Genomics and Prognosis Analysis of N6-Methyladenosine Regulators in Lung Adenocarcinoma

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Objective: N6-methyladenosine (m6A) modification is involved in modulating various biological processes in human cancers. But the implication of m6A modification in lung adenocarcinoma (LUAD) is still unclear. Hence, this study conducted a comprehensive analysis of the expression and clinical implication of m6A regulators in LUAD.

Methods: Consensus clustering analysis of 502 LUAD samples in the TCGA dataset was presented based on the expression profiles of 20 m6A regulators using ConsensusClusterPlus package. Overall survival (OS), activation of signaling pathways and tumor immunity (immune/stromal score, tumor purity, expression of HLA and immune checkpoints, and immune cell infiltration) were compared between m6A modification patterns. The m6A-related genes between patterns were identified and prognostic m6A-related genes were imported into LASSO-cox regression analysis. The m6A risk score was developed and its prognostic implication was evaluated and externally verified in the GSE30219 and GSE72094 dataset. Furthermore, a nomogram that contained independent prognostic indicators was established, followed by external verification.

Results: Two m6A modification patterns were clustered across LUAD based on the expression similarity of the m6A regulators via consensus clustering analysis, with distinct OS, activation of signaling pathways and tumor immunity. Totally, 213 m6A-related genes that were identified by comparing two patterns were significantly related to LUAD prognosis. By LASSO method, we constructed the m6A risk score that was a reliable and independent prognostic factor for LUAD. Patients with low m6A risk score displayed a prominent survival advantage. After incorporating independent clinical features, we developed the prognostic nomogram that exhibited high predictive accuracy and the best clinical net benefit for OS.

Abbreviations: NSCLC, non-small cell lung cancer; LUAD, lung adenocarcinoma; m6A, N6-methyladenosine; TCGA, The Cancer Genome Atlas; GEO, Gene Expression Omnibus; GSVA, Gene set variation analysis; ESTIMATE, Estimation of Stromal and Immune Cells in Malignant Tumors Using Expression Data; HLA, human leukocyte antigen; ssGSEA, single-sample gene set enrichment analysis; DEGs, differentially expressed genes; FC, fold change; FDR, false discovery rate; LASSO, least absolute shrinkage and selection operator; OS, overall survival; ROC, receiver operating characteristic; AUC, area under curve; HR, hazard ratio.
INTRODUCTION

Lung cancer has the high incidence and mortality globally, occupying almost 20% of cancer-related deaths in 2018 (Bray et al., 2018). It was estimated that there were 2.1 million new lung cancer cases and 1.8 million deaths in 2018 (Bray et al., 2018). This disease mainly includes two histological subtypes: non-small cell lung cancer (NSCLC; 85%) and small cell lung cancer (SCLC). NSCLC contains lung adenocarcinoma (LUAD) and lung squamous cell carcinoma (Zhu et al., 2019). LUAD is the main histology, and its incidence is constantly on the rise. Conventional therapeutic options against NSCLC include surgery resection, chemotherapy, and radiotherapy. Despite the progress in combined and personalized therapies such as tyrosine kinase inhibitors and immunotherapies (PD1/PD-L1 inhibitors), the 5-year survival rate is only 16% (Zhang C. et al., 2020). Diagnosis of LUAD usually occurs at an advanced stage, and most patients experience badly toxic treatment and poor clinical benefit (Schmidt et al., 2019). Hence, it is of importance to explore specific prognostic models for predicting patients’ survival, which can assist design appropriate therapeutic strategies and management choice for distinct LUAD subgroups.

N6-methyladenosine (m6A) is the most abundant type of RNA post-transcriptional modification in eukaryotes, which plays a key role in a variety of biological processes by regulating the translation, splicing, stabilization, and degradation of mRNAs (Zhang H. et al., 2020). Typically, m6A regulators contain three types: writers (including VIRMA, METTL14, METTL3, RBM15, RBM15B, RBMX, WTAP, and ZC3H13), erasers (including ALKBH5 and FTO), and 10 readers (HNRNPA2B1, HNRNPC, IGF2BP1, IGF2BP2, IGF2BP3, WTAP, and ZC3H13), 2 erasers (ALKBH5 and FTO), and 10 readers (HNRNPA2B1, HNRNPC, IGF2BP1, IGF2BP2, IGF2BP3, YTHDC1, YTHDC2, YTHDF1, YTHDF2, and YTHDF3) were collected from the published literature. The location of these m6A regulators on the human chromosomes was plotted through Circos package (version 1.2.1) (Zhang et al., 2013). Protein-protein interaction analysis of the m6A regulators was performed by the STRING online database (version: 11.0; https://string-db.org/) (Szklarczyk et al., 2017).

