

### Development and validation of a DNA damage repair-related genebased prediction model for the prognosis of lung adenocarcinoma

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**Background:** Lung cancer is the leading cause of morbidity and mortality among all cancer types, with lung adenocarcinoma (LUAD) being the most prevalent subtype. DNA damage repair (DDR)-related genes are closely associated with cancer progression and treatment, with emerging evidence highlighting their correlation with tumor development. However, the relationship between LUAD prognosis and DDR-related genes remains unclear.

**Methods:** RNA sequencing (RNA-seq) data and clinical information were obtained from The Cancer Genome Atlas (TCGA) database. The GSE31210 dataset, utilized for external validation, was retrieved from the Gene Expression Omnibus (GEO) database. Differentially expressed DDR genes were identified, and a DDR-related prognostic model was established and validated using Kaplan-Meier (KM) survival analysis, time-dependent receiver operating characteristic (ROC) curves, gene set enrichment analysis (GSEA), tumor mutational burden (TMB) analysis, and immune cell infiltration. A P value of less than 0.05 was considered statistically significant.

**Results:** A total of 514 patients with LUAD from TCGA database were divided into distinct subtypes to characterize the diversity within the DDR pathway. DDR-activated and DDR-suppressed subgroups showed distinct clinical characteristics, molecular characteristics, and immune profiles. Nine genes were identified as hub DDR-related genes, including *CASP14*, *DKK1*, *ECT2*, *FLNC*, *HMMR*, *IGFBP1*, *KRT6A*, *TYMS*, and *FCER2*. By using the expression levels of these selected genes, the corresponding risk scores for each sample was predicted. In the training group, KM survival analysis revealed that the high-risk group exhibited significantly diminished overall survival (OS) [hazard ratio (HR) =3.341, P=1.38e-08]. The corresponding area under the curve (AUC) values for the 1-year follow-up periods was 0.767, respectively. Upon validation in the external cohort, patients with higher risk scores manifested significantly reduced OS (HR =2.372, P=1.87e-03). The AUC values of the ROC curves for the 1-year OS in the validation cohort was 0.87, respectively. Moreover, advanced DDR risk score was correlated with increased TMB scores, a heightened frequency of *TP53* mutations, an increased abundance of cancer-testicular antigens (CTAs), and a lower tumor immune dysfunction and exclusion (TIDE) score in patients with LUAD (P<0.05).

**Conclusions:** A nine-gene risk signature associated with DDR in LUAD was effectively developed, demonstrating its potential as a robust and reliable classification tool for clinical practice. This model exhibited the capability to accurately predict the prognosis and survival outcomes of LUAD patients.

Keywords: DNA damage repair (DDR); lung adenocarcinoma (LUAD); prognosis; risk score model; tumor

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### Introduction

Lung cancer, particularly non-small cell lung carcinoma (NSCLC), is the leading cause of morbidity and mortality among all cancer types (1,2). NSCLC is commonly classified into histological subtypes, with lung adenocarcinoma (LUAD) being the most prevalent subtype (3). The development of LUAD is influenced by various factors, including smoking, alcohol consumption, and metabolic disorders. Despite significant advancements in multimodal treatment approaches, such as immunotherapy, radiotherapy, and noninvasive surgical resection, the outcomes for patients with lung cancer remain unsatisfactory, with a 5-year relative overall survival (OS) rate of approximately 18% (3,4). This can be attributed to the limitations of the traditional histological classification of LUAD, given its high heterogeneity and complexity (5,6). Additionally, the existing staging system fails to accurately predict the prognosis of lung cancer, which can result in some patients with early-stage disease failing to receive appropriate adjuvant therapy after surgery,

#### Highlight box

#### Key findings

• A nine-gene risk signature associated with DNA damage repair (DDR) in lung adenocarcinoma (LUAD) was effectively developed, demonstrating its potential as a robust and reliable classification tool for clinical practice.

#### What is known and what is new?

- Lung cancer is the leading cause of morbidity and mortality among all cancer types, with LUAD being the most prevalent subtype. DDR-related genes are closely associated with cancer progression and treatment, with emerging evidence highlighting their correlation with tumor development. However, the relationship between LUAD prognosis and DDR-related genes remains unclear.
- We successfully developed a nine-gene risk signature associated with DDR in LUAD, demonstrating its potential as an effective and stable classification tool for clinical practice.

### What is the implication, and what should change now?

• This study has significant implications for LUAD patient care and highlights the importance of considering DDR subtypes in personalized treatment approaches. leading to cancer recurrence or metastasis (7,8). Therefore, there is a crucial need to identify more effective prognostic indicators for patients with LUAD.

DNA damage repair (DDR) genes play a pivotal role in maintaining the stability of the human genome, while the loss of DDR function can contribute to the initiation and progression of cancer (9,10). As a result, there is a growing appreciation for treatment strategies that target aberrant DDR function. One such example is poly (ADPribose) polymerase (PARP), a nuclear enzyme involved in recognizing DNA damage, which has emerged as a therapeutic target for cancer treatment (11). DDR genes can be categorized into specific functional pathways based on their roles in DNA damage response (12). This categorization has provided valuable insights into underlying mechanisms and therapeutic analyses. DDRrelated genes are closely associated with cancer progression and treatment, with emerging evidence highlighting their correlation with tumor development (13,14). However, the prognostic significance of these genes in LUAD has not been thoroughly investigated. In this study, we conducted a comprehensive evaluation and developed a novel signature and nomogram based on DDR genes to predict the outcomes of LUAD. Despite this progress, there remains a limited understanding of the dysregulation and heterogeneity of DDR genes in LUAD, particularly in terms of transcriptomic and proteomic analysis. Recent studies by Gu et al. demonstrated the prognostic value of a 15-feature gene signature in improving outcomes for patients with LUAD (15). Wu et al. also investigated the survival benefits associated with high tumor mutational burden (TMB) or DDR gene mutations in patients with LUAD with high stromal or immune scores (16). These findings underscore the potential of DDR genes as not only oncogenes but also promising biomarkers for prognosis prediction and treatment in patients with LUAD.

In this study, we analyzed a dataset of gene expression in LUAD obtained from The Cancer Genome Atlas (TCGA) and identified nine DDR genes through screening. The identified DDR gene signature proved to be a reliable predictor of survival prognosis in patients with LUAD. Moreover, our DDR subtype signature demonstrated its potential as a robust and clinically applicable classification

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tool in LUAD. These DDR genes hold additional promise as biomarkers for guiding immunotherapies in patients with LUAD. We present this article in accordance with the TRIPOD reporting checklist (available at https://jtd. amegroups.com/article/view/10.21037/jtd-23-1746/rc).

### Methods

### Data collection

RNA sequencing (RNA-seq) data along with clinical information including age, clinical stage, mutations, copy number variations, days to death, vital status, and more, were obtained from TCGA database (https://portal.gdc. cancer.gov/). Samples lacking clinical information were excluded from the analysis. Additionally, the GSE13213 dataset retrieved containing 117 LUAD samples from the Gene Expression Omnibus (GEO) database (https://www. ncbi.nlm.nih.gov/geo/) was utilized for external validation. To analyze the expression levels, count per million (CPM) read values were calculated using the edgeR software package (17). Ethics approval was deemed unnecessary for this phase of the study given that TCGA and GEO databases are publicly accessible resources. The study was conducted in accordance with the Declaration of Helsinki (revised in 2013).

### Identification of differentially expressed DDR genes

Initially, the list of DDR-associated genes was retrieved from the Molecular Signatures Database (https://www.gseamsigdb.org/gsea/msigdb). Ensemble IDs were transformed into gene symbols, selecting median values in cases where a gene had multiple symbols. To assess the expression levels of DDR genes in each sample, we conducted cluster analysis using the ConsensusClusterPlus package. This analysis facilitated the grouping of samples based on the expression patterns exhibited by DDR genes (18).

### Establishment and validation of a prognostic model

The cancer samples of messenger RNA (mRNA)-seq in LUAD were randomly divided into two equal groups: a training group and a test group. A regression model was built according to the training group, and the test group and the total samples were used to verify the model results. The glmnet R package (The R Foundation for Statistical Computing, Vienna, Austria) was used to perform least absolute shrinkage and selection operator (LASSO) Cox regression for the purpose of identifying prognostic genes. To prevent overfitting, 10-fold cross-validation was employed to determine the penalized regularization parameter  $\lambda$  in the model. Based on the constructed model, the risk score for each LUAD sample was calculated. Additionally, the survival R package was used to conduct univariate Cox regression analysis of OS (19). Using the median of the risk score, we divided the patients with LUAD into two groups: a high-risk and low-risk group. Subsequently, receiver operating characteristic (ROC) curve analysis was conducted separately in the training dataset, testing dataset, and the entire dataset to assess the accuracy of the DDR signature. The differences between the highrisk and low-risk groups were evaluated through Kaplan-Meier (KM) curve analysis and the log-rank test. A P value of less than 0.05 was considered statistically different.

### Functional and pathway enrichment analysis

Gene set enrichment analysis (GSEA) is a computational method used to identify sets of genes that are statistically enriched for a specific observable variable. In this study, we conducted GSEA using gene expression data obtained from TCGA and the Gene Oncology (GO) or Kyoto Encyclopedia of Genes and Genomes (KEGG) databases (20,21). The objective was to determine if particular gene sets exhibited enrichment based on their expression levels. Additionally, GSEA was performed using the gsva R package (22). We analyzed the differentially regulated genes between the high-risk and low-risk groups. A P value of less than 0.05 was considered statistically significant.

## Relationship between the DDR signature and immune infiltration

The wilcox.test function in R was used to compare the differential expression of immune checkpoint genes between different groups of patients with LUAD. To visualize the mutational profiles of the low-risk and high-risk groups, the maftools package in R was employed (23). Cancer-testicular antigen (CTA) levels were obtained from the CTdatabase (http://www.cta.lncc.br/) (24), and the number of CTAs in each patient was calculated. Differential analysis of CTA numbers was performed using the wilcox.test function in R. Additionally, the response of patients with LUAD to immune checkpoint blockade (ICB) was predicted on pretreatment genomics using the tumor immune dysfunction and exclusion (TIDE) program (http://tide.dfci.



**Figure 1** Clustering result plot. Group 1: cluster 1 (n=227, 44.2% of all LUAD), designated as the DDR-suppressed subgroup; group 2: cluster 2 (n=287, 55.8% of all LUAD), designated as the DDR-activated subgroup. LUAD, lung adenocarcinoma; DDR, DNA damage repair.

harvard.edu/). A P value of less than 0.05 was considered statistically significant.

### Statistical analysis

The RNA-seq data, along with clinical information pertaining to LUAD samples, were acquired from the Genomic Data Commons Portal or the GEO database. Statistical analyses were performed using GraphPad Prism 9.0 and R packages. Differences between the high-risk and low-risk groups were evaluated through KM curve analysis and the log-rank test. ROC curve analysis was separately conducted on the training, testing, and entire datasets to assess the accuracy of the DDR signature. All statistical procedures were carried out using R software (v4.0; The R Foundation for Statistical Computing). The Student's *t*-test and paired *t*-test were applied for independent and paired groups, respectively. Continuous variables were presented as mean  $\pm$  standard deviation. Statistical significance was established for a P value below 0.05.

### Results

### DDR gene alteration profiles in LUAD

To investigate the heterogeneity of DDR gene expression in

LUAD, a total of 514 patients with LUAD were included in the analysis. Based on the expression profiles of 429 DDR genes, these patients were divided into distinct subtypes to characterize the diversity within the DDR pathway. Through consensus clustering and consideration of clinical features, two DDR subgroups were identified (*Figure 1*). Cluster 1 consisted of 227 patients, accounting for 44.2% of all LUAD cases and was designated as the DDR-suppressed subgroup, exhibiting comparative downregulation of DDR genes. In contrast, cluster 2 comprised 287 patients, representing 55.8% of all LUAD cases and was designated as the DDR-activated subgroup, showing significant upregulation of most DDR-related genes.

## DDR gene-based subtypes exhibited distinct clinical characteristics

The two subgroups exhibited contrasting clinical outcomes. KM plots revealed that patients classified into the DDRactivated subtypes had poorer OS [hazard ratio (HR) =1.516, P=5.9e-03; Figure 2A] and disease-free survival (DFS) (HR =1.312, P=6.92e-02; Figure 2B). Furthermore, we investigated clinical parameters between the two subgroups and found that the DDR-activated subgroup was associated with more diverse factors. Specifically, we observed a higher incidence of advanced M stage ( $0.01 \le P < 0.05$ , Figure 2C), higher grade N stage  $(0.01 \le P < 0.05, Figure 2D)$ , advanced pathologic stage (P<0.01, Figure 2E), and progressive T stage (0.01 $\leq$ P<0.05, *Figure 2F*) in the DDR-activated subgroup. Additionally, we observed a higher frequency of females and older individuals in the DDR-suppressed subgroups (P<0.01, Figure 2G), while patients in the DDR-activated subgroup were significantly younger compared to those in cluster 1 (P<0.01, *Figure 2H*). These results indicated that the expression level of the DDR gene has a significant impact on the clinical parameters and prognosis of patients with LUAD.

## DDR genes-based subtypes show distinct molecular characteristics

We conducted gene set variation analysis (GSVA) analysis to explore DDR-related pathways. The results revealed that the DDR-suppressed subgroups exhibited a higher frequency of involvement in the base excision repair pathway (P<0.01), Fanconi anemia pathway (P<0.01), homologous recombination pathway (P<0.01), and mismatch repair pathway (P<0.05). However, no significant



**Figure 2** Clinical prognosis analysis. (A) DDR-activated subtypes exhibited poorer OS. (B) DDR-activated subtypes exhibited poorer DFS. (C-F) The DDR-activated subgroup had a higher incidence of advanced M stage, advanced pathologic stage, and higher grade N stage. (G,H) Females and older individuals were more frequently observed in the DDR-suppressed subgroups. \*\*, 0.01≤P<0.05; \*\*\*, P<0.01. OS, overall survival; HR, hazard ratio; DFS, disease-free survival; DDR, DNA damage repair.

difference was observed between the two groups in the nonhomologous end-joining and nucleotide excision repair pathways (*Figure 3*).

When examining genomic alterations, we conducted a comparison of gene mutation differences between the two DDR subtypes using maftools. The analysis revealed a tumor median mutation burden of 3.39 mutations per megabase (MB) (*Figure 4A*), with a significantly higher TMB observed in the DDR-activated subgroups (P<0.01, *Figure 4B*). This finding suggests a potentially enhanced response to immunotherapy in the DDR-activated subgroups. Among the top 10 genes with the highest frequencies of driver mutational genes in patients from the training cohort were *TP53*, *TTN*, *CSMD3*, *RP1L1*, *XIRP2*, *STAB2*, *MMRN1*, *DCHS2*, *MTCL1*, and *LRP2* (*Figure 4C*). Notably, given the significance of *TP53*, we



**Figure 3** GSEA of the DDR subtype-specific pathways. \*\*, 0.01≤P<0.05; \*\*\*, P<0.01; ns, not significant. GSEA, gene set enrichment analysis; DDR, DNA damage repair.

further investigated and found a higher frequency of *TP53* mutations in the DDR-activated subgroup, which correlated with a poor prognosis (189/283 vs. 52/225, P<0.01) (*Figure 4C*). Missense mutations were the most prevalent type among these mutations (*Figure 4D*).

# DDR subtypes were characterized by different immune profiles

Immune cell infiltration has a significant impact on tumor progression and the response to immunotherapy. Therefore, we investigated the differences in immune cell infiltration between the DDR-activated and DDR-suppressed subgroups. Our analysis revealed that naive B cells (P<0.01),



**Figure 4** Genomic alterations between the DDR-activated and DDR-suppressed subgroups. (A) Tumor median mutation burden of 3.39/MB. (B) The TMB was significantly higher in cluster 2 (n=283) compared to cluster 1 (n=225) (P<0.01). (C) Mutual exclusion/cooccurrence analysis of mutations in each cohort. (D) Landscape of mutation profiles in the LUAD samples. The waterfall plot displays the mutation information for each gene in each sample. The data were analyzed based on TCGA data portal. \*\*\*, P<0.01. TMB, tumor mutational burden; MB, megabase; OR, odds ratio; CI, confidence interval; DDR, DNA damage repair; LUAD, lung adenocarcinoma; TCGA, The Cancer Genome Atlas.



**Figure 5** Immune profile alterations between the DDR-activated and DDR-suppressed subgroups. (A) Comparison of immune infiltration scores between the two subgroups. (B) Differential expression of PD-L1 (CD274) between the two subgroups. \*, 0.05≤P<0.1; \*\*, 0.01≤P<0.05; \*\*\*, P<0.01. NK, natural killer; FC, fold change; DDR, DNA damage repair; PD-L1, programmed death-ligand 1.

resting dendritic cells (P<0.01), resting mast cells (P<0.01), monocytes (P<0.01), plasma cells (P<0.05), and resting memory CD4 T cells (0.01 $\leq$ P<0.05) were significantly upregulated in the DDR-suppressed subgroup (*Figure 5A*). On the other hand, M0 macrophages, M1 macrophages, activated memory CD4 T cells, CD8 T cells, follicular helper T cells, resting natural killer (NK) cells (all P values <0.01), and neutrophils (0.01 $\leq$ P<0.05) were significantly upregulated in the DDR-activated subgroup (*Figure 5A*). Furthermore, we observed a significant upregulation of programmed death-ligand 1 (PD-L1; CD274) in the DDRactivated subgroup [log<sub>2</sub>fold change (log<sub>2</sub>FC) =-1.161, P=4.36e-22<0.01] (*Figure 5B*). This result is consistent with the prediction of TMB for immunotherapy mentioned above.

## Construction and validation of the prognostic DDR-related gene pair signature

Differentially expressed DDR-related genes were identified based on the mRNA-seq count data and sample clustering information in LUAD. We observed 604 upregulated genes in the DDR-activated subgroup and 933 downregulated genes in the DDR-suppressed subgroup (*Figure 6A*,6B).



**Figure 6** Analysis of differentially expressed DDR-related genes in LUAD. (A) Volcano plot depicting differential gene expression between cluster 1 and cluster 2, revealing 604 upregulated genes and 933 downregulated genes. (B) Abundance of highly expressed genes observed in the DDR-activated subgroups (cluster 2). (C) Upregulation of *UBD*, *EID3*, *SMC1B*, *DMC1*, and *CDKN2A* in the DDR-activated subgroups in contrast to the DDR-suppressed subgroups. DDR, DNA damage repair; LUAD, lung adenocarcinoma.

Furthermore, a comparison of differentially expressed DDR-related genes between the two subgroups was performed, with the results being presented in *Figure 6C*.

After the differentially expressed genes were screened, univariate Cox regression analysis was performed for each gene according to its expression level in LUAD the mRNAseq expression data (CPM) and clinical information of the samples. Finally, a total of 223 differentially expressed DDR-related genes were found to be significantly associated with the OS of patients with LUAD in the training set (P<0.001; table available at https://cdn.amegroups.cn/static/ public/jtd-23-1746-1.xlsx). To mitigate overfitting, LASSO regression was applied, and lambda.min ( $\lambda$ .min) was selected as the optimal regularization parameter, ensuring a more accurate model. Figure 7A,7B show the results of LASSO regression and cross-validation. The genes whose regression coefficient was not equal to 0 in LASSO regression analysis were then selected as marker genes. We observed that  $\lambda$ .min =0.095. From this analysis, nine genes were identified as hub DDR-related genes: CASP14, DKK1, ECT2, FLNC, HMMR, IGFBP1, KRT6A, TYMS, and FCER2 (Figure 7C). These hub genes were considered independent prognostic indicators of tumor prognosis. By using the expression levels of these selected genes, the corresponding risk scores for each sample was predicted (Figure 7D). Higher risk scores were associated with worse survival outcomes for the patients (Figure 7E, 7F).

To validate the robustness of the constructed model, the

patients were divided into high-risk and low-risk groups based on the median cutoff (Figure 8). In the training group, KM survival analysis revealed that the high-risk group exhibited significantly poorer OS compared to the low-risk group (HR =3.341, P=1.38e-08, Figure 8A). The area under the curve (AUC) values for the 1-, 3-, and 5-year follow-up periods were 0.767, 0.699, and 0.665, respectively, indicating the predictive ability of the model (Figure 8B). To further assess the prognostic performance of the model, individual risk scores were calculated using the aforementioned method, and patients in TCGA testing set and the entire TCGA set were classified accordingly. The predictions of the signature in these datasets were consistent with the previous findings. Specifically, the highrisk patients in all cohorts and the test group exhibited a significantly shorter OS compared to those in the low-risk group (Figure 8C,8E). The AUCs of the ROC curves for the 1-, 3-, and 5-year OS are presented in Figure 8D,8F and further supported the predictive ability of the model. The clustering heat map of marker genes (Figure 8G) indicated that patients with higher risk scores of FCER2 showed lower risk score status and superior survival. However, CASP14, DKK1, ECT2, FLNC, HMMR, IGFBP1, KRT6A, and TYMS exhibited opposing results.

In the external validation cohort (GSE13213), consistent with the previous findings, patients with higher risk scores exhibited significantly shorter OS compared to those with lower risk scores (*Figure 9A*). The AUC values of the ROC



**Figure 7** Construction of the LASSO regression model. (A,B) Graphs depicting the results of LASSO regression and cross-validation. (C) Results of one-way Cox regression analysis of marker genes (genes with nonzero LASSO regression coefficients). (D) Plot showing the sorting of risk values. (E,F) Graphs illustrating the relationship between risk values and survival status. HR, hazard ratio; CI, confidence interval; LASSO, least absolute shrinkage and selection operator.

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Figure 8 Internal validation of a prognostic model for predicting the prognosis of LUAD. (A) KM survival analysis showing that the highrisk subtypes in the training group had poorer OS. (B) Time-dependent ROC curves demonstrating the predictive performance of the model in the training group, with AUCs of 0.767, 0.699, and 0.665 for 1, 3, and 5 years, respectively. (C) OS analysis in the test group indicating that patients classified as high-risk subtypes had worse survival outcomes. (D) Time-dependent ROC curves in the test group, displaying AUCs of 0.743, 0.594, and 0.536 for 1, 3, and 5 years, respectively. (E) OS analysis in the total sample group revealing that patients classified as high-risk subtypes had inferior survival. (F) Time-dependent ROC curves in the total sample group, with AUCs of 0.745, 0.655, and 0.607 for 1, 3, and 5 years, respectively. (G) Clustering heat map of marker genes. OS, overall survival; HR, hazard ratio; TP, true positive; FP, false positive; AUC, area under the curve; LUAD, lung adenocarcinoma; KM, Kaplan-Meier; ROC, receiver operating characteristic.

FCER2

CASP14

KRT6A

FCT2

HMMR

TYMS

IGFBP1

DKK1

FLNC

0

-2

Status Dead

Alive



**Figure 9** External validation of a prognostic model for predicting the prognosis of LUAD. (A) External validation of OS. (B) External validation of ROC. (C) Correlation between age and risk scores. (D) Box plot showing the relationship between pathologic stage and risk scores. (E) Box plot showing the relationship between sex and risk scores. \*,  $0.05 \le P < 0.1$ . OS, overall survival; HR, hazard ratio; CI, confidence interval; TP, true positive; FP, false positive; AUC, area under the curve; F, female; M, male; LUAD, lung adenocarcinoma; ROC, receiver operating characteristic.

curves for the 1-year OS in the validation cohort were 0.87, respectively, indicating the favorable predictive ability of our model (*Figure 9B*). In addition, our verification results revealed significant correlations between the risk score and gender, T stage, age, and pathologic stage (*Figure 9C-9E*).

## Advanced DDR risk score was correlated with poor clinical factors and lower OS in patients with LUAD

We investigated the association between the DDR risk score and clinical factors in patients with LUAD. The high-risk group exhibited poor clinical characteristics, including advanced M stage (P<0.01, *Figure 10A*), advanced N stage (P<0.01, *Figure 10B*), lower pathologic stage (P<0.01, *Figure 10C,10D*), and advanced T stage (P<0.01,

*Figure 10E*, *10F*). Additionally, higher risk scores were observed in male patients (P<0.01, *Figure 10G*) and showed a negative correlation with age (correlation =-0.028, P=0.531). Tumor purity was found to have a positive correlation with the risk values (correlation =0.098, P=0.0277, *Figure 10H*, *10I*). Univariate Cox regression analysis revealed significant associations between the risk score and various clinical variables, including risk (HR =2.1, P=4.23e-15), age (HR =1.01, P=0.299), sex (HR =1.05, P=0.753), T stage (HR =1.52, P=8.91e-06); N stage (HR =1.7, P=1.39e-09), M stage (HR =2.13, P=0.00583), and pathologic stage (HR =1.66, P=8.08e-13, *Figure 10f*). Subsequently, multivariate Cox regression analysis revealed that the DDR risk score could serve as an independent prognostic factor (HR =1.96, P=1.33e-15, *Figure 10K*).



**Figure 10** Correlation analysis between DDR risk score and clinical factors of LUAD. (A) Difference in risk scores based on M stage. (B) Difference in risk scores based on N stage. (C,D) Difference in risk scores based on pathologic stage. (E,F) Difference in risk scores based on T stage. (G) Difference in risk scores based on sex. (H,I) Correlation between risk scores and age and between risk scores and tumor purity. (J) Single-factor Cox regression results for risk values and clinical information. (K) Multifactor Cox regression results for risk values and clinical information. \*,  $0.05 \le P < 0.1$ ; \*\*,  $0.01 \le P < 0.05$ ; \*\*\*, P < 0.01. HR, hazard ratio; CI, confidence interval; DDR, DNA damage repair; LUAD, lung adenocarcinoma.



Figure 11 Boxplot depicting the differential expression of immune checkpoint genes. \*,  $0.05 \le P < 0.1$ ; \*\*,  $0.01 \le P < 0.05$ ; \*\*\*, P < 0.01; ns, not significant. CPM, count per million.

## DDR signature is a promising predictor for immunotherapy

Firstly, the differential expression of immune checkpoint genes was demonstrated. Notably, ADORA2A, BTLA, CD200, LAIR1, and TIGIT were significantly up-regulated in the low-risk score subgroup. CD274, CD276, IDO1, and *PVR* were significantly up-regulated in the high-risk score subgroup (Figure 11). The TMB of each sample in TCGA-LUAD was then calculated by maftools. The median TMB was 3.4/MB (Figure 12A). TMB scores were significantly higher in the high-risk group (P<0.01, Figure 12B), indicating a better response to immunotherapy. The top 10 genes with the highest frequencies of driver mutations are shown in Figure 12C,12D. TP53 had a more frequent mutation in the high-risk group, indicating a poorer prognosis. The difference in the number of CTA between different risk groups was tested using wilcox.test. The result showed a higher CTA number in the high-risk group than the low-risk group (P<0.01, Figure 13). Moreover, the lowrisk group had a higher TIDE score compared to the highrisk group (P<0.01, Figure 14). Taken together, patients with a high-risk score may have a better immunotherapy effect.

### Discussion

DDR plays a crucial role in the development of various cancers by regulating multiple pathways involved in the interaction between tumor and immune cells (25-28). DNA damage can occur through endogenous events, such as oxidative damage, replication fork collapse, or errors that naturally occur during DNA replication or immune cell maturation, as well as exogenous factors such as ultraviolet rays, ionizing radiation, or chemical reagents. DNA repair is a pivotal mechanism for preserving genome stability and repairing DNA lesions. Deficiencies in the DNA repair pathway can influence tumor development, metastasis, and prognosis (14,29,30). The objective of this study was to investigate the predictive function of a DDR gene-related signature in the prognosis and immunotherapy response of LUAD.

Distinct clinical and molecular characteristics were observed in association with different DDR signatures. Specifically, patients belonging to the DDR-activated subgroup exhibited aggressive clinical manifestations, including advanced stage, poor differentiation, and an unfavorable prognosis. In contrast, the DDR-suppressed subgroup showed a higher frequency of alterations in base excision repair, Fanconi anemia pathway, homologous recombination, and dislocation repair. TMB is an important indicator that affects the treatment response and prognosis of lung cancer. Our results demonstrated that the DDRactivated subgroup had a significantly higher amount of variations in TMB compared to the DDR-suppressed subgroup. The TP53 gene, a critical DNA repair factor implicated in various cancers, has been reported to be more frequently mutated in the DDR-activated subtype (31,32). Notably, we found a higher frequency of troponin (TNN) variants in the DDR-activated subgroup, which has been associated with poor OS, increased immunogenicity, and altered immunotherapy prognosis in LUAD (33). Furthermore, CