A Hypoxia-Related Signature for Predicting Prognosis, Cellular Processes, Immune Microenvironment and Targeted Compounds in Lung Squamous Cell Carcinoma

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Background: Lung squamous cell carcinoma (LUSC) is a malignant tumour of the lung epithelium. A hypoxic environment can promote tumour cell proliferation and invasion. Therefore, this study aims to explore hypoxia-related genes and construct reliable models to predict the prognosis, cellular processes, immune microenvironment and target compounds of lung squamous carcinoma.

Methods: The transcriptome data and matched clinical information of LUSC were retrieved from The Cancer Genome Atlas (TCGA) database. The GSVA algorithm calculated each LUSC patient’s hypoxia score, and all LUSC patients were divided into the high hypoxia score group and low hypoxia score group. Weighted gene co-expression network analysis (WGCNA) and differential expression analysis were performed to screen out differentially expressed hypoxia-related genes (DE-HRGs) in LUSC microenvironment, and the underlying regulatory mechanism of DE-HRGs in LUSC was explored through Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analyses. Hereafter, we established a prognosis-related genetic signature for DE-HRGs using univariate and multivariate Cox regression analyses. The relationship between gene signature and immune cells was further evaluated. Finally, the Comparative Toxicogenomics Database (CTD) was utilized to predict the targeted drugs for the prognostic genes.

Results: We obtained 376 DE-HRGs. Functional enrichment analysis indicated that the DE-HRGs were involved in the cell cycle-related regulatory processes. Next, we developed and validated 3 HRGs-based prognostic signature for LUSC, including HEMS, GPRIN1, and FAM83A. Risk score is an independent prognostic factor for LUSC. Functional enrichment analysis and immune landscape analysis suggested that the risk scoring system might be involved in altering the immune microenvironment of LUSC patients to influence patient outcomes. Ultimately, a total of 92 potential compounds were predicted for the three prognostic genes.

Conclusion: In summary, we developed and validated a hypoxia-related model for LUSC, reflecting the cellular processes and immune microenvironment characteristics and predicting the prognostic outcomes and targeted compounds.

Keywords: lung squamous cell carcinoma, hypoxia, prognosis, immune, targeted compounds

Introduction

Lung cancer is the major leading cause of tumour-related deaths throughout the world, while lung squamous cell carcinoma (LUSC) as the second most common histological type of lung cancer.1 Each year, almost 1.8 million people are diagnosed with lung cancer worldwide and 400,000 of these die from LUSC.2,3 Due to the lack of early symptoms of LUSC, patients are often diagnosed at an advanced stage.4,5 In the past, there was a lack of effective targeted therapeutic options for patients with LUSC, and therefore the first choice for patients with LUSC remained traditional radiotherapy and chemotherapy.6 In recent years, although immune checkpoint blocker (ICB) therapy has been gradually used for...
LUSC patients, some LUSC patients still experience excessive progression after ICB therapy. Given the limitations of existing therapies such as immunotherapy and targeted therapy, the prognosis of LUSC remains poor.\textsuperscript{7,8} Therefore, there is an urgent need to investigate the underlying biological mechanisms and identify new therapeutic targets.

Hypoxia is one of the essential features of the tumor microenvironment (TME). During cancer progression, tumor cells induce hypoxia through various mechanisms, such as increasing metabolic rate and oxygen consumption leading to endothelial dysfunction or disrupting oxygen delivery due to various effects on blood vessels, forming a chronic hypoxic environment, activating hypoxia-inducible factor HIF signaling, accelerating tumor growth, increasing tumor aggressiveness, and promoting tumor metastasis.\textsuperscript{9–11} Not only that, tumor hypoxia has been reported to contribute to heterogeneous changes, genetic instability, angiogenesis, and resistance to chemoradiotherapy and targeted therapy.\textsuperscript{12–14} Hypoxia stimulates tumor progression and metastasis through physiological and genomic mechanisms and has become a poor prognostic factor for cancer assessment.\textsuperscript{15–17} Several studies have shown that genes associated with hypoxia have prognostic value in a variety of cancers. For example, overexpression of LBH in gliomas under hypoxic conditions is associated with poor prognosis.\textsuperscript{18} In lung adenocarcinoma, NLUCA1 is a transcript that is strongly upregulated in hypoxia and has been shown to have an important prognostic value.\textsuperscript{19} However, the prognostic value of hypoxia-related genes have not been reported in LUSC.