Gene Set Variation Analysis

Consensus clustering analysis was carried out utilizing ConsensusClusterPlus package (version 1.48.0) to assign LUAD patients in the TCGA dataset into different m6A modification patterns with 50 iterations and resample rate of 80% based on the expression matrix of the 20 m6A regulators (Willkerson and Hayes, 2010). Kaplan-Meier curves of overall survival (OS) were conducted between two m6A modification patterns. The survival difference was compared with log-rank test. The t-distributed stochastic neighbor embedding (t-SNE) was presented to validate the accuracy of this classification.

Conclusion: Collectively, our study may provide a clinically useful tool for precise prognostic management and optimization of immunotherapeutic strategies for LUAD patients.

Keywords: lung adenocarcinoma, N6-methyladenosine, molecular subtypes, risk score, prognosis, tumor immunity

MATERIALS AND METHODS

Dataset Preparation

The Cancer Genome Atlas database (TCGA) RNA-seq data (FPKM values) and matched clinical features of 502 LUAD patients were retrieved from the Genomic Data Commons website (https://portal.gdc.cancer.gov/). The FPKM values were normalized with transcripts per million (TPM) method, followed by log2 conversion. Microarray expression profiling and clinical information of 274 LUAD samples and 398 LUAD samples were separately obtained from the GSE30219 dataset (Roussaux et al., 2013) and the GSE72094 dataset (Schabath et al., 2016) in the Gene Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/geo/). All data were obtained from the publicly available databases. Therefore, it was not applicable for the ethical approval. A total of 20 m6A regulators including 8 writers (VIRMA, METTL14, METTL3, RBM15, RBM15B, RBMX, WTAP, and ZC3H13), 2 erasers (ALKBH5 and FTO), and 10 readers (HNRNPA2B1, HNRNPC, IGF2BP1, IGF2BP2, IGF2BP3, YTHDC1, YTHDC2, YTHDF1, YTHDF2, and YTHDF3) were collected from the published literature. The location of these m6A regulators on the human chromosomes was plotted through Circos package (version 1.2.1) (Zhang et al., 2013). Protein-protein interaction analysis of the m6A regulators was performed by the STRING online database (version: 11.0; https://string-db.org/) (Szklarczyk et al., 2017).

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Gene Set Variation Analysis

The activation of pathways was quantified in each LUAD sample from the TCGA dataset by single-sample gene set enrichment analysis (ssGSEA) method derived from GSVA package (version 1.32.0) in an unsupervised manner (Hänzelmann et al., 2013). The gene set of “c2. cp.kegg.v7.2. symbols” was obtained from the Molecular Signatures Database, which was used as the reference set (Liberzon et al., 2015).
Estimation of Tumor Immunity
According to the normalized expression matrix, stromal and immune scores across LUAD samples in the TCGA dataset were estimated via the Estimation of Stromal and Immune Cells in Malignant Tumors Using Expression Data (ESTIMATE) method (https://sourceforge.net/projects/estimateproject/). (Yoshihara et al., 2013) that was applied for inferring the overall infiltrations of stromal and immune cells in LUAD tissues based on gene symbols. The tumor purity was calculated via ESTIMATE and consensus measurement of purity estimations methods. Tumor immune signatures were assessed in LUAD samples, including the mRNA expression of human leukocyte antigen (HLA) family genes and immune checkpoints. The infiltration levels of immune cells were quantified across LUAD samples based on the published gene signatures utilizing the ssGSEA algorithm (Charoentong et al., 2017; Jia et al., 2018).

Identification of m6A-Related Differentially Expressed Genes
The DEGs were screened between two m6A modification patterns in the TCGA dataset through limma package (version 3.40.6) (Ritchie et al., 2015). The cut-off was [log2 fold change (FC)] > 1 and false discovery rate (FDR) < 0.001. FDR was calculated with Benjamin–Hochberg method. The m6A-related DEGs were visualized into volcano and heat maps via pheatmap package (version 1.0.12).

Construction of a Least Absolute Shrinkage and Selection Operator-Cox Regression Model
LASSO represents a regularization and descending dimension method that has been applied for prognostic Cox models. Univariate-cox regression analysis was utilized to assess the correlation between overall survival (OS) of LUAD patients in the TCGA dataset and the m6A-related DEGs. The genes with p < 0.05 were input into the LASSO-cox regression model through glmnet package (version 2.0.16) (Engebretsen and Bohlin, 2019). Variable selection was presented for penalizing the data fitting criteria, which reduced the complexity and made the model more interpretable. The coefficient of each variable was the average estimate of the coefficient obtained from 10-fold cross-validation. The m6A risk score was developed following the formula: risk score = \sum_{i=1}^{n} \text{Coef } f(i) \times X(i), where n indicated the number of variables in this LASSO model, Coef 1) represented the regression coefficient, and X 1) meant the mRNA expression levels of variables in LUAD samples. To evaluate the prediction utility of the LASSO model, time-dependent receiver operating characteristic (ROC) curves were conducted by survivalROC package (version 1.0.3) in the TCGA, GSE30219 and GSE72094 datasets, followed by calculation of one, three and 5-year area under curve (AUC). In the two datasets, LUAD patients were separately split into two groups according to the median m6A risk score through survminer (version 0.4.9) and survival (version 3.2–13) packages. Kaplan-Meier curves of OS were depicted for two groups via survival package and OS difference was compared with log-rank test. The distribution of survival status in two groups was then visualized. By heatmap package, heatmap was established to visualize the mRNA expression pattern of each variable in the LASSO model.

Estimation of the Prediction Independency of the m6A Risk Score
To estimate whether the m6A risk score independently predicted LUAD patients’ OS, univariate- and multivariate-cox regression analysis was carried out following adjusting clinical features (gender, stage, T, N and M) in the TCGA, GSE30219 and GSE72094 datasets. Hazard ratio (HR) and p value were calculated for each variable.

Construction of a Nomogram Model
To better apply the m6A risk score in clinical practice, a nomogram that included independent prognostic indicators was conducted to predict LUAD patients’ one, three and 5-year OS in the TCGA, GSE30219 and GSE72094 datasets via rms package (version 6.2–0). Calibration plot was presented to evaluate predictive performance of the m6A risk score. Furthermore, decision curve analysis was carried out for calculating the clinical net benefit of every model in comparison to all or none strategies. The none plots indicated the assumption that no subjects had one, three or 5-year OS. Meanwhile, all plots indicated the assumption that each subject had one, three or 5-year OS at specific threshold probabilities. The best model was the one with the highest net benefit.

Statistical Analysis
All statistical analysis was implemented through the R software (version 3.6.3). Wilcoxon test was used for comparison between two groups. p < 0.05 was statistically significant.

RESULTS
Landscape of Expression and Prognostic Implications of m6A Regulators in Lung Adenocarcinoma
Totally, 20 m6A regulators including 8 writers (VIRMA, METTL14, METTL3, RBM15, RBM15B, RBMX, WTAP, and ZC3H13), 2 erasers (ALKBH5 and FTO), and 13 readers (HNRNPA2B1, HNRNPC, IGF2BP1, IGF2BP2, IGF2BP3, IGF2BP3, YTHDC1, YTHDC2, YTHDF1, YTHDF2, and YTHDF3) were compared between LUAD and normal tissues. There were close direct (physical) or indirect (functional) interactions between the m6A regulators (Figure 1A). Pan-cancer analysis revealed the prognostic implication of the m6A regulators in the TCGA cohort (Figure 1B). For LUAD, IGF2BP1, IGF2BP2, IGF2BP3, HNRNPAP2B1, HNRNPC, VIRMA, RBM15 and ALKBH5 were significant risk factors. The mRNA expression of the 20 m6A regulators was compared between LUAD and normal tissues.
We found that METTL3, VIRMA, RBM15, RBMX, YTHDF1, YTHDF2, IGF2BP1, IGF2BP3, HNRNPA2B1 and HNRNPC displayed higher mRNA expression in LUAD compared to normal specimens (Figure 1C). Meanwhile, METTL14, WTAP, ZC3H13, ALKBH5 and FTO were significantly down-regulated in LUAD than normal tissues. These data were indicative of the important implication of m6A regulators in the progression of LUAD.

