higher in AD than in NC. Difference in the editing level between AD and NC was largest for miR-99a-5p. Then, we examined the editing level of miR-99a-5p in 50 surgically resected lung adenocarcinoma cases at our institution by a conventional sequence-based method, and its association with clinical outcomes. The editing level of miR-99a-5p was significantly lower in 19 cases of AD (38%) than in corresponding NC. These cases showed a shorter overall survival as assessed using the log-rank test (P = .047). This trend was consistent with previous analyses of TCGA dataset. The altered editing level of microRNAs in lung adenocarcinoma could serve as a potential biomarker.

KEYWORDS biomarker, computational biology, lung adenocarcinoma, microRNA, RNA editing

1 | INTRODUCTION

Lung cancer is the leading cause of cancer-related death worldwide.1 Adenocarcinoma, which is a category of non-small-cell lung cancer (NSCLC), is the most common histological type among all lung cancer cases.2 Treatment outcome of unresectable or recurrent NSCLC has been improving with the introduction of novel molecular-targeted drugs3,4 and immune checkpoint inhibitors;5,6 nevertheless, these epoch-making drugs do not cure NSCLC. In the American Society of Clinical Oncology (ASCO) guidelines for completely resected NSCLC, adjuvant chemotherapy is recommended depending on the pathological stage alone.8 To pursue individualized treatments, a novel biomarker for predicting the clinical outcome after surgery is desired.

Adenosine-to-inosine (A-to-I) RNA editing is a posttranscriptional modification of specific RNAs, catalyzed by proteins of the adenosine deaminases acting on the RNA (ADAR) family, ADAR1, and ADAR2.9 As inosine base-pairs with cytosine in double strand nucleotides, A-to-I editing is functionally equivalent to an A-to-G mutation. The editing level largely varies among organs: The most frequently edited is brain mRNA, whereas lung mRNA is the second...
most frequently edited. Since 2004, when primary microRNAs, precursors of microRNAs were first demonstrated to undergo A-to-I editing, some studies have shown the role of microRNA editing in normal organs and various types of cancers. MicroRNAs are single-stranded RNA molecules that are approximately 22 nucleotides in length and function as posttranscriptional gene regulators. They bind to the 3′ UTR of mRNAs which contain a sequence complementary to the seed sequence (positions 2-8) on the microRNAs, resulting in degradation or downregulation of the target mRNAs. As a consequence of microRNA editing, altered base complementation of a primary microRNA molecule affects the cleavage activity of microRNA processing proteins, DROSHA and DICER. Moreover, when the seed sequence of a microRNA undergoes RNA editing, the edited microRNA targets another set of mRNAs. Therefore, the alteration in the editing levels of microRNAs affects the phenotype of a cancer through the alterations of the gene expression profile. Recently, in silico analyses of small RNA deep sequencing data of The Cancer Genome Atlas (TCGA) dataset showed the comprehensive microRNA editing profile of 32 types of cancer. In the cases of lung adenocarcinoma, significant differences in the editing levels of 5 microRNAs (miR-200b-3p, miR-379-5p, miR-411-5p, miR-497-5p, and miR-99a-5p) were found between adenocarcinoma (AD) samples and the corresponding normal counterpart (NC) samples, and correlations were found between the editing levels of 4 of these microRNAs (miR-376c-3p, miR-379-5p, miR-381-3p, and miR-99a-5p) in AD and the survival of the patients.

In the present study, we analyzed the published small RNA deep sequencing data of 74 cases of lung adenocarcinoma, including the data of both AD and NC in silico, and detected 3 microRNAs showing loss of editing in AD (the editing level was significantly lower in AD than in NC) and 1 microRNA showing gain of editing in AD (the editing level was significantly higher in AD than in NC). Regarding miR-99a-5p, which showed the largest difference of the editing level between AD and NC, we set up a conventional sequence-based method to quantify the editing levels. We examined surgically resected specimens from 50 cases of lung adenocarcinoma at our institution to detect differences of the editing level of miR-99a-5p between AD and NC. In our dataset, loss of editing of miR-99a-5p was significantly associated with shorter overall survival (OS), which was consistent with the analyses of TCGA dataset.

2 | MATERIALS AND METHODS

2.1 | In silico analysis of deep sequencing data

We searched the Sequence Read Archives (SRA) at the National Center for Biotechnology Information (https://www.ncbi.nlm.nih.gov/sra) using a search phrase, “miRNA lung adenocarcinoma.” We included paired data of lung adenocarcinomas and the corresponding normal tissues published before March 2015. The downloaded SRA files were analyzed using perl scripts as previously described. In short, 5′ and 3′ adaptors were removed by Process_reads.pl. The reads were mapped to the human reference genome (hg38) using Bowtie. Here, we trimmed 2 bases at the 3′ end that undergo massive modifications. We allowed a maximum of 1 mismatch per read with a quality score of at least 30, best unique alignment, and no cross-mapping. Then, the reads were remapped to the pre-microRNA sequence database by Analyze_mutation.pl, to count the read depth and the number of reads with mismatch. The original script used release 18 from miRBase (http://www.mirbase.org) as reference. Instead, we used the latest version, release 21. The data format differs slightly according to versions; revised scripts are shown in Docs S1 and S2, and reference files are shown in Docs S3-S5. Finally, sequencing errors were removed for each position of each pre-microRNA by Binomial_analysis.pl. Output data included name of the pre-microRNA, mismatch position, mismatch type (such as C-to-U), number of mismatched reads, total number of reads in the position, and P-value with Benjamini-Hochberg correction with a false discovery rate (FDR) of 0.05. Known single-nucleotide polymorphisms (SNP) were discarded by reference to miRNASNP Release 2.0 (http://bioinfo.life.hust.edu.cn/miRNASNP2/).

2.2 | Patient population and collection of samples

The study protocol was approved by the Institutional Ethical Review Board at the University of Tokyo Hospital (Tokyo, Japan), and written informed consent was obtained from each patient. Samples were obtained from patients who underwent lung resection for primary lung adenocarcinoma at the University of Tokyo Hospital between August 2007 to September 2011. We included all 27 patients with pathological stage IIA to IIIA disease and consecutive 23 patients with pathological stage IB disease. Pathological stage was defined according to the seventh edition of the TNM classification by the UICC. Recurrence-free survival (RFS) was defined as the period from the date of resection for the lung cancer to the date of diagnosis of disease recurrence or death from any cause. OS was defined as the period from the date of resection to the date of death from any cause. Characteristics of the 50 patients are listed in Table 1.

2.3 | Total RNA extraction from tissue

Fresh tissue specimens were immersed into 500 µL RNAlater solution (Ambion, Austin, TX, USA) overnight at 4°C. After removing the RNAlater solution, tissue specimens were frozen and stored at −20°C. Finely divided tissue pieces were placed in the ceramic beads tube of the Precellys Lysing Kit (Bertin Technologies, Saint-Quentin en Yveline, France) containing 1000 µL RNAiso plus (TaKaRa, Shiga, Japan), and homogenized twice using Precellys24 (Bertin Technologies) at the oscillation frequency of 6500 per minute for 23 seconds each. The homogenized solution was centrifuged at 20 000 × g for 5 minutes and total RNA in the supernatant was extracted with chloroform, precipitated with isopropanol and eluted with 20 µL DEPC (diethylpyrocarbonate)-treated water (Wako Pure Chemical Industries, Osaka, Japan).
2.4 | Quantification of the editing level of miR-99a-5p and validation of the method

An overview of this original method is shown in Figure 1. First-strand cDNA was synthesized from 5 ng total RNA using the TaqMan Advanced miRNA cDNA Synthesis Kit (Thermo Fisher Scientific, Waltham, MA, USA), in accordance with the manufacturer’s instructions.28 Then, 2 μL cDNA was amplified using 50 μL AmpliTaq Gold 360 Master Mix (Thermo Fisher Scientific) and primers at a final concentration of 400 nmol/L (F, 5′-GAT CGA ATT CCA TCT CAT CCC TGC GTG TCT-3′; R, 5′-GAT CGA ATT CTT TTT CAC AAG ATC GGA TCT AC-3′). Thermal cycling conditions consisted of preheating at 95°C for 10 minutes, followed by 37-40 cycles (optimized for each sample) at 95°C for 15 seconds, and 60°C for 60 seconds. The PCR product was purified using Wizard SV Gel and PCR Clean-Up System (Promega, Madison, WI, USA), in accordance with the manufacturer’s instructions. The eluate was digested with EcoRI-HF (New England Biolabs, Ipswich, MA, USA) at 37°C for 120 minutes, followed by heat inactivation at 65°C for 20 minutes. Then, T4 DNA ligase (New England Biolabs) was added, followed by incubation at room temperature for 30 minutes to concatenate the amplicons.29 Formation of a ladder with intervals of 73 base pairs was confirmed by electrophoresis. Taq DNA Polymerase (New England Biolabs), dATP, and dTTP were added, followed by incubation at 70°C for 20 minutes. The solution was loaded onto 1.2% Low Melting Agar (PH Japan, Hiroshima, Japan) with 1% Gel Indicator (BioDynamics Laboratory, Tokyo, Japan), and electrophoresed for 30 minutes. The ladder between approximately 300-700 base pairs was cut out, and melted at 65°C. β-Agarase (New England Biolabs) was added, followed by incubation at 42°C for 60 minutes. The solution was salted out, precipitated with ethanol, and eluted with nuclease-free water. The concatenated amplicons were ligated into the pGEM-T Easy Vector System (Promega), and clones with inserts of the expected length were sequenced using BigDye Terminator v3.1 (Thermo Fisher Scientific) with a primer on the vector, 5′-GCC AAG CTA TTT AGG TGA C-3′. We repeated the process of cloning and sequencing until we obtained in excess of 100 amplicons for each sample. The editing level in each sample was calculated as the number of edited reads divided by total reads.

To confirm that the proportion of edited reads reflected the editing level of miR-99a-5p, the unedited amplicon which consists of the 5′ adaptor, unedited miR-99a-5p and polyA was ligated into the pGEM-T Easy Vector System (Promega). The edited amplicon was ligated similarly. Edited and unedited plasmids were mixed, so that the proportion of the copy number of the edited plasmids was 5%, 15%, and 25%. These 3 templates were amplified, concatenated, and sequenced by the same method described above until amplicons in excess of 100 were obtained for each template.

2.5 | Quantitative RT-PCR

For mRNA analysis, 1 μg total RNA treated with DNase I (Invitrogen, Carlsbad, CA, USA) was reverse transcribed with Superscript III (Invitrogen) using random hexamer primers, in accordance with the manufacturer’s instructions. The cDNA was amplified with 10 μL Thunderbird SYBR qPCR Mix (Toyobo, Tokyo, Japan) and primers at a final concentration of 400 nmol/L, using the 7500 Fast Real-Time PCR System (Applied Biosystems, Carlsbad, CA, USA). Thermal cycling conditions consisted of preheating of 95°C for 5 seconds, followed by 40 cycles at 95°C for 15 seconds and 60°C for 60 seconds. Specificity of the PCR was confirmed by analyzing the melting curves. PCR was carried out in triplicate, and mean Ct values of 3 samples were used for the analysis. Relative expression level of each gene was determined after normalization to the expression level of GAPDH. For survival analysis, the cutoff values for high or low expression level were determined using receiver operating characteristic (ROC) curves.20 The following primer sets were used: ADAR1 (F, 5′-ACC GGT GCT TCA ACA CTA TGA-3′; R, 5′-CGG GAG ATT TCT GCA TGG-3′), ADAR2 (F, 5′-TGT CAA CTG GAC GGT AGG CGA CT-3′; R, 5′-TGC CGG CAG CTG GCA AT-3′), and GAPDH (F, 5′-CAC CAC CAA TGC TTA GCA C-3′; R, 5′-TGG CAG TTT CTA GAC GG-3′).

2.6 | Statistical analysis

Student’s t test (age) and Fisher’s exact test (the others) were used to compare patient characteristics between the two groups. Statistical significance of the difference in the editing level between AD and NC was evaluated using the chi-squared test with Benjami-
Hochberg correction with a FDR of 0.05 (in silico analysis) and Fisher’s exact test (conventional sequence-based analysis). Survival analyses were carried out using the Kaplan-Meier method, and survival curves were compared using the log-rank test and a Cox proportional-hazards model. Correlation between the editing level of each sample and the relative expression level of each gene (ΔCt value) was evaluated using Spearman’s rank-order correlation test. All statistical analyses outside the perl scripts were carried out using IBM SPSS Statistics version 24 (IBM SPSS, Chicago, IL, USA). All the tests in this study were two sided, and the statistical significance level was set at P < .05.

3 | RESULTS

3.1 | In silico analysis of deep sequencing data

We downloaded the SRA files of 94 cases (188 samples) as candidates for analysis. They were from a single study which examined the relationship between smoking history and alteration of microRNA expression level in lung adenocarcinoma. Twenty samples had an average spot length of 30 bases, and further analysis showed that these samples had extremely low alignment proportions of approximately 1% (mapped reads divided by total reads). Therefore, these 20 samples and corresponding tissue samples were excluded, and the remaining 148 samples were analyzed. Profiles of the 74 patients are shown in Table 2.

The perl scripts output, including all mismatch types, included a total of 4518 significant mismatch events in 148 samples. The events were detected in 186 mismatch spots. We detected 64 mismatch hotspots, in which 1-base mismatch events were detected in

| TABLE 2 | Characteristics of the 74 patients in the published database |
|----------|-------------------|
| n = 74   | Proportion, %     |
| Gender   |                   |
| Male     | 20                | 27                |
| Female   | 54                | 73                |
| Smoking  |                   |
| Current  | 38                | 51                |
| Former   | 20                | 27                |
| Never    | 16                | 22                |
| Stage    |                   |
| I        | 45                | 61                |
| II       | 19                | 26                |
| III      | 7                 | 9                 |
| IV       | 3                 | 4                 |
FIGURE 2  Distribution plot of the mismatch hotspots in AD (left panel) and NC (right panel). Each dot represents the proportion of the number of samples (out of 74) in which statistically significant mismatch events were observed (horizontal axis) and the median editing level among the cases (vertical axis). AD, adenocarcinoma; NC, normal counterpart

FIGURE 3  Distribution of A-to-I microRNA editing levels by in silico analysis. These 8 microRNAs showed significant A-to-I editing in more than 5 cases each of both AD and NC. Editing sites are shown in red letters. Each dot represents the editing level of AD and NC of each case showing (red) or not showing (blue) a statistically significant difference of the editing level. A, miR-379-5p (n = 65), miR-99a-5p (n = 60), and miR-497-5p (n = 23): loss of editing was seen in most cases. B, miR-200b-3p (n = 41): nearly half of the cases showed gain of editing. C, miR-381-3p (n = 71), miR-589-3p (n = 68), miR-411-5p (n = 22), and miR-6503-3p (n = 7): no significant differences of the editing level were observed in most cases. AD, adenocarcinoma; NC, normal counterpart
a common mismatch spot in at least 10 samples. However, most of the non-A-to-G mismatch hotspots had extremely low mismatch frequencies (the number of reads with a mismatch divided by the number of total mapped reads) and were detected in fewer samples than the A-to-G mismatch hotspots (Figure 2). Therefore, we regarded the non-A-to-G mismatch hotspots as noises, random sequencing errors which could not be eliminated by statistical procedure in the perl

| Mixture fraction | 5%   | 15%  | 25% |
|------------------|------|------|-----|
| Total amplicon   | 103  | 105  | 110 |
| Edited amplicon  | 8    | 12   | 23  |
| Observed proportion | 7.8% | 11.4% | 20.9% |
| 95% CI           | 3.4%-14.7% | 6.0%-19.1% | 13.7%-29.7% |
| P-value (binomial test) | 0.144 | 0.189 | 0.19 |

scripts. In regard to the 10 A-to-G mismatch hotspots, all of them were also detected in the previous study of TCGA dataset.22 These mismatch hotspots can be regarded as true editing sites, because they were firmly reproducible; we call them “editing hotspots” from here.

For the 10 A-to-I editing hotspots, we compared the editing levels of 8 microRNAs (paired editing hotspots) between AD and NC samples; for the remaining 2 microRNAs (miR-100-5p, miR-24-2-5p), only 1 editing event was detected in 74 AD samples. Distribution of the editing levels of the 8 microRNAs is shown in Figure 3. Loss of editing was detected in position 5 of miR-379-5p (52 cases out of the 65 cases in which RNA editing was detected in both AD and NC: 52/65), position 1 of miR-99a-5p (60/60), and position 2 of miR-497-5p (20/23) (Figure 3A). Gain of editing was detected in position 5 of miR-200b-3p (19/41) (Figure 3B). Among these 4 microRNAs, median value of the editing level difference between AD and NC was the largest for miR-99a-5p. Most cases showed no significant difference of editing level at position 4 of miR-381-3p (53/70), position 6 of miR-589-3p (67/68), position 5 of miR-411-5p (14/22), or position 7 of miR-6503-3p (7/7) (Figure 3C).

**TABLE 3** Results of the validation test for quantifying the editing level of miR-99a-5p. Mixture fraction indicates the proportion of the template with edited amplicon

All patients (n = 50)

| Mixture fraction | 5%   | 15%  | 25% |
|------------------|------|------|-----|
| Total amplicon   | 103  | 105  | 110 |
| Edited amplicon  | 8    | 12   | 23  |
| Observed proportion | 7.8% | 11.4% | 20.9% |
| 95% CI           | 3.4%-14.7% | 6.0%-19.1% | 13.7%-29.7% |
| P-value (binomial test) | 0.144 | 0.189 | 0.19 |

**FIGURE 4** A, Editing levels of miR-99a-5p in the 50 cases at our institution. The format of the scatter plot is the same as that in Figure 2. B-D, Kaplan-Meier curves of the overall survival and the recurrence-free survival, stratified by loss of editing of miR-99a-5p. D, represents the subgroup of pathological stage IB disease (n = 23)
3.2 | Quantification of the editing level of miR-99a-5p

Our improved method to quantify the editing level of miR-99a-5p shown in Figure 1 is based on two assumptions: (i) PCR efficiency of edited and unedited miR-99a-5p is equivalent; and (ii) the proportion of edited and unedited amplicons in concatemers is equivalent to that before ligation. To confirm that the proportion of edited reads reflected the editing level of the sample, we examined the standard template which contains 5%, 15%, and 25% edited amplicons (5′ adaptor, miR-99a-5p, and polyA). As shown in Table 3, each template showed results consistent with its mixture fractions.

Next, we extracted total RNA from 100 resected tissue specimens obtained from a cohort of 50 cases of lung adenocarcinoma and examined the editing level of miR-99a-5p in these specimens. In 43 cases (86%), the editing level in AD was lower than that in NC. After statistical analysis, 19 cases (38%) showed significant loss of editing and 1 case (2%) showed significant gain of editing (Figure 4A). Loss of editing was not significantly associated with any unique clinical features (age, \( P = 0.750 \); gender, \( P = 0.560 \); smoking history, \( P = 1 \); stage, \( P = 0.247 \); epidermal growth factor receptor (EGFR) mutation, \( P = 0.736 \); adjuvant chemotherapy, \( P = 0.389 \)).

Kaplan-Meier curves stratified by loss of editing are shown in Figure 4B-D. Cases with loss of editing showed significantly shorter OS (\( P = 0.047 \); hazard ratio [HR], 3.25; 95% confidence interval [CI], 0.95-11.13) and a tendency toward shorter RFS (\( P = 0.103 \); HR 1.91; 95% CI, 0.87-4.19). In the subgroup analysis, loss of editing was significantly associated with shorter RFS in the subgroup of pathological stage IB disease (\( n = 23; P = 0.028 \); HR, 5.08; 95% CI, 1.01-25.58).

Then, we compared the editing level of the 8 microRNAs with paired editing hotspots among our hospital dataset, downloaded SRA dataset, and previously reported TCGA dataset by Pinto et al.22 (Table 4). For miR-99a-5p, the editing level ranges largely overlapped between our hospital dataset and the SRA dataset. Overall editing level (total number of edited reads of all cases divided by the total number of mapped reads of all cases) of each microRNA in TCGA dataset was within the corresponding range in the SRA dataset, except for miR-497-5p. For miR-497-5p, the overall editing level in AD from TCGA dataset reported by Wang et al.21 (approximately 2.8% for AD; \( n = 483 \), raw data not available) was somehow within the corresponding range in the SRA dataset.

3.3 | Analysis of ADAR1 and ADAR2 expressions

Finally, we carried out quantitative RT-PCR for expression of ADAR1 and ADAR2, with normalization to the expression of GAPDH. Amplification efficiencies of ADAR1, ADAR2, and GAPDH calculated from the slope of the standard curves from standard samples were 91.4%, 95.5%, and 93.0%, respectively. Expression levels of both ADAR1 and ADAR2 were downregulated in AD as compared to NC (median fold change (\( 2^{-\Delta\Delta Ct} \)) of ADAR1, 0.52; ADAR2, 0.16). There was a moderate positive correlation between the editing level in each sample and the expression level of ADAR2 (\( r_s = 0.258; P = 0.009 \) for ADAR1, and \( r_s = 0.424; P < 0.001 \) for ADAR2) (Figure 5A), consistent with a previous study in which miR-99a-5p was reported to be edited by ADAR2.21

Then, the patients were divided into 2 groups according to the expression level (\( \Delta\Delta Ct \) value) of ADAR1 (high group, \( n = 24 \); low group, \( n = 26 \)) and ADAR2 (high group; \( n = 17 \), low group; \( n = 33 \)).

### Table 4 Comparison of the editing levels of the 8 microRNAs among the 3 datasets: the University of Tokyo Hospital, SRA, and TCGA

|                        | Our dataset | SRA | TCGA |
|------------------------|-------------|-----|-------|
|                        | n = 50      | n = 74 | n = 36 |
| miR-99a-5p             |             |       |       |
| Range                  | 0-12.3      | 0.79-6.49 | 3.57 |
| Overall                | 4.37        | 2.34 | 12.37 |
| miR-379-5p             |             |       |       |
| Range                  | 0.35-10.53  | 1.40 | 1.54  |
| Overall                | 1.07        | 5.76 | 4.50  |
| miR-497-5p             |             |       |       |
| Range                  | 0.30-0.37   | 0.30-5.05 | 5.56 |
| Overall                | 1.30        | 2.10 | 17.43 |
| miR-200b-3p            |             |       |       |
| Range                  | 9.48-57.14  | 24.32-87.02 | 70.73 |
| Overall                | 25.75       | 60.9 | 67.98 |
| miR-381-3p             |             |       |       |
| Range                  | 3.30-36.36  | 8.67 | 19.17 |
| Overall                | 23.08-76.92 | 28.27 | 35.32 |
| miR-589-3p             |             |       |       |
| Range                  | 2.10        | 2.10 | 6.40  |
| Overall                | 1.30        | 6.40 | 14.63 |
| miR-6503-3p            |             |       |       |
| Range                  | 47.72       | 47.72 | 39.26 |
| Overall                | 42.33       | 42.33 | 32.00 |

AD, adenocarcinoma; NC, normal counterpart; SRA, Sequence Read Archives; TCGA, The Cancer Genome Atlas.
Kaplan-Meier curves are shown in Figure 5B. The ADAR1-high group ($P = 0.087; \text{HR}, 0.331; 95\% \text{ CI}, 0.087-1.251$) and the ADAR2-high group ($P = 0.061; \text{HR}, 0.176; 95\% \text{ CI}, 0.023-1.376$) showed a tendency toward a longer OS.

4 | DISCUSSION

In the present study, in silico analysis identified 8 microRNAs with paired A-to-I editing hotspots. Furthermore, we demonstrated an improved method to quantify the editing level of miR-99a-5p. In our patient cohort, loss of editing was associated with poor prognosis after complete resection of lung adenocarcinoma. This tendency was consistent with the results of previous studies of TCGA dataset$^{21,22}$ reproducibility was confirmed by a well-validated conventional RT-PCR method. Along with these studies, we believe that the alteration of microRNA editing in lung adenocarcinoma could serve as a potential biomarker for predicting patient outcomes.

Before this biomarker can be applied practically in clinical medicine, the method for quantifying the editing level of microRNA needs to be further refined. In the first study to report microRNA editing, the authors amplified the precursor of miR-22, pri-miR-22, cloned the PCR product and sequenced at least 100 amplicons.$^{12}$ However, because editing of a primary microRNA affects the processing efficiency of the microRNA catalyzed by the double-strand RNase enzymes, DROSHA and DICER, the editing level of primary microRNA is not always equal to the editing level of mature microRNA, which finally functions as the posttranscriptional regulator.$^{32,33}$ A recent comprehensive analysis showed that the difference in the editing level between a primary microRNA and mature microRNA is >10% in approximately half of the microRNAs.$^{34}$ An improved method to amplify mature microRNA was published in 2012.$^{28}$ In this study, mature microRNA was 3’ polyadenylated and 5’ adaptor ligated, reverse-transcribed with oligo-dT primer, specifically amplified, and cloned. At least 100 amplicons were sequenced. Use of the TaqMan Advanced miRNA cDNA Synthesis Kit (Thermo Fisher Scientific) reduced the operation hours of this method. Furthermore, to reduce cost entailed by the vast amount of sequencing, we made use of the concept of serial analysis of gene expression (SAGE)$^{29}$ and established a protocol to concatenate the amplicons, as described above. Our improved method takes approximately 6 extra hours to concatenate the amplicon, while the number of sequenced samples was almost one-half (5322 samples used to sequence 11 110 amplicons). Moreover, our validation experiment showed the reliability of this method.

Another candidate method, which would be less time-consuming and less expensive, is to make use of TaqMan MGB (minor groove binder) Probe with 2 colors of fluorescent dye. After synthesizing first-strand cDNA with the 5’ adaptor and polyA using TaqMan Advanced miRNA cDNA Synthesis Kit, the expression level of each microRNA can be quantified with a specific primer set and TaqMan MGB Probe,
which are contained in TaqMan Advanced miRNA Assays (Thermo Fisher Scientific). Moreover, a method involving the use of the TaqMan Probe with 2 colors of fluorescent dye, FAM and VIC, to quantify the editing level of mRNA was established in a previous study. Combining these methods, both the expression and editing levels of micro-RNAs can be measured. A significant difficulty is that while securing the specificity of amplifying the microRNA, there is little space left for the TaqMan Probe to anneal on the editing spot, even with an MGB.

Previous studies have shown that miR-99a-5p itself functions as a tumor-suppressing microRNA through repressing invasion, migration, and stemness. However, the functional impact of editing miR-99a-5p is not clear, because it undergoes RNA editing at position 1. The seed sequence of the microRNAs, spanning positions 2-8, must be completely complementary to the 3’ UTR part of the target mRNA. A previous study reported that miR-17-5p and miR-106a-5p, whose sequences differ only in position 1, are regulated and function in parallel. Although the previous study of TCGA dataset showed that the editing level of miR-99a-5p was correlated with the mutation frequency of 14 important genes, including TP53, HRAS, NRAS and PIK3CA in 6 cancer types, this cannot be explained by the known function of microRNA editing. Therefore, we assume that the difference in the editing level of miR-99a-5p being the largest among the paired editing hotspots caused a “passenger change” in lung adenocarcinoma, rather than alteration of a specific function of miR-99a-5p.

In a previous study of TCGA dataset, expression level of ADAR1 in lung adenocarcinoma was reported to be upregulated as compared with that in the corresponding normal tissue. Moreover, amplification of the ADAR1 region and higher expression level of ADAR1 in cancerous tissue were associated with a tendency toward a poorer prognosis, although the association was not statistically significant for either. Our dataset showed an inverse tendency. We did not identify any clinical features (age, gender, smoking history, stage, EGFR mutation status) that were significantly associated with the expression levels of ADAR1 in our dataset. The largest difference between our cohort and TCGA cohort was in the ethnicity of the patients (100% East Asian vs 67.2% White and 1.4% Asian), which may have affected tumor biogenesis. We consider that study of more cases is needed to verify the function of ADAR1 in lung adenocarcinoma. Regarding ADAR2, our findings were consistent with previous studies; ADAR2 was downregulated in lung adenocarcinoma tissues, and a lower ADAR2 expression level was significantly associated with a poorer prognosis. In summary, we analyzed the alterations of microRNA editing in lung adenocarcinoma both in silico and by a conventional RT-PCR method. Loss of editing of miR-99a-5p in cancerous tissue was associated with poor survival in surgically resected lung adenocarcinoma specimens. Further investigations are required to elucidate the cause and effect of this association and to refine the method for quantification of the editing level.

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CONFLICTS OF INTEREST

Authors declare no conflicts of interest for this article.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.