In this study, we performed a systematic and comprehensive analysis of the hypoxia scores of LUSC. We found that a high hypoxia score implied a worse clinical outcome. Meanwhile, functional enrichment analysis and CIBERSORT demonstrated that a risk scoring system based on 3 HRGs was associated with the immune microenvironment. Moreover, we predicted potential compounds for prognostic genes by CTD, providing a theoretical basis for clinical translation efforts of the prognostic signature. In summary, we developed and validated a Hypoxia-related model containing 3 genes for LUSC, reflecting the cellular processes and immune microenvironment characteristics and predicting the prognostic outcomes and targeted compounds.

**Materials and Methods**

**Data Collection**

Transcriptome data in FPKM format and corresponding clinical information of 49 para-cancerous samples and 496 LUSC samples were collected from the TCGA database (https://portal.gdc.cancer.gov/). In this study, 20 LUSC samples with missing survival information were excluded, and the remaining 476 LUSC samples were utilized for hypoxia score estimation, WGCNA, and prognostic model construction. One set of independent microarrays, GSE73403 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE73403), were extracted from the Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/) database. The 69 LUSC patients with complete survival data and clinical information included in this dataset were used in the validation analysis of the prognostic signature. Specific sample information would be described carefully in the corresponding methods as well. The 200 hypoxia-related genes (Supplementary Table 1) were obtained from the HALLMARK_HYPOXIA attributed to the hallmark gene set according to MsigDB (http://www.gsea-msigdb.org/gsea/msigdb/).

**GSVA Calculation of the Hypoxia Score**

Hypoxia scores were analyzed with 200 hypoxia-associated genes and gene set variation analysis (GSVA), which is a non-parametric unsupervised algorithm that estimates changes in pathway activity in a sample population.\textsuperscript{20} In this study, the GSVA scores obtained by GSVA for each sample (n = 476) based on hypoxia-related genes were considered as hypoxia scores, representing the hypoxic status of each sample. LUSC patients were divided into high- (n = 193) and low- (n = 283) hypoxia score groups based on the cut-off value (cut off = 0.04) calculated using the best separation tool in the R package “survminer”. Subsequently, the difference in OS between the two groups was assessed by K-M curves. Moreover, Wilcox test was employed to determine the differences in clinical indicators between the groups. To avoid randomness, we used the “sample ()” function to randomly divide the TCGA-LUSC sample into two groups with the same sample size as the previously divided high-low hypoxia group and compared the difference in OS between the two groups. Also, the procedure was repeated 1000 times, and then a probability of P<0.05 was imputed to determine that the
difference in OS between the high and low hypoxia scoring groups we identified was not a random effect. In addition, the TCGA-LUSC samples were also randomly divided into training and testing sets based on a 5:5 ratio. The hypoxia scores of the samples in the training and testing sets were calculated using the same method. The samples were divided into high and low hypoxia score groups according to the best cutoff value of hypoxia score in each data set, and OS differences were calculated.

Weighted Gene Co-Expression Network Analysis (WGCNA)
Genes with mean expression values (FPKM) >1 in all high- and low-hypoxia group samples (476 LUSC samples in total) were subjected to construction of co-expression networks using R package “WGCNA”. The high- and low-hypoxia scores were served as trait data. The “goodSamplesGenes” function was also used to perform sample clustering to identify and remove outliers. For making the co-expression network contented the distribution of scale-free network, a soft-thresholding power was computed with the pickSoftThreshold function. The dynamic tree cutting method was used to identify different modules, with the minimum number of genes in each module of 30. Subsequently, set a merging threshold of 0.2 to merge similar modules. Subsequently, correlations of modules and traits were calculated by Pearson correlation analysis. |correlation (cor)| > 0.3 and P < 0.05 was considered significantly correlated. The module that possessed the highest correlation coefficient with the trait was considered as the interesting module, and the genes in this module were considered as hub genes.

Identification of DEGs
The “limma” package of R 3.4.3 software was performed to identify the DEGs in LUSC (n = 496) vs para-cancerous (n = 49) samples and high-risk group vs low-risk group. The screening criteria were as follows: |log₂ (fold change) FC| > 1 and adjust P-value < 0.05. The volcano map developed by the R package “ggplot2” was applied to exhibit the distribution of DEGs, while the heatmap generated by the “pheatmap” package displayed the expression pattern of DEGs. Overlapping genes of DEGs and hub genes were recognized as DE-HRGs.

Functional Enrichment Analysis
To further understand the potential biological functions of DE-HRGs and risk score-related DEGs (high-risk group vs low-risk group), we performed GO enrichment analysis, including cellular component (CC), molecular function (MF), and biological process (BP). Then, we performed KEGG pathway analysis. If the P-value is less than 0.05, the result is statistically significant. The functional enrichment analysis was implemented in the R package “clusterProfiler”.

DE-HRGs-Related Gene Signature Construction and Validation
Here, 476 TCGA-LUSC samples were randomly divided into a training set (n = 333) and an internal testing set (n = 143) in a 7:3 ratio. TCGA-training set was used to construct and evaluate the prognostic predictive validity of the prognostic signature. TCGA-internal testing set was employed for internal validation of the prognostic signature. Besides, the GSE3403 dataset was treated as an external validation set to verify the general applicability of the constructed prognostic signature. Feature genes were screened using K-M analysis and Cox analysis based on the identified DE-HRGs. Briefly, we performed univariate Cox analysis incorporating K-M analysis on DE-HRGs and selected variables with p-values less than 0.05 in both K-M analysis and univariate Cox analysis for inclusion in the multivariate Cox analysis with stepwise regression to determine the best variables for the prognostic signature. This study used a risk scoring system to assess the ability of the prognostic signature. The risk score was calculated as follows: Risk score = β1* Exp1 + β2* Exp2 + .... + βi* Expi, where β represents the coefficient value, and Exp represents the level of gene expression. The corresponding risk scores were calculated for the LUSC samples in each dataset (TCGA-training set, TCGA-internal testing set, and external validation set).

Based on the above formula, the risk score of the LUSC samples in each dataset was calculated by R package “predict” to obtain the median value that could classify the samples in each dataset into high-risk and low-risk groups. Subsequently, K-M curve analysis generated by the R package “survminer” was carried out in the TCGA-training set and the internal testing set to assess OS, Event Free Survival (EFS), and Cumulative Incidence in Relapse differences.
between high- and low-risk groups. Prognostic receiver operating characteristic (ROC) curves (“survival ROC” package) were then created at 1, 3, and 5 years. To test the accuracy of the prognostic model, we repeated the above analysis in an external validation set (GSE73403 dataset).

To further determine the necessity of our method for screening the 3 DE-HRGs used to construct the prognostic signature, we randomly selected 3 genes from 376 DE-HRGs to construct the control prognostic signature (random prognostic signature) and assessed their prognostic predictive validity in the TCGA-LUSC dataset using the same method. The coefficients of each gene were calculated using multivariate Cox analysis.

Independent Prognostic Analysis
To investigate whether prognostic characteristics could be independent of other clinical parameters (including sex, age, pathologic stage, pathologic T, pathologic N, and pathologic M), univariate and multivariate Cox regression model analyses were performed in TCGA-training set. P < 0.05 was considered statistically significant. Ultimately, variables with P < 0.05 in the multivariate Cox analysis were identified as independent prognostic factors for LUSC.

CIBERSORT
CIBERSORT is a deconvolution algorithm using a gene expression signature consisting of 547 genes. This algorithm can determine the genetic composition of each cell by calculating the expression level of each gene in each immune cell, thus performing gene expressionism analysis of 22 immune cells. In the present study, we proposed to use CIBERSORT to analyze the composition and proportion of 22 immune infiltrating cells in the high- and low-risk groups of TCGA set. It is important to note that CIBERSORT derives inverse convolution p-values for each sample. Only samples meeting P < 0.05 were included in the follow-up analysis. A total of 438 samples met the above requirement, 225 in the high-risk group and 213 in the low-risk group. Additionally, Spearman correlation analysis examined the relationship between 7 prognostic genes and 22 immune cells. |correlation coefficient (cor)| ≥ 0.3 and P < 0.05 was considered significant.

Prediction of Chemotherapy
Chemotherapy response was predicted for each LUSC patient in TCGA database according to the Genomics of Drug Sensitivity in Cancer (GDSC) database. The GDSC database collects data on the sensitivity and response of a large number of tumor cells to drugs. In the present study, the measurement to assess drug sensitivity was the half maximal inhibitory concentration (IC50). We performed a ridge regression model based on the expression profile of GDSC cancer cell lines using the R package pRRophetic to predict the IC50 values of each drug. We then compared the differences between the low-risk and high-risk groups to determine whether the two groups exhibited different drug sensitivities. For drug selection, we mainly considered the current third-generation chemotherapeutic agents for LUSC, including gemcitabine, paclitaxel, and pemetrexed. Unfortunately, the GDSC database does not record information about pemetrexed.

Comparative Toxicogenomics Database (CTD)
CTD (http://ctdbase.org/) serves as an innovative digital ecosystem capable of linking a given gene to toxicological information about chemicals, genes, phenotypes, diseases and exposures. In this study, we predicted drug candidates for selected prognostic genes in Homo sapiens only.

MetaboAnalyst
MetaboAnalyst (version 5.0; https://www.metaboanalyst.ca/home.xhtml) is a web-based platform for comprehensive analysis of quantitative metabolomic data. This study was based on this platform for functional analysis of compounds.

Statistical Analysis
The R programming language was used for all analyses. All survival analysis was implemented in the R package “survival”. The results of K-M plots were displayed as p-values of Log rank tests. Overlap analysis and Venn diagram were implemented in the Jvenn (http://bioinfo.genotoul.fr/jvenn) online tool. The Wilcoxon assay was used to reveal
differences in the abundance of immune infiltrating cells between high- and low-risk groups. The visualization of the prognostic gene-compound-pathway network was performed by Cytoscape. P < 0.05 was considered to be consistent with statistical significance if not otherwise stated.

**Results**

**Evaluation of the Degree of Hypoxia**

The distribution of hypoxia scores of TCGA-LUSC (n = 476) samples calculated by GSVA based on 200 hypoxia-related gene expression profiles ranged from −0.487 to 0.560 (Supplementary Table 2). The 476 TCGA-LUSC patients were categorized into 193 in the hypoxia score-high group and 283 in the hypoxia score-low group according to the optimal cut-off value of the hypoxia score (cut-off = 0.04, Figure 1A). We initially focused on the prognostic impact of hypoxia. K-M curves suggested that patients with high hypoxia scores were associated with worse OS (P = 0.04, Figure 1B). As shown in Figures 1C–H, the differences in hypoxia scores between patients stratified according to different clinical characteristics were not statistically significant. Random effects analysis showed that after performing 1000 replications, the TCGA-LUSC sample was randomly divided into two groups containing 238 and 193 cases with a significant probability of OS difference (P < 0.05) at 0.048 (Supplementary Figure 1A and B). Furthermore, we randomly divided the 476 LUSC samples into a training set (n = 238) and a testing set (n = 238) according to a 5:5 ratio. And the respective optimal cutoff values were calculated based on the hypoxia scores of all samples in the training and testing sets (cutoff value training set = 0.28, cutoff value testing set = −0.15; Supplementary Figure 1C and D). Based on the respective cutoff values, the clinical outcomes of populations with high- and low-hypoxia scores could be well distinguished between the training and testing sets (P < 0.05; Supplementary Figure 1E and F). This evidence indicated that hypoxia affected the outcome of patients with LUSC.

**Identification of the Closely Connected Module Genes for Hypoxia**

In the R package WGCNA, we constructed co-expression networks using genes with FPKM > 1 in 476 TCGA-LUSC cases with high- and low-hypoxia scores as clinical traits, aiming to assess the modules of interest and hub genes most associated with hypoxia scores. The obvious outliers samples (n = 33; Supplementary Table 3) above the red line were removed by clustering, and β = 4 (scale-free R² = 0.9) was selected to construct a scale-free network (Figure 2A and B). Ultimately, 24 modules were identified according to the average hierarchical clustering and dynamic tree clipping (Figure 2C). Subsequently, we assessed each module’s correlation with two clinical traits (hypoxia score-high and hypoxia score-low). The results indicated that the black module had the highest correlation not only with the high hypoxia score (cor = 0.5, P = 2e-29) but also with the low hypoxia score (cor = −0.5, P = 2e-29) (Figure 2D). Thus, we extracted 3510 genes from the black module for subsequent analysis (Figure 2E; Supplementary Table 4).

**Identification and Functional Enrichment Analysis of DE-HRGs**

In the TCGA-LUSC dataset, 3055 DEGs were identified between LUSC (n = 476) and normal (n= 49) samples, among which 1459 DEGs were upregulated and 1596 DEGs were downregulated (Figure 3A; Supplementary Table 5). The heatmap displayed the expression patterns of the top 50 up- and down-regulated genes. (Figure 3B). By taking the intersection of DEGs and black module genes, we obtained 376 overlapping genes, namely DE-HRGs (Figure 3C; Supplementary Table 6).

To further comprehend the latent functions of 376 DE-HRGs in LUSC, we performed GO and KEGG enrichment analyses. The top 10 components of BP, CC, and MF were illustrated in Figure 3D. Interestingly, we discovered that the DE-HRGs were remarkably enriched in numerous cell cycle-related processes, such as nuclear division (BP), condensed chromosome (CC), and microtubule binding (MF) (Supplementary Table 7). KEGG pathway analysis demonstrated that the DE-HRGs were mainly involved in cell cycle-related pathways (Figure 3E, Supplementary Table 8). Combining the above results, we speculated that the DE-HRGs might participate in the occurrence and development of LUSC by regulating the cell cycle process.
Construction and Verification of the Prognostic Model

In the TCGA-training set, univariate Cox regression with K-M survival analyses were performed on 376 DE-HRGs, and 6 genes (HELIS, GPRIN1, TARBP1, CST3, FAM83A, and LY6D) related to the OS of LUSC were selected ($P < 0.05$; Figure 4A; Supplementary Table 9). Subsequently, stepwise multivariate Cox regression analysis was used to construct a prognostic model for LUSC, which was composed of HELLS, GPRIN1, and FAM83A (Figure 4B; Supplementary Table 10). The risk score of each LUSC patient in the TCGA-training set was calculated according to the prognosis model: Risk score = $-0.10399 \times$ expression of HELLS + $0.07685 \times$ expression of GPRIN1 + $0.00738 \times$ expression of

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**Figure 1** Hypoxia affected the outcome of TCGA-LUSC patients. (A) 476 LUSC patients were divided into two groups: hypoxia score-high group (n = 193) and hypoxia score-low group (n = 283), according to their optimal cut-off value (0.04). (B) Analysis of overall survival (OS) based on hypoxia score-high group, n = 193; hypoxia score-low group, n = 283. Wilcoxon analysis was performed to evaluate the differences of hypoxia score in different clinical characteristics. (C) Distribution of hypoxia score of patients younger than 65 and those older than 65 years of age. (D) Hypoxia score of different gender. (E) Hypoxia score of different stages. (F) Distribution of hypoxia score of patients with different T stage. (G) Distribution of hypoxia score of patients with (N1-3) or without (N0) lymph node metastasis. (H) Distribution of hypoxia score of patients with (M1) or without (M0) distant metastasis.
FAM83A. According to the median value of the risk score, the LUSC cases in the training set were assigned into a low-risk group and a high-risk group.

We performed a survival analysis of 333 LUSC patients from the TCGA-training set and found that OS (P = 0.013; Figure 5A) and EFS (P = 0.009; Figure 5B) were significantly lower in the high-risk group than in the low-risk group, and conversely, Cumulative Incidence in Relapse was significantly lower in the low-risk group than in the high-risk group (P = 0.018; Figure 5C). The AUC for 1-, 3-, and 5-year survival rate were 0.614, 0.660, and 0.650, respectively (Figure 5D). As the risk score increased, the mortality of patients also increased. Compared with the low-risk group, the expression levels of FAM83A and GPRIN1 in the high-risk group were higher, while the expression level of HELLS was lower (Supplementary Figure 2A). Furthermore, we adopt analogous methods to further evaluate the prognostic performance of the 3 gene features in TCGA-internal testing set and GEO-external validation set. It was consistent with the result of analysis in the TCGA-training set, the patients in the high-risk group were significantly associated with a poorer survival rate compared to the patients in the low-risk group (P < 0.05, Figure 5E and 5I).

Similarly, in the TCGA-internal testing set, patients in the high-risk group had a significantly lower EFS (P = 0.01; Figure 5F) than in the low-risk; the difference in Cumulative Incidence in Relapse between the two groups was not significant, but overall, patients in the high-risk group had a slightly higher Cumulative Incidence in Relapse than in the low-risk group (P = 0.9; Figure 5G). Unfortunately, information on patients’ EFS and Cumulative Incidence in Relapse was not recorded in the external validation set. The AUC for survival rate was 0.686 at 1 year, 0.614 at 3 years, and 0.604 at 5 years in TCGA-internal testing set (Figure 5H); 0.565 at 1 year, 0.632 at 3 years, and 0.647 at 5 years in the GEO-external validation set (Figure 5J). The Supplementary Figure 2B and 2C were displayed the distribution of the risk score, survival status, and 3 gene expression profiles between the two risk groups in TCGA-internal testing set and GEO-external validation set, respectively. The above evidence suggests that our prognostic characteristics based on the 3 DE-HRGs constructed by Cox regression analysis are effective in predicting patient prognosis.
Independent Prognostic Analysis of Risk Model

Univariate and multivariate Cox regression analyses were conducted in the training set to ascertain the independence of the 3-gene signature in clinical application. It was exhibited that pathologic stage (HR = 1.266, 95% CI = 1.071–1.495, \( P < 0.001 \)), pathologic T (HR = 1.307, 95% CI = 1.089–1.568, \( P = 0.004 \)), pathologic M (HR = 1.277, 95% CI = 1.054–1.547, \( P = 0.012 \)), and risk score (HR = 2.263, 95% CI = 1.566–3.271, \( P < 0.001 \)) were obviously connected with OS in the univariable Cox analysis (Figure 6A; Supplementary Table 11). The results of multivariate Cox regression analysis displayed that risk score (HR = 2.235, 95% CI = 1.635–3.392, \( P < 0.001 \)) and pathologic M (HR = 1.280, 95% CI = 1.051–1.558, \( P = 0.014 \)) were the independent prognostic indicators for LUSC (Figure 6B; Supplementary Table 12).

Revealing the Potential Functions of Risk Score-Related DEGs

To further understand the potential mechanisms by which risk score predicted the outcome of LUSC patients, we first identified a total of 14 up-regulated genes and 9 down-regulated genes between high- and low-risk groups based on the R package limma (Figure 7A and B; Supplementary Table 13), and these DEGs were considered as risk score-associated DEGs. Subsequently, GO analysis was used to reveal the potential functions of risk score-related DEGs (Supplementary Table 14). Figure 7C was illustrated the top 10 enriched terms in the three categories of the GO system. We focused on the results of the GO analysis and found that immune response (“leukocyte aggregation” and “leukocyte migration involved in inflammatory response”) related terms were significantly enriched. KEGG analysis indicated that the IL-17...
A signaling pathway was the most enriched pathway (Figure 7D; Supplementary Table 15). This evidence suggested that the risk scoring system might be mediating the immune microenvironment of LUSC.

Furthermore, interestingly, we found that the results of the GO (Supplementary Table 16) and KEGG (Supplementary Table 17) enrichment analyses of the 39 DEGs (Supplementary Table 18) between the high and low hypoxia score groups were strikingly similar to those of the risk score-related DEGs (Supplementary Figure 3A–D). Subsequently, Sankey plots showed that most of the samples in the high hypoxia score group were also classified in the high-risk group (Supplementary Figure 4).

**Risk Scoring System-Mediated Alterations in the Immune Landscape of LUSC**

Inspired by the above results, we explored the effect of a risk scoring system constructed based on 3 HRGs on the immune microenvironment of LUSC. The abundance of 22 immune cell subtypes in each LUSC sample for the high (n = 225) low (n = 213) risk group was shown in Figure 8A. Further, the violin plot revealed that patients in high-risk group had a significant decrease in the fraction of “B cells naive”, “T cells CD8”, “T cells follicular helper”, and “Macrophages M1”, while a remarkably increased in the proportion of “T cells CD4 memory resting”, “Macrophages M0”, “Dendritic cells activated”, and “Neutrophils” (Figure 8B). Subsequently, Spearman correlation analysis pointed out that the prognostic genes GPRIN1 (cor = −0.301, P = 1.28E-10) and FAM83A (cor = 0.356, P = 1.52E-14) were significantly and weakly correlated with Neutrophils (Figure 8C; Supplementary Table 19).
Prediction of Potential Compounds Targeting Prognostic Genes

Next, to improve the ability of the clinical utility of the risk scoring system, we identified the sensitivity of patients in the high- and low-risk groups to current third-generation chemotherapeutic agents for LUSC using the GDSC database. The results showed significantly lower IC50 values for gemcitabine in the low-risk group, indicating that patients with LUSC exhibiting low risk scores were more sensitive to gemcitabine (Figure 9A). However, the IC50 values of paclitaxel did not differ significantly between the high- and low-risk groups (Figure 9B). Due to limited database information, our risk scoring system seems to be currently associated with Gemcitabine only. To facilitate the development of new LUSC drugs, we used the CTD database to predict compounds capable of targeting prognostic genes. Ultimately, a total of 92 compounds were predicted, of which HELLS had 81 acting relationships with 65 compounds; 21 compounds constituted 23 interacting relationships with GPRIN1; and FAM83A had 39 pairs of relationships with 25 compounds. Benzo(a)pyrene, Estradiol, Tobacco Smoke Pollution, and Valproic Acid were the common compounds for the 3 prognostic genes, but their interaction relationships were not identical. In detail, Benzo(a)pyrene led to increased HELLS and GPRIN1 mRNA expression, increased GPRIN1 5' UTR methylation, decreased FAM83A exon methylation, and also affected FAM83A promoter methylation; Estradiol resulted in increased HELLS mRNA expression, and GPRIN1 and FAM83A mRNA expression was due to co-treatment of Estradiol with TGFβ1 protein; Tobacco Smoke Pollution contributed to decreased HELLS mRNA expression but led to increased GPRIN1 and FAM83A mRNA expression and also affected FAM83A protein expression; Valproic Acid was able to affect HELLS mRNA expression and lead to increased methylation of GPRIN1 and FAM83A genes. Detailed information on the above results was presented in Supplementary Table 20. Further, we performed a functional network analysis of these compounds via the MetaboAnalyst 5.0 online network. From Figure 9C, Aflatoxin B1, which targeted FAM83A, and Bazedoxifene and 7,8-Dihydro-7,8-dihydroxybenzo(a)pyrene 9,10-oxide, which targeted HELLS, were all involved in the Metabolism of xenobiotics by cytochrome P450; Testosterone, Progesterone, and Estradiol, which could target HELLS, were associated with Steroid hormone biosynthesis; Steroid biosynthesis and Glyoxylate and dicarboxylate metabolism were associated with Calcitriol and Oxygen, which can target HELLS, respectively; Tretinoin, which would target FAM83A, was involved in Retinol metabolism.
Lung squamous cell carcinoma (LUSC) is one of the leading causes of cancer-related death in the world, and the development of LUSC is a complicated process influenced by various factors.\(^1\)\(^-\)\(^3\) Although molecularly targeted drugs have made great progress in the treatment of LUAD due to the presence of EGFR mutations and ALK fusions.\(^30\) Unfortunately, these phenomena are not present in LUSC,\(^31\)\(^-\)\(^33\) resulting in the difficulty of many effective target drugs to exert therapeutic effects in LUSC, so LUSC patients have been lacking effective targeted therapeutic options.

Hypoxia is an important factor in the development and progression of malignant tumours, and the hypoxic microenvironment of tumour tissue can exacerbate disease progression and metastasis through physiological and genomic mechanisms.\(^15\) Meanwhile, several studies have demonstrated that hypoxia-related genes can be used in the prognosis of other types of cancer, such as melanoma,\(^34\) breast cancer,\(^35\) and hepatocellular carcinoma.\(^36\) However, systematic studies on the prognostic role of hypoxia-related genes in LUSC are still scarce. Therefore, there is urgent to explore biomarkers and therapeutic targets potentially associated with the hypoxic tumor environment to develop better individualized treatment plans and improve LUSC patients’ prognosis.

To confirm the correlation between hypoxia and LUSC, we verify hypoxia-related genes’ prognostic value, the GSVA algorithm calculated gene expression profiles and hypoxia scores for all LUSC samples. Then patients were divided into high and low hypoxia groups based on hypoxia scores. Patients with higher hypoxia scores had poorer survival rates, and we identified 376 DE-HRGs as significantly hypoxia-related genes correlated with LUSC by intersecting DEGs and critical modules. Hypoxic activity in the tumour microenvironment is complex and previous studies have demonstrated that the hypoxic microenvironment of tumours can promote tumour cell proliferation.\(^37\)\(^-\)\(^39\) In this study, we found that
these DE-HRGs were significantly enriched by GO enrichment analysis in several cell cycle-related processes, including nuclear division, chromosome condensation and microtubule binding. KEGG pathway analysis indicated that DE-HRGs were mainly involved in cell cycle-related pathways. Combining the above results, we speculated that the hypoxia might participate in the occurrence and development of LUSC by regulating the cell cycle process.

In this study, we identified three genes (HELLS, GPRIN1 and FAM83A) associated with hypoxia that are closely associated with LUSC. The DNA helicase HELLs, which is central to tumour proliferation and progression, has been shown to be expressed in many tumours and plays a relevant role in the transcriptional and genomic stabilisation of cancer by transcriptionally regulating genes involved in cleavage furrow regulation to promote tumour cell division. Furthermore, it has been shown that G protein-regulated neuronal growth inducer 1 (Gprin1) may promote lung cancer proliferation and migration by affecting the epithelial-mesenchymal transition of lung cancer cells and may be an effective target for the treatment of lung cancer. As a member of sequence similarity 83, FAM83A has also been found to promote the progression of various cancers such as breast, cervical and lung cancers by activating signalling pathways such as epidermal growth factor receptor EGFR/PI3K/AKT. However, the above three signature genes have rarely been studied in the context of hypoxia and immune binding. To further understand the potential mechanisms...
Figure 8 Immune infiltrating cells profile in LUSC and correlation analysis. (A) Barplot showing the proportion of 22 kinds of immune infiltrating cells in the high- and low-risk LUSC samples. Column names of plot were sample ID. (B) Violin plot of 22 immune cells content in the high-risk and low-risk group. Red color represented high-risk group while blue color represented low-risk group. Differential immune cell type expression was observed between the high- and low-risk groups. (C) Heatmap showing the correlation between 22 kinds of immune cells and prognostic genes. The shade of each tiny color box represented corresponding correlation value between two features, and Spearman coefficient was used for significance test. ***P < 0.001.
underlying the outcome of hypoxic LUSC patients, we performed GO and KEGG enrichment analysis on DEGs. The results both showed that DEGs were involved in the immune response. This analysis reveals an association between hypoxia and the immune microenvironment of LUSC and provides insight into potential therapeutic mechanisms that may underlie individualised treatment of LUSC patients.

To date, many studies have confirmed that most tumors have hypoxic zones and that the secondary formation of hostile metabolic and physical microenvironments leads to an imbalance of positive and negative regulators of processes such as activation and dysregulation of angiogenesis, demyelination and inflammation. Thus the development of abnormal vascular and hypoxic microenvironments promotes abnormal angiogenesis, demyelination and inflammation, all of which contribute to tumor development and resistance to therapy. It has been reported that inflammation may promote cellular transformation and that approximately 25% of cancer cases are associated with chronic inflammation of different origins. Not only that, the hypoxic state of tumor tissue is important in promoting immunosuppression of tumors. Tumor hypoxic areas can recruit immunosuppressive cells, such as tumor-associated macrophages (TAM), and can inhibit the activation of immune cells such as T cells. These reports coincide with our analysis, where we found significantly higher rates of “T-cell CD4 memory quiescence”, “macrophage M0”, “dendritic cell activation” and “neutrophils” in hypoxic high-risk patients, and a strong correlation between the model genes GPRIN1 and FAM83A for neutrophils. Given the central role of hypoxia in regulating LUSC progression and immunosuppression, it may provide translational value for the clinical management of LUSC patients. Finally, after determining the relationship between hypoxia and the immune microenvironment of LUSC, it may be necessary to provide potential therapeutic agents. We used the GDSC database to screen hypoxic LUSC for sensitive and selective drugs. The screening results showed that gemcitabine showed potential sensitivity and selectivity for LUSC with low risk of hypoxia. To facilitate the development of new LUSC drugs, we used the CTD database to predict compounds that could target prognostic genes. Ultimately, a total of 92 compounds were predicted, with Benzo(a)pyrene, Estradiol, Tobacco Smoke Pollution, and Valproic Acid as common compounds for these 3 prognostic genes, which provides new insights into future gene targeting therapy for hypoxic LUSC patients.

Fortunately, this is the first comprehensive study to develop and validate hypoxia-related gene signatures to predict the prognosis of LUSC. Our research aimed to explore the hypoxia-related genes and construct reliable models for predicting prognosis, cellular processes, immune microenvironment, and targeted compounds in lung squamous cell carcinoma. Our research provides new insights into the clinical management of patients with LUSC.

Figure 9 Drug sensitivity analysis and potential compound prediction. (A) IC50 values of gemcitabine in high- and low-risk groups. (B) IC50 values of paclitaxel in high- and low-risk groups. (C) The prognostic gene-compound-pathway network consisting of 15 nodes and 18 edges. Red rounded rectangles represent functional pathways, yellow diamonds indicate compounds, and light blue ovals indicate prognostic genes.
Data Sharing Statement
The datasets generated during the current study are available in the TCGA database (https://portal.gdc.cancer.gov/) and GEO database (https://www.ncbi.nlm.nih.gov/geo/).

Ethics Approval and Consent to Participate
The Ethics Committee of Affiliated Hospital of Nantong University has granted exemptions from approval for research related to the use of such public databases.

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Disclosure
The authors report no conflicts of interest for this work and declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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