**Construction of m6A Regulators-Mediated m6A Modification Patterns in Lung Adenocarcinoma**

A total of 502 LUAD samples were clustered based on the expression similarity of the m6A regulators via consensus clustering analysis. Our data found that when the number of groups (k) = 2, there was an excellent clustering among LUAD samples in the consensus matrix (Figure 2A). Consensus
FIGURE 2 | Construction of m\textsuperscript{6}A regulator-mediated m\textsuperscript{6}A modification patterns in LUAD from the TCGA cohort. (A) Consensus matrix when number of groups \(k = 2\). In the consensus matrix, white meant that samples were impossibly clustered together, and dark blue meant that samples were always clustered together. Both rows and columns of the matrix represented samples. (B) Consensus cumulative distribution function (CDF) diagram when different \(k\) values. (C) Delta area plot for relative change in the area under CDF curve for \(k\) compared to \(k-1\). (D) Tracking plot for sample clustering when different \(k\) values. (E) Kaplan-Meier curves of OS between two m\textsuperscript{6}A modification patterns. The survival probabilities were compared with log-rank test. (F) Heatmap for the expression patterns of 20 m\textsuperscript{6}A regulators in different m\textsuperscript{6}A modification patterns, age, survival status, gender, stage, T, N and M. (G) The t-SNE plots for verifying the differences between two m\textsuperscript{6}A modification patterns. (H) GSVA for the activation of signaling pathways in LUAD samples between two m\textsuperscript{6}A modification patterns.
cumulative distribution function (CDF) diagram showed that when k value = 2, CDF reached an approximate maximum (Figure 2B). Delta area plot depicted the relative change in the area under CDF curve for k compared to k-1 (Figure 2C). As shown in tracking plot, when k value = 2, sample classification was stable (Figure 2D). Hence, we clustered LUAD patients into two m6A modification patterns, named as C1 (N = 318) and C2 (N = 184). To further understand the characteristics of m6A modification patterns clustered by consensus clustering analysis in LUAD, we firstly analyzed the difference in OS. The data showed that C2 exhibited a more unfavorable OS in comparison to C1 (p = 0.00054; Figure 2E). Furthermore, we visualized the...
expression patterns of the m6A regulators in two m6A modification patterns. As shown in Figure 2F, IGF2BP2, IGF2BP1, IGF2BP3 had distinctly higher expression in C2 compared to C1. The t-SNE was carried out for verifying whether the categories were appropriate. Our results showed that most of samples from C1 and C2 were separately gathered (Figure 2G), indicating that the clustering of two m6A modification patterns was a relatively good choice. By applying GSVA algorithm, activation of several signaling pathways was quantified in each LUAD sample. We found that E2F targets, G2M checkpoint, MYC targets, mTORC1 signaling, DNA repair and unfolded protein response had higher activations in C2 than C1 (Figure 2H).

**Two m6A Modification Patterns Characterized by Distinct Tumor Immunity**

The overall infiltration levels of immune and stromal cells were estimated in 502 LUAD samples from the TCGA dataset via ESTIMATE algorithm. Compared to C1, C2 pattern had a significantly decreased immune score (p = 0.0025; Figure 3A). But no significant difference in stromal score was detected between m6A modification patterns (Figure 3B). There was significantly increased tumor purity in C2 than C1 (p = 0.049; Figure 3C). The mRNA expression of HLA genes and immune checkpoints was compared between two m6A modification patterns. Most of HLAs were highly expressed in C1 compared to C2, including HLA-E, HLA-DPB2, HLA-J, HLA-DQB1, HLA-DQB2, HLA-DQA1, HLA-DMA, HLA-DOB, HLA-DRB1, HLA-DRB5, HLA-DOA, HLA-DRB1, HLA-DRB6, HLA-F, HLA-DMB and HLA-DPA1 (Figure 3D). We also evaluated the differences in mRNA expression of common immune checkpoint between two m6A modification patterns. Our results showed that BTLA, CD200R1, CD40LG, CD48, HHLA2, IDO2, LGALS9, TNSFRSF14, TNSFRSF14 and TNSFRSF15 displayed higher mRNA expression in C1 compared to C2 (Figure 3E). Meanwhile, C2 pattern had increased mRNA expression of CD200, CD274, CD276, IDO1, LAG3, PDCD1, PDCD1LG2, TNSFRSF25, TNSFRSF8, TNSFRSF9, TNSFRSF4 and VTCN1 in comparison to C1. The infiltration levels of immune cells were quantified in each LUAD specimen via ssGSEA algorithm. Compared to C2, there were increased infiltration levels of activated B cell, activated CD8 T cell, central memory CD4 T cell, effector memory CD8 T cell, immature B cell, T follicular helper cell, type 17 T helper cell, activated dendritic cell, eosinophil, immature dendritic cell, mast cell and monocyte in C1 (Figure 3F). The higher infiltration levels of activated CD4 T cell, central memory CD8 T cell, memory B cell, type 2 T helper cell and plasmacytoid dendritic cell were found in C2 than C1.

**Identification of DEGs Between Two m6A Modification Patterns**

To explore the molecular mechanisms underlying two m6A modification patterns, we presented differential expression analysis. With the cutoff of |log2FC|>1 and adjusted p < 0.001, a total of 297 genes exhibited abnormal expression between two m6A modification patterns (Figures 4A,B). Among them, 111 genes were down-regulated and 186 were up-regulated in C1 compared to C2 (Supplementary Table S2). These genes could be affected by m6A methylation modification in LUAD.

**Development of an m6A Risk Score for Lung Adenocarcinoma**

Prognostic implications of the 297 m6A-related DEGs were assessed via univariate-cox regression analysis. As a result, 213 genes had significant correlations to LUAD prognosis in the TCGA dataset (Supplementary Table S3). Candidate prognostic m6A-related DEGs were further screened with LASSO-Cox regression analysis (Figures 5A,B). As a result, 12 candidate m6A-related DEGs were identified for constructing a LASSO-cox regression model. Non-zero coefficients and the expression of 12 m6A-related DEGs in this LASSO regression model were calculated in the TCGA dataset. The m6A risk score formula was as follows: m6A risk score = 0.0301860610758127 * mRNA expression of ANLN + 0.0263443393604996 * mRNA expression of PLK1 + 0.0576834829109261 * mRNA expression of IGF2BP1 + 0.023636868862302 * mRNA expression of HMMR + 0.035623995104486 * mRNA expression of NEIL3 + (-0.00157287764972551) * mRNA expression of SFTA3 + (-0.0250630515726943) * mRNA expression of CXCL17 + (-0.0244060686771379) * mRNA expression of IRX5 + 0.027706067741147 * mRNA expression of PKP2 + 0.028163642957098 * mRNA expression of LYPD3 + 0.0091755940795636 * mRNA expression of ABCCC2 + 0.157371504648263 * mRNA expression of DKK1. ROC curves were conducted to evaluate whether the m6A risk score accurately and sensitively estimated the survival likelihood of LUAD patients in the TCGA dataset. The AUC values of one, three and 5-year OS were separately 0.751, 0.690 and 0.611 (Figure 5C). These indicated the good predictive performance of the m6A risk score. Figure 5D depicted the distribution of the m6A risk score across LUAD patients. According to the median m6A risk score, patients were split into high- and low-m6A risk score groups. The OS difference was compared between groups. As shown in Figure 5E, low m6A risk score group displayed a potential survival advantage in comparison to high m6A risk score group (p < 0.0001). The distribution of survival status was visualized in Figure 5F. We found that high m6A risk score group had the relatively increased number of dead status than low m6A risk score group. Figure 5G showed the mRNA expression of 12 variables in the model between high- and low-m6A risk score groups. DKK1, PKP2, LYPD3, NEIL3, HMMR, ANLN, PLK1, IGF2BP1 and ABCCC2 displayed higher mRNA expression in high-m6A risk score group compared to low-m6A risk score group.

**External Verification of the m6A Risk Score in Lung Adenocarcinoma Prognosis**

To externally verify the prognostic implication of the m6A risk score, we acquired the transcriptome data and follow-up information of 274 LUAD patients from the GSE30219 cohort.
The AUC values under one, three and 5-year OS were separately 0.663, 0.677 and 0.694 (Figure 6A). According to the median m6A risk score, we clustered these LUAD patients into two groups (Figure 6B). High m6A risk score group had more patients with dead status than low m6A risk score group (Figure 6C). Consistently, high m6A risk score was markedly associated with worse prognosis of LUAD in comparison to low m6A risk score (p < 0.0001; Figure 6D). Furthermore, DKK1, PKP2, LYPD3, NEIL3, HMMR, ANLN, PLK1, IGF2BP1, and ABCC2 were highly expressed in high m6A risk score group than low m6A risk score group (Figure 6E). We also verified the prognostic significance of the m6A risk score in the GSE72094 dataset. The AUC values at one, three and 5-year OS were 0.699, 0.635, and 0.663 (Figure 6F). As expected, high m6A risk score indicated poorer OS than low m6A risk score (Figure 6G). Therefore, the m6A risk score possessed the potential in accurately predicting survival outcomes of LUAD patients.

The m6A Risk Score Acts as an Independent Prognostic Indicator of Lung Adenocarcinoma

In the TCGA dataset, univariate-cox regression analysis showed that the m6A risk score (p < 0.001; HR: 5.227 (3.347–8.163)), stage (p < 0.001; HR: 1.674 (1.458–1.923)), T (p < 0.001; HR: 1.530 (1.271–1.843)), N (p < 0.001; HR: 1.705 (1.437–2.023)) and M (p = 0.007; HR: 2.106 (1.229–3.609)) were significantly associated with LUAD prognosis (Figure 7A). These prognostic factors were input into multivariate-cox regression analysis. In Figure 7B, m6A risk score [p < 0.001; HR: 4.373 (2.618–7.306)] and stage [p = 0.029; HR: 1.533 (1.046–2.246)] were independent prognostic factors of LUAD. The prognostic implication of the m6A risk score was externally verified in the GSE30219 and GSE72094 cohorts. Our results confirmed that the m6A risk score could independently predict LUAD prognosis both in the GSE30219 cohort (Figures 7C,D) and GSE72094 cohort (Figures 7E,F).

Establishment and Verification of a Prognostic Nomogram for Lung Adenocarcinoma Patients

A nomogram was built for predicting one, three and 5-year OS likelihood of LUAD patients in the TCGA dataset, including the m6A risk score and stage that were obtained from multivariate-cox regression analysis (Figure 8A). Calibration diagram demonstrated that there was a high consistency in this nomogram-predicted and actual one, three and 5-year OS probabilities (Figures 8B–D). Moreover, decision curves suggested that the nomogram showed the best clinical net benefit for one, three and 5-year OS (Figures 8E–G). The nomogram was externally verified in the GSE30219 cohort (Supplementary Figure S1A–G) and GSE72094 cohort (Supplementary Figure S2A–G).

DISCUSSION

LUAD is usually diagnosed at an advanced stage, characterized by a high mortality (Zhu et al., 2020). The development of more effective therapeutic strategies requires an in-depth understanding of factors that impact the initiation and
progression of LUAD. Recently, several studies have reported the key implications of m6A regulators during LUAD development (Li F. et al., 2020; Chao et al., 2020; Li Y. et al., 2020). For example, Zhuang et al. constructed a diagnostic score model and a prognostic model for LUAD based on m6A regulators (Zhuang et al., 2020). Zhou et al. characterized two molecular subtypes with diverse prognosis and tumor microenvironment in LUAD based on m6A RNA methylation modification (Zhou et al., 2021). Wu et al. developed a five-m6A regulatory gene signature as a prognostic biomarker in LUAD patients (Wu X. et al., 2021). Xu et al. proposed m6A-related lncRNAs as potential biomarkers for prediction of prognosis and immune response in patients with LUAD (Xu et al., 2021). However, more studies should be presented for investigating the biological significance of m6A regulators in LUAD progression and prognosis. Hence, this study systematically analyzed the abnormal expression and clinical implications of m6A regulators.

FIGURE 5 | Development of an m6A risk score for LUAD by LASSO-cox regression analysis in LUAD samples from the TCGA cohort. (A) Partial likelihood deviance in each lambda value for the LASSO-cox regression analysis. (B) Process of variable selection in the LASSO-cox regression model by 10-fold cross-validation. (C) ROC curves under one, three and 5-year OS based on the m6A risk score. (D) The distribution of the m6A risk score and identification of the median m6A risk score indicated by vertical dotted line. (E) Kaplan-Meier curves of OS for high- and low-m6A risk score groups, followed by log-rank test. (F) The distribution of survival status in high- and low-m6A risk score groups. Red dots indicated dead status and blue dots indicated alive status. (G) Hierarchical clustering analysis for the mRNA expression patterns in LUAD samples from high- and low-m6A risk score groups.
By consensus clustering analysis, we established two m6A modification patterns with distinct survival outcomes based on the expression matrix of 20 m6A regulators for LUAD (Figure 2). Immune evasion represents a "hallmark of cancer," which reflects that immune effector constitutes a key determinant in the tumor microenvironment (Liu J. et al., 2020). Exploring the interactions between tumors and corresponding immune cells may reveal powerful and novel treatment options against LUAD.
Immunotherapies like PD1/PD-L1 inhibitors have become the standard-of-care therapeutic strategy against NSCLC (Cui et al., 2019). But, only 20–30% patients respond to such therapy (Rittmeyer et al., 2017). Limited data are available concerning the interactions between markers and immunotherapy response. Hence, it is necessary to explore and identify effective tumor immunity-related markers for LUAD, thereby reducing the mortality and developing innovative targeted therapeutic options. Compared to C2, C1 had an increased immune score and most of HLAs were highly expressed in C1, indicating that patients in C1 pattern exhibited higher tumor immunity (Figure 3).

Much progress in genome-wide methods like RNA-seq and microarrays has accelerated the evolution of cancer biomarker-related research. Numerous genetic markers of LUAD have been discovered, which are significantly correlated to diagnosis, survival outcomes, and drug resistance. But most of studies are limited to a single marker or a small sample population, leading to the limited accuracy and availability of markers. Furthermore, due to tumor heterogeneity, conventional clinical parameters like TNM staging are difficult to meet the requirement of accuracy and individuation in predicting prognosis. Thus, combined multiple markers or large sample analysis are necessary. Here, we established an m6A risk score...
containing ANLN, PLK1, IGF2BP1, HMMR, NEIL3, SFTA3, CXCL17, IRX5, PKP2, LYPD3, ABCC2, and DKK1 for LUAD prognosis (Figure 5). High m^6^A risk score indicated reduced OS duration of LUAD patients after external validation in two cohorts (Figure 6). Following multivariate Cox regression analysis, m^6^A risk score was an independent prognostic indicator of LUAD (Figure 7). ROC curves confirmed the excellent performance in predicting LUAD patients’ outcomes.

Previously, ANLN up-regulation is in relation to LUAD metastasis (Xu et al., 2019). ANLN promotes LUAD progression via activating RHOA and involving the PI3K/AKT signaling (Suzuki et al., 2005). PLK1/vimentin pathway promotes immune escape
through recruitment of Smad2/3 to PD-L1 promoter in LUAD metastasis (Jang et al., 2021). PLK1 induces migration of LUAD epithelial cells through STAT3 (Yan et al., 2018). IGF2BP1 induces LUAD progression via interaction with circXPO1 (Huang et al., 2020). Up-regulation of IGF2BP1 contributes to an unfavorable prognosis of LUAD (Huang et al., 2019). HMMR acts as an oncogene of LUAD and induces tumor progression (Li W. et al., 2020). NEIL3 that is correlated to immune infiltrations serves as an independent indicator for prediction of LUAD survival (Zhao et al., 2021). CXCL17 is an important determinant for LUAD spine metastasis (Liu W. et al., 2020). IRX5 as an oncogene is in relation to LUAD outcomes (Zhang et al., 2018). PKP2 accelerates the development of LUAD through increasing focal adhesion and EMT (Wu Y. et al., 2021). Elevated expression of LYPD3 contributes to LUAD carcinogenesis and unfavorable survival outcomes (Hu et al., 2020). ABC2C, a multidrug resistance-associated protein, displays an increased expression in LUAD (Maruhashi et al., 2018). DKK1 is an immune-associated prognostic marker in LUAD (Zhang et al., 2019). Above findings revealed the critical biological implications of the variables in the m6A risk score in the progression of LUAD.

Furthermore, our data indicated that in comparison to the nomogram established by a single prognostic indicator, the nomogram established by the m6A risk score and clinical features might become the best model in prediction of short- and long-term survival of LUAD patients, thereby possibly assisting clinical management and therapy (Figure 8). However, there are some limitations in our study, as follows: firstly, more information should be provided for internal mechanisms of m6A modification; secondly, the prognostic value of the m6A risk score should be verified in prospective research.

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CONCLUSION
Collectively, this study comprehensively characterized the expression and clinical implication of m6A regulators in LUAD. Two m6A modification patterns were conducted, with different OS and tumor immunity. Furthermore, we developed the m6A risk score, which had high accuracy in predicting LUAD prognosis. Thus, our data may provide a reliable tool for prediction of prognosis and optimization of immunotherapy for LUAD patients.

DATA AVAILABILITY STATEMENT
The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS
HZ conceived and designed the study. YM conducted most of the experiments and data analysis, and wrote the manuscript. All authors reviewed and approved the manuscript.

SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2021.746666/full#supplementary-material

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