

Identification of prognostic indicator based on hypoxia-related lncRNAs analysis in lung adenocarcinoma

Jiaojiao Zhang¹, Blessed Kondowe², Hui Zhang³, Xinming Xie⁴, Qiang Song⁵, Bo Guo^{6*}, Jin Shang^{3*}

¹ Department of Pathology, the First Affiliated Hospital of Xi'an Jiaotong University, Xi'an, P.R. China

² Radiology Department, Mzuzu Central Hospital, Mzuzu, Malawi

³ Department of Medical Imaging, the First Affiliated Hospital of Xi'an Jiaotong University, Xi'an, P.R. China

⁴ Department of Respiratory and Critical Care Medicine, the First Affiliated Hospital of Xi'an Jiaotong University, Xi'an, P.R. China

⁵ Department of Cardiovascular Medicine, the First Affiliated Hospital of Xi'an Jiaotong University, Xi'an, P.R. China

⁶ Institute of Genetics and Developmental Biology, Translational Medicine Institute, School of Basic Medical Sciences, Xi'an Jiaotong University Health Science Center, Xi'an, P.R. China

*Corresponding Author: Bo Guo and Jin Shang contributed equally to this work; E-mail: bo_guo@xjtu.edu.cn; shangjin01@qq.com

Abstract

Introduction

There were no systematic studies about hypoxia-related long noncoding RNAs (lncRNAs) signatures to predict the survival of patients with lung adenocarcinoma (LUAD). Setting up matching hypoxia-related lncRNA signatures was necessary.

Objective

This study aimed to establish hypoxia-related lncRNAs signatures and to seek new biomarkers to predict the prognosis of the patients with lung adenocarcinoma.

Methodology

The Cancer Genome Atlas (TCGA) database provided the expression profiles of lncRNAs that includes 535 lung adenocarcinoma samples. The coexpression network of lncRNAs and hypoxia-related different expression genes (DEGs) was utilized to select hypoxia-related lncRNAs. The lncRNAs were further screened using univariate Cox regression. In addition, Lasso regression and multivariate Cox regression were used to develop a hypoxia-related lncRNAs signature. A risk score based on the signature was established, and Cox regression was used to test if it was an independent prognostic factor.

Results

Nine prognostic hypoxia-related lncRNAs (LINC01150, AC010980.2, AL606489.1, AL034397.3, LINC00460, LINC02081, FAM83A-AS1, AL365181.2, and AC026355.1) were identified to be significantly different, which made up a hypoxia-related lncRNAs signature. The high-risk group had shorter OS compared with the low-risk group ($P = 3.329e - 09$, log-rank test). A risk score based on the signature was a significantly independent factor for the patients with LUAD (HR = 1.449, 95% CI = 1.312 - 1.602, $P < 0.001$).

Conclusion

The nine hypoxia-related lncRNAs and their signature might be molecular biomarkers and therapeutic targets for the patients with LUAD.

Key words: Hypoxia-related long noncoding RNAs; prognostic; lung adenocarcinoma

Introduction

Lung cancer is the second most common cause of death (> 1.3 million people world-wide every year)^{1,2}. In China, lung cancer incidence in both men and women has increased rapidly in recent years, imposing a great threat to human health³. Non-small cell lung cancer (NSCLC) accounts for approximately 85% of all lung cancers, and lung adenocarcinoma (LUAD) and squamous cell carcinoma (LUSC) are two major histologic subtypes of NSCLC^{4,6}. LUAD is the most common type of NSCLC, which shows distinct genetic drivers and divergent prognostic profiles versus other types of lung cancer⁷⁻⁹. Therefore, further research on the pathogenesis, development and prognosis of LUAD will help to discover new targets and therapeutic drugs.

Evidence has been found that hypoxia is one of the common characteristics in rapidly growing solid tumors. The early phase of solid tumor growth can be divided into two steps in hypoxia conditions: First, malignant cells form

small solid tumors, resulting in tumor hypoxia due to relative lag of vascular growth and rapid proliferation of tumor cells. Second, hypoxia triggers fundamental changes in gene expression, leading to neovascularization and tumor growth and metastasis¹⁰. A large number of findings revealed that hypoxia had multiple functions in occurrence, maintenance, and development of tumors, including LUAD¹¹⁻¹⁴. However, there are no systematic studies about hypoxia-related gene signatures to predict the survival of patients with LUAD.

Long noncoding RNAs (lncRNAs) are a group of noncoding RNAs with more than 200 bp in length with no or limited protein-coding function, which were first discovered in mice in 2002 and lack of specific and complete open reading frame¹⁵. Recently, several studies reported that lncRNAs represent some of the most differentially expressed transcripts between lung tumor and normal lung tissues, highlighting their potential in lung cancer initiation and progression^{16,17}. It has been found that lncRNAs are involved in the development, invasion and metastasis, prognosis, and the chemoresistance of lung cancer via

modulating hypoxia^{13,18}. These studies focused on single or a few lncRNAs for LUAD, while the expression of lncRNAs of The Cancer Genome Atlas (TCGA) datasets was not performed to explore novel biomarkers for forecasting the prognosis of LUAD. Therefore, we aimed to utilize TCGA databases to establish hypoxia-related lncRNAs signatures and seek new biomarkers to predict the prognosis of the patients with lung adenocarcinoma.

Materials and Methods

Datasets and sample extraction

We followed the methods of Weige Zhou, et al¹⁹. The RNA sequencing (RNA-seq) data of LUAD was acquired from The Cancer Genome Atlas (TCGA) database (<https://portal.gdc.cancer.gov/>). Patients with a LUAD diagnosis and intact lncRNA data as well as clinical information qualified as inclusion criteria. The exclusion criteria: patients with follow-up time less than 30 days. Finally, 535 patients with LUAD were absorbed. Furthermore, the clinical data for the patients was retrieved from the TCGA database. The LIMMA package for R software was used to examine the difference expression genes (DEGs) between LUAD samples and normal samples²⁰. Adjusted P value and $|\log \text{fold change}|$ ($|\log \text{FC}|$) were used to evaluate the significance of DEGs, adjusted $P < 0.05$ and $|\log \text{FC}| > 1$ were set as the cutoff criteria. R software was used to analyze the DEGs' heatmap and volcano (version 4.0.4).

Screening of lncRNAs and hypoxia-related genes

The RNAseq dataset was used to obtain the lncRNA profiles, and the log₂ transformation was used to standardize the total RNA expression data. The list of genes related to hypoxia was downloaded using Gene Set Enrichment Analysis (GSEA) (<http://www.gsea-msigdb.org/gsea/msigdb/index.jsp>)²¹. The correlation between lncRNAs and hypoxia-related genes was calculated using Pearson correlation. The square of correlation coefficient $|R^2| > 0.3$ and $P < 0.001$ was considered to be hypoxia-related lncRNAs. Finally, Cytoscape software 3.6.1 was employed to picture co-expression networks²².

Identification of prognostic hypoxia-related lncRNAs

The prognostic value of hypoxia-related lncRNAs was assessed by univariate Cox regression. Hypoxia-related lncRNAs with $P < 0.05$ were included in the absolute minimum shrinkage and selection operator (Lasso) regression in the univariate analysis²³. Then, in order to establish a risk score, the results of Lasso were included into a multivariate Cox model. We found a risk score based on a linear combination of the hypoxia-related lncRNAs expression levels multiplied with a regression coefficient (β): risk score = $\sum_{i=1}^n \beta_i * \text{expression of lncRNA}_i$. Based on the median risk score, the patients were divided into two groups: high-risk and low-risk. Log-rank test was used to compare the survival differences between those two groups.

Development of prognostic model

An independent prognostic model was generated from Cox regression. The nomogram was used to predict patient survival²⁴. Index of agreement (C-index), calibration curves, and receiver operating characteristic (ROC) curves were used to examine model accuracy. To confirm whether the risk score was an independent prognostic indicator, demographic data were entered into a multivariate Cox regression.

Statistical analysis

The Kaplan-Meier method was used to generate the survival curves and the log-rank test was used for comparison. Cox regression and Lasso regression were utilized to estimate the prognostic impact of the hypoxia-related lncRNAs signature and clinicopathological data. The statistical analyses were conducted in R language (version 4.0.4). $P \leq 0.05$ were considered statistically significant.

Results

Identification of DEGs in LUAD and construction of a coexpression network

In TCGA-LUAD, 14142 lncRNAs and 19658 mRNAs were found together. Then, in TCGA-LUAD datasets, an additional 3061 DEGs for lncRNAs (Figure 1A and 1B) and 4661 DEGs for mRNAs (Figure 1C and 1D) were discovered. 64 hypoxia-related genes in total, of which 64 genes (38 down-regulated and 26 up-regulated) were expressed in lung cancer, were retrieved from GSEA (Table 1).

Nine mRNAs (NLU, CRLF1, NQO1, COL3A1, TPBG, KCNK1, SLC16A3, MEST, and MAP2K6) were discovered to be shared by LUAD up-regulated genes and hypoxia up-regulated genes (Figure 2A), while seven genes (ID3, CADM1, IL18R1, LPL, EPAS1, IL6, and RGCC) were discovered to be shared by both LUAD down-regulated genes and hypoxia down (Figure 2B). These hypoxia-related genes (9 up-regulated and 7 down-regulated) and lncRNA coexpression network was constructed to identify the hypoxia-related lncRNAs. Finally, 227 hypoxia-related lncRNAs were selected ($|R^2| > 0.3$ and $P < 0.001$) (Table 2).

Identification of prognostic hypoxia-related lncRNAs signature

According to the results of univariate Cox, 20 hypoxia-related lncRNAs had a prognostic value for the patients with LUAD ($P < 0.05$, Table 3). After Lasso regression, 9 hypoxia-related lncRNAs were discovered (Figure 3, Table 4). Figure 4 showed the coexpression network and Sankey diagram of prognostic hypoxia-related lncRNAs. As shown in Figure 5, we observed a positive correlation between hypoxia-related genes and these lncRNAs. Among them, six lncRNAs (LINC00460, AL365181.2, AL606489.1, LINC02081, AC010980.2, and FAM83A-AS1) were harmful prognostic factors, and three (LINC01150, AC026355.1, and AL034397.3) were favorable prognostic factors (Figure 6, Table 5). These nine lncRNAs were utilized to establish a hypoxia-related lncRNAs signature. The formula of the risk score was as follows: risk score = $(0.32223 * AC010980.2) - (0.42795 * LINC01150) + (0.138432 * AL606489.1) - (0.20391 * AL034397.3) + (0.02445 * LINC00460) + (0.16306 * LINC02081) + (0.02140 * FAM83A-AS1) + (0.05533 * AL365181.2) - (0.30018 * AC026355.1)$.

The prognostic influence of the established signature

The risk score was significantly associated with the overall survival (OS) of patients with LUAD. The high-risk group had shorter OS compared with the low-risk group ($P = 3.329e - 09$, log-rank test) (Figure 7). Cox regression indicated significant prognostic impact of the risk score for the patients with LUAD (Figure 8).

Clinical value of the hypoxia-related lncRNAs signature

Univariate Cox regression revealed that risk score and stage

Table 1 Hypoxia-related genes of lung cancer in Gene Set Enrichment Analysis (GSEA) database

Type	Genes
Up-regulated	ALDH3A1 ANP32A BAG1 CADM1 CALM1 COL15A1 COL3A1 CRLF1 DMD GSTA4 HMOX1 HSPA1B ID3 IGFBP7 KCNK1 LPL MAP2K6 MARCKS MEST NMU NNMT NQO1 RGCC SLC16A3 TPBG TPM1
Down-regulated	ALDH3A1 ANP32A CADM1 CD55 COL15A1 COL3A1 CRLF1 DMD EPAS1 HSPA1B ID3 IER3 IGFBP3 IGFBP7 IL18R1 IL1A IL6 KCNK1 LPL MARCKS MEST MMP1 NMU NNMT NRCAM NT5E PLA2G4A PLAUR RGCC SDC4 SLC16A3 SOD2 STC2 TFPI2 TPBG TPM1 UGCG VEGFC

Table 2 The 227 hypoxia-related lncRNAs of lung adenocarcinoma (LU)

AL133243.2 AC133644.2 LINC02100 AC090739.1 AC010547.2 AC090579.1 AP003119.1 LINC00513 LINC00941 LINC02544 LINC01150 AP001432.1 AC083900.1 FP671120.4 AC010980.2 AC138207.4 AP001528.2 AC009686.2 AC096921.2 AC245014.3 TMPO-AS1 AC018653.3 LINC01936 AP001453.2 AC011442.1 AP001429.1 LINC00467 AC007991.2 AL590666.2 AL162724.1 AP003119.2 AL132780.1 AC083949.1 AC025287.3 AC138393.3 EP300-AS1 AC007546.1 AL033397.1 AC018682.1 MIR3945HG AC011815.1 AC078778.1 AC147067.2 TBX2-AS1 AP005899.1 AC005021.1 AC004832.5 AP002907.1 AL606489.1 LINC00216 AL158166.1 AL024508.2 AL662844.3 AC048341.2 MAL2-AS1 AC025917.1 PITPNA-AS1 AC131971.1 LINC01269 AC067817.2
AC130650.2 AC104984.4 FENDRR AC124319.1 AC079684.1 TBX5-AS1 AC027277.2 AC091057.1 DRAIC AC027228.2 AC007038.2 LINC01607 AC011676.1 AC093278.2 AL645608.8 MIR31HG AP001160.4 AL365356.5 KCNMB2-AS1 DLEU2 MACC1-AS1 AC114488.1 SAP30L-AS1 AC006017.1 AP000695.2 AC026785.3 AC010168.2 AC020913.3 AL034397.3 AC005856.1 CR936218.1 LINC00460 LINC01614 AC079907.1 LUCAT1 AC104695.3 AC015813.1 AC004596.1 AL109914.1 AC090772.3 AC010834.3 HM13-IT1 AL157838.1 LINC01943 AP000866.6 AC010186.3 AC107959.3 AL355488.1 AC021016.2 LINC00942 LINC02081 AC245884.8 AC093110.1 MYO16-AS1 AC004908.2 MYOSLID MBNL1-AS1 AC022784.5 LINC00894 AP000692.1 AC016590.2 AC084117.1 AP001189.3 AC253576.2 PCAT19 AC008115.3 AL049869.3 AC099850.3 AP003170.3 AC108134.3 AC026202.2 AC005519.1 LINC02122 AL136115.2 AC138932.5 MAGI2-AS3 AL031717.1 AL162724.2 LINC00630 AC108727.1 AC004253.1 AC093788.1 AC007014.2 AC012073.1 AF131215.5 AL513327.1 LINC00973 AL512353.1 AC078883.1 MCM3AP-AS1 AC010542.5 AL590723.1 AP002336.2 AC026369.3 AC037198.1 ALMS1-IT1 AC010201.2 HIF1A-AS2 AC004466.3 AC008870.2 MIR193BHG AC127024.4 AC022150.4 LINC00511 SNHG1 LANCL1-AS1 AC079384.1 AC092687.3 FAM83A-AS1 AL365277.1 MIR155HG AC009275.1 AP000786.1 AL133355.1 AC018755.4 AC145207.8 Z82243.1 AL365181.3 FOXP4-AS1 AL022067.1 AC002128.2 AL109761.1 P001033.2 NEAT1 AC063965.1 LINP1 NARF-IT1 LINC01836 AC027288.3 AL365181.2 AL359697.1 AL109614.1 AC026355.1 AL442125.2 AL513365.2 AP000873.2 MIR22HG AC099343.2 DLEU1 ALG13-AS1 AC007249.1 AL683807.1 SMIM25 AC025171.3 AC092115.3 AL928654.1 AL353804.2 AC022211.1 AL354989.1 ANKRD10-IT1 HSPC324 AC002128.1 LINC01355 AC004884.2 AC097641.2 AC026356.1 C20orf197 AF117829.1 AL354719.2 AL035409.1
AC073316.2 AP000695.1 AL391427.1 ELF3-AS1 AL355075.2 PRRT3-AS1 AC008669.1

Table 3 Univariate cox results of hypoxia-related lncRNA based on TCGA-LUAD data

lncRNA	KM	B	SE	HR	HR.95L	HR.95H	p-value
LINC01150	0.0154	-0.50957	0.190111	0.600755	0.41388	0.872007	0.007354
AC010980.2	0.003607	0.282903	0.086847	1.326977	1.11928	1.573214	0.001124
AP001453.2	0.01057	0.057147	0.02399	1.058811	1.010179	1.109785	0.017214
AL606489.1	0.000851	0.182138	0.046646	1.19978	1.094954	1.31464	9.44E-05
FENDRR	0.017209	-0.36127	0.13778	0.696792	0.531894	0.912812	0.00874
AC093278.2	0.035723	-0.21798	0.109936	0.804142	0.648269	0.997493	0.047391
AP000695.2	0.00327	0.335042	0.092891	1.397999	1.1653	1.677166	0.00031
AL034397.3	0.000174	-0.29612	0.108249	0.743699	0.601527	0.919473	0.006228
AC005856.1	0.032812	-0.66277	0.280845	0.515424	0.297241	0.893757	0.01828
LINC00460	0.035424	0.022921	0.00831	1.023186	1.006656	1.039987	0.00581
LINC02081	0.006379	0.113238	0.041578	1.119898	1.032255	1.214983	0.00646
AC099850.3	0.015386	0.045118	0.011985	1.046152	1.021863	1.071018	0.000167
AF131215.5	0.005534	-0.33959	0.133239	0.712064	0.548411	0.924553	0.010812
AC026369.3	0.009687	-0.21522	0.089717	0.806364	0.676338	0.961388	0.016446
FAM83A-AS1	0.009499	0.035622	0.010357	1.036264	1.015441	1.057515	0.000583
AL365181.2	0.048168	0.061963	0.01839	1.063923	1.026259	1.10297	0.000753
AC026355.1	0.000146	-0.28476	0.08401	0.752192	0.637997	0.886826	0.0007
HSPC324	0.009531	-0.62362	0.204554	0.536002	0.358963	0.800356	0.002299
C20orf197	0.001484	-0.11343	0.054822	0.892769	0.801816	0.99404	0.038546
AP000695.1	0.007595	0.223142	0.067199	1.249998	1.095745	1.425965	0.000898

Table 4 LASSO regression coefficients of nine hypoxia-related lncRNAs

lncRNA	Coefficient	HR	HR.95L	HR.95H	p-value
LINC01150	-0.42795	0.651846	0.426544	0.996153	0.047954
AC010980.2	0.322259	1.380243	1.14761	1.660032	0.000622
AL606489.1	0.138432	1.148471	1.008348	1.308066	0.037052
AL034397.3	-0.20391	0.815536	0.670968	0.991252	0.040539
LINC00460	0.024453	1.024755	1.007387	1.042422	0.005049
LINC02081	0.163063	1.177111	1.041602	1.33025	0.008971
FAM83A-AS1	0.021398	1.021629	0.994715	1.049271	0.116194
AL365181.2	0.055328	1.056887	1.014126	1.101453	0.00865
AC026355.1	-0.30018	0.740687	0.631686	0.868496	0.000219

Table 5 Multivariate Cox results of lncRNAs based on TCGA-LUAD data

lncRNA	Coefficient	HR	HR.95L	HR.95H	p-value
LINC01150	-0.42795	0.651846	0.426544	0.996153	0.047954
AC010980.2	0.322259	1.380243	1.14761	1.660032	0.000622
AL606489.1	0.138432	1.148471	1.008348	1.308066	0.037052
AL034397.3	-0.20391	0.815536	0.670968	0.991252	0.040539
LINC00460	0.024453	1.024755	1.007387	1.042422	0.005049
LINC02081	0.163063	1.177111	1.041602	1.33025	0.008971
FAM83A-AS1	0.021398	1.021629	0.994715	1.049271	0.116194
AL365181.2	0.055328	1.056887	1.014126	1.101453	0.00865
AC026355.1	-0.30018	0.740687	0.631686	0.868496	0.000219

Table 6 Clinical characteristics and risk scores of lung adenocarcinoma using univariate cox regression

Variable	B	SE	Z	HR	HR.95L	HR.95H	p-value
Age	-0.00347	0.009599	-0.36166	0.996534	0.977961	1.015461	0.71761
Gender	1.17E-05	0.186484	6.28E-05	1.000012	0.693857	1.441253	0.99995
Stage	0.499618	0.084733	5.896378	1.648091	1.395907	1.945835	3.72E-09
T	0.470248	0.112135	4.193574	1.600391	1.284625	1.993775	2.75E-05
M	0.558732	0.306235	1.824523	1.748454	0.959372	3.186556	0.068073
N	0.580724	0.104857	5.538258	1.787332	1.455295	2.195126	3.05E-08
RiskScore	0.410106	0.049371	8.306557	1.506977	1.367986	1.660089	9.85E-17

Table 7 Clinical characteristics and risk scores of LUAD using multivariate Cox regression

Variable	B	SE	Z	HR	HR.95L	HR.95H	p-value
Age	0.000858	0.00961	0.089306	1.000859	0.982184	1.019888	0.928838
Gender	-0.15414	0.19139	-0.80536	0.857153	0.589043	1.247297	0.42061
Stage	0.498721	0.255521	1.951786	1.646615	0.997912	2.717012	0.050964
T	0.0711	0.127703	0.556762	1.073689	0.835945	1.379047	0.57769
M	-0.59991	0.674064	-0.88999	0.548862	0.146453	2.056967	0.373473
N	0.079815	0.220549	0.361892	1.083087	0.702961	1.668764	0.717432
RiskScore	0.360784	0.052757	6.838604	1.434454	1.29354	1.590719	8.00E-12

Table 8 Clinical influences of risk score signature for TCGA-LUAD data

Clinical	Group	n	Mean	SD	t	P
Age	> 65	168	1.635	1.984	0.729059	0.467
	≤ 65	157	1.493	1.493		
Gender	Femal	164	1.523	1.547	-0.4459	0.656
	Male	161	1.611	1.962		
Stage	I-II	245	1.283	1.113	-3.56944	0.001
	III-IV	80	2.434	2.812		
T	T1-2	281	1.38	1.193	-2.56003	0.014
	T3-4	44	2.755	3.531		
M	M0	304	1.506	1.661	-1.53458	0.140
	M1	21	2.444	2.768		
N	N0	205	1.349	1.654	-2.84986	0.005
	N1-3	120	1.939	1.883		

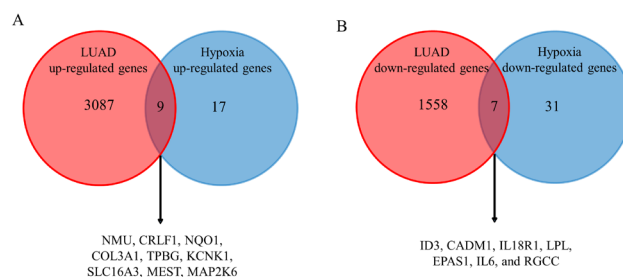


Figure 2 Number of intersecting genes between the (A) up-regulated and (B) down-regulated different expression genes (DEGs) of lung adenocarcinoma and hypoxia-related genes

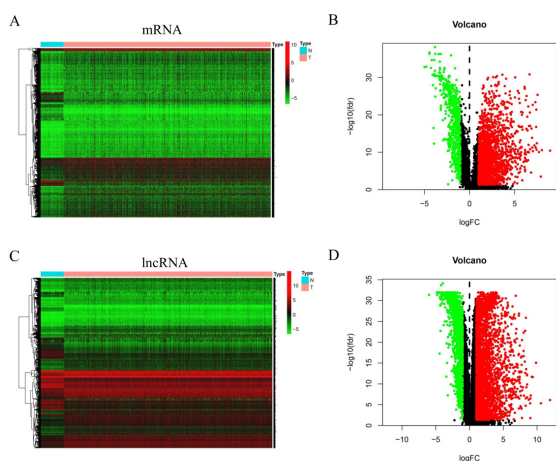


Figure 1 Identification of differentially expressed genes (DEGs) in lung adenocarcinoma (LUAD) from TCGA datasets. (A) The heatmap and (B) volcano plot of DEGs of mRNA in LUAD. (C) The heatmap and (D) volcano plot of DEGs of long noncoding RNAs (lncRNAs) in LUAD. T: tumor; N: control

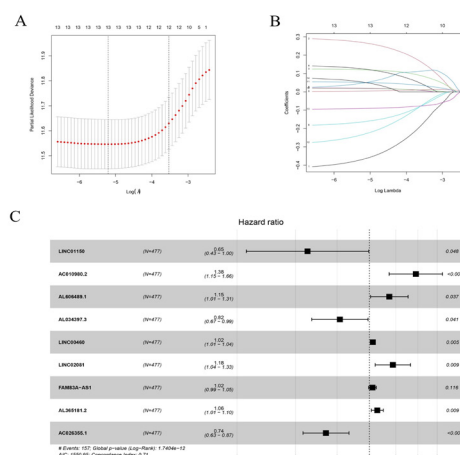


Figure 3 Hypoxia-related lncRNAs selection utilizing Lasso model. (A) Lasso coefficient values of 9 hypoxia-related lncRNAs in lung adenocarcinoma (LUAD). The vertical dashed lines are at the optimal log (lambda) value. (B) Profiles of Lasso coefficients. (C) Forest plot for 9 hypoxia-related lncRNAs in LUAD.

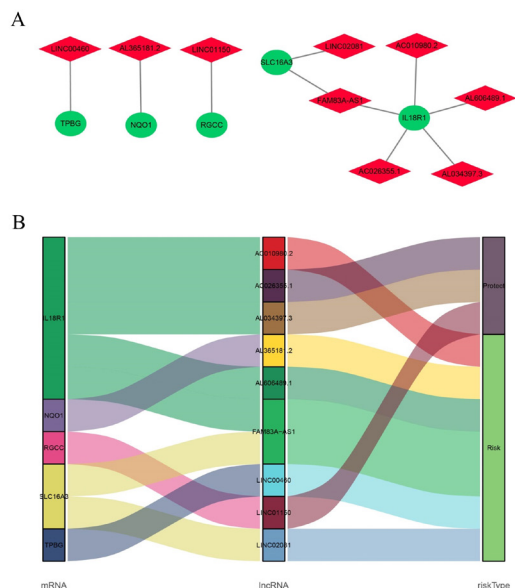


Figure 4 The coexpression network and Sankey diagram of prognostic hypoxia-related lncRNAs. (A) The coexpression network between prognostic lncRNAs and hypoxia-related genes in lung adenocarcinoma. Red diamond nodes represent prognostic lncRNAs, and the sky blue round nodes represent hypoxia-related genes. The coexpression network was visualized using Cytoscape 3.6.1 software. (B) Sankey diagram showed the association between prognostic hypoxia-related lncRNAs, hypoxia-related genes, and risk types

FAM83A-AS1 were independent unfavorable factors. Three lncRNAs (LINC01150, AC026355.1, and AL034397.3) were independent beneficial factors for lung adenocarcinoma

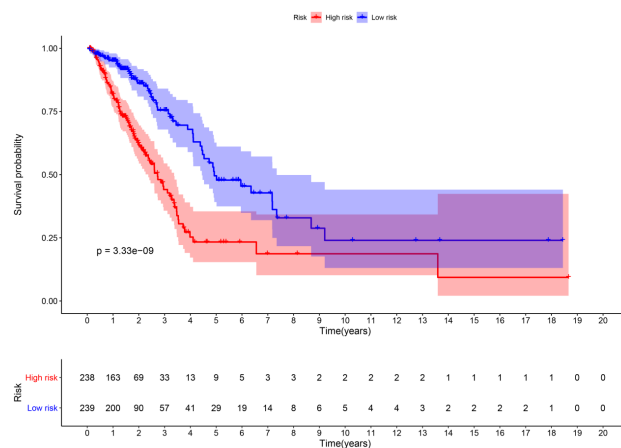


Figure 7 The KM survival curve of risk score based on nine hypoxia-related lncRNAs

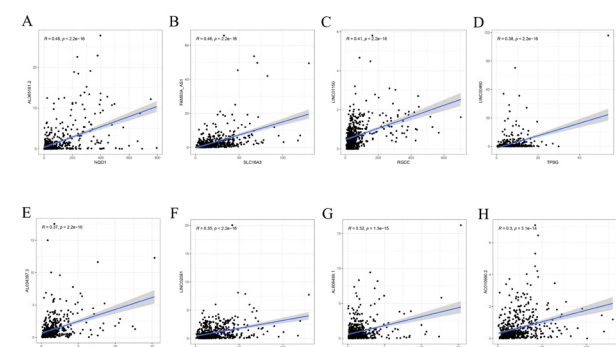


Figure 5 Correlation between hypoxia-related lncRNAs and hypoxia-related genes. (A) AL365181.2 and NQO1. (B) FAM83A-AS1 and SLC16A3. (C) LINC01150 and RGCC. (D) LINC00460 and TPBG. (E) AL034397.3 and IL18R1. (F) LINC02081 and SLC16A3. (G) AL606489.1 and IL18R1. (H) AL606489.1 and IL18R1. (I) AC010980.2 and SLC16A3

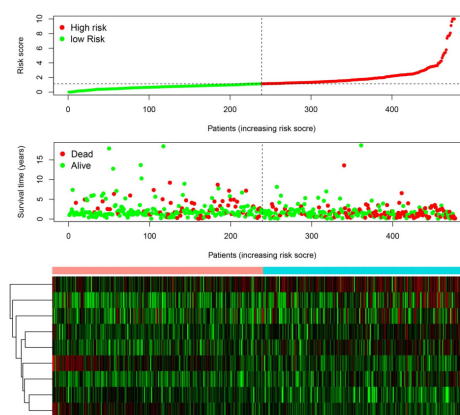


Figure 8 The analysis of hypoxia-related lncRNA signature for patients with lung adenocarcinoma. (A) The risk score between the high-risk group and the low-risk group. (B) The survival time of the patients. (C) Heat map of nine hypoxia-related lncRNAs' expression. The color from green to red reveals a rising tendency from low to high levels

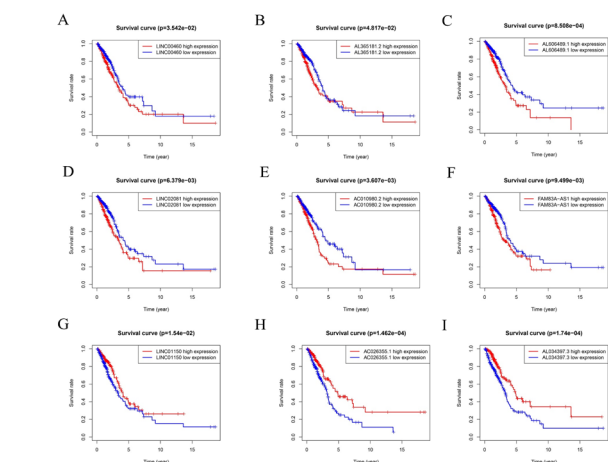


Figure 6 The KM survival curves of nine prognostic hypoxia-related lncRNAs. Six hypoxia-related lncRNAs (LINC00460, AL365181.2, AL606489.1, LINC02081, AC010980.2, and

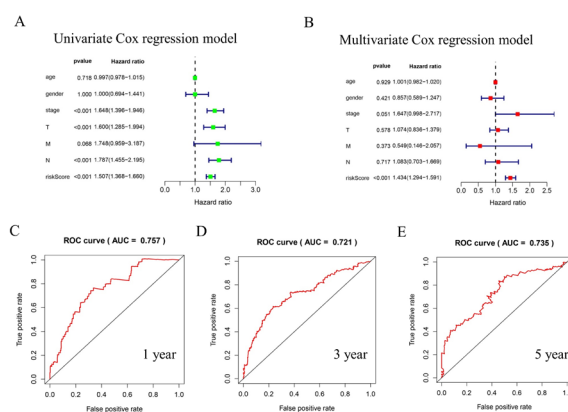


Figure 9 Prognostic indicators based on hypoxia-related lncRNAs showed great predictive performance. The forest plots for univariate (A) and multivariate (B) Cox regression analysis in lung adenocarcinoma. The areas under the ROC curve about (C) 1 year, (D) 3 years, and (E) 5 years

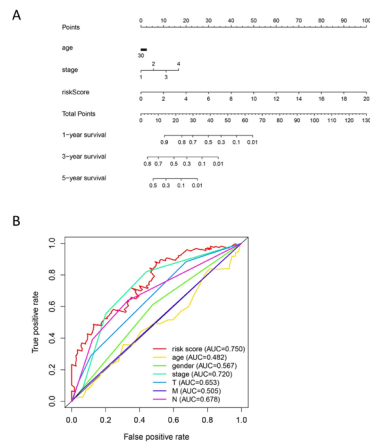


Figure 10 The evaluation of prognostic models based on nine hypoxia-related lncRNAs. (A) The nomogram of 1-year, 3-year or 5-year OS based on risk score, age, and TNM stage. (B) The ROC curves analysis based on risk score and the clinicopathologic parameters

were independent prognostic indicators, and HR of risk score was 1.507 (95% CI: 1.368–1.660, $P < 0.001$, Table 6, Figure 9A). After controlling clinical features, risk score remained an independent prognostic indicator in multivariate analysis (HR = 1.434, 95% CI = 1.294 – 1.591, $P < 0.001$, Table 7, Figure 9B). The areas under the ROC curve corresponding to 1 year, 3 years, and 5 years of survival were 0.757, 0.721, and 0.735, respectively (Figure 9C, 9D and 9E). Risk score, age, and TNM stage were included in the nomogram. As indicated in the nomogram, risk score and TNM stage were the largest contribution to 1-, 3- and 5-year OS of patients with lung adenocarcinoma (Figure 10A). The C-index of the prognostic model was 0.759 (95% CI: 0.710-0.808). The AUC of five-year survival rate showed that risk score (0.750) and stage (0.720) had a certain prediction ability (Figure 10B). The risk scores increased with stage, demonstrating that this hypoxia-related lncRNAs signature may be related to the progression of lung adenocarcinoma (Table 8).

Discussion

Dysregulation of hypoxia has been associated with a number of diseases, including cancer, myocardial infarction, chronic obstructive pulmonary disease, obesity, Coronavirus Disease 2019 (COVID-19), and others^{20,25-27}. lncRNAs played an indispensable role in different aspects of tumorigenesis which were considered a new type of biomarkers in cancer diagnosis, prognosis and therapeutic^{28,29}. However, most researches focused on the function of specific genes involved in hypoxia^{30,31}. There are no systematic studies about hypoxia-related lncRNAs signatures to predict the survival of patients with LUAD. Therefore, it was necessary to establish a hypoxia-related lncRNAs signature to predict the prognosis of patients with LUAD based on the large-scale databases.

In the present study, lncRNAs were screened for hypoxia-relatedness by constructing a coexpression network of lncRNA and genes important for hypoxia. Lasso regression and Cox regression were also used to identify the nine predictive hypoxia-related lncRNAs: LINC00460, AL365181.2, AL606489.1, LINC02081, AC010980.2, FAM83A-AS1, LINC01150, AC026355.1, and AL034397.3. The nine hypoxia-related lncRNAs may act as prognostic molecular markers of prognosis and therapeutic targets for LUAD patients. According to our knowledge, this is the first

study to identify a signature of hypoxia-related lncRNAs that can be used to predict the prognosis of LUAD patients utilizing huge databases.

Four hypoxia-related lncRNAs (LINC00460, FAM83A-AS1, AC026355.1, and AL034397.3) were reported to be associated with LUAD³²⁻³⁵. For example, LINC00460 promotes tumor growth of human lung adenocarcinoma by targeting miR-302c-5p/FOXA1 axis³⁴. LINC00460 can also promote cell migration and invasion through regulating epithelial-mesenchymal transition (EMT) in non-small cell lung cancer^{35,36}. A recent study shows that LINC00460 plays a pivotal role in gefitinib resistance of NSCLC cells by targeting EGFR through sponging miR-769-5p³⁷. Wang et al. revealed that FAM83A-AS1 increased FAM83A expression by enhancing FAM83A pre-mRNA stability and promoted the tumorigenesis of LUAD, revealing that FAM83A-AS1 was a risk factor and possessed oncogenic functions in LUAD³⁸. It was found that FAM83A-AS1 enhances cell migration, invasion and EMT by modulating the miR-150-5p/MMP14 pathway in LUAD³³. For the five remaining hypoxia-related lncRNAs (AL365181.2, AL606489.1, LINC02081, AC010980.2, and LINC01150), there were no studies to report their prognostic roles in cancer, as well as LUAD. Thus, more researches were necessary to explore how these lncRNAs affect the prognosis of patients with LUAD through hypoxia exactly.

The prognosis of LUAD patients was significantly predicted by a signature made up of 9 hypoxia-related lncRNAs. The areas under the ROC curve corresponding to 1 year, 3 years, and 5 years of survival were 0.757, 0.721, and 0.735, respectively. This outcome suggested that the risk score signature had some potential in survival prediction. The signature might be employed as an independent prognostic factor, according to both univariate and multivariate Cox analyses. The C-index, ROC curve, and calibration curve findings showed that the model had superior discrimination and accuracy, suggesting that it might be used as a possible prognostic tool for LUAD patients. These findings are helpful in our investigation of how hypoxia-related lncRNAs function. There were several restrictions on the current investigation. Firstly, the data source of this study is single, and the amount of data included is not large, so the analysis results may have certain deviation. Secondly, our study is a retrospective study, and more prospective studies will be required to prove the prognostic function of hypoxia-related signals. Thirdly, in order to ensure the robustness of the prognostic model, the prognostic model of our established model is required to be further confirmed in other independent cohorts to ensure its robustness. Finally, the functional experiments should be conducted to further indicate the potential molecular mechanisms for predicting the effect of hypoxia-related lncRNAs.

Conclusion

The hypoxia-related lncRNAs signature which was composed of nine hypoxia-related lncRNAs was used to differentiate patients at different risks, and it was a significantly independent factor for the patients with LUAD. Therefore, the nine hypoxia-related lncRNAs and their signature might be molecular biomarkers and therapeutic targets for the patients with LUAD.

Acknowledgements

This work was supported by the Natural Science Foundation

of Shaanxi Province (Grant No. 2024JC-YBQN-0927 and No. 2023-JC-QN-0979), the Key Research and Development Programs of Shaanxi Province (Grant No. 2021SF-322), and the Integration of Basic and Clinical Science Project of School of Basic Medical Sciences, Xi'an Jiaotong University (Grant No. YXJLRH2022034).

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability statement

The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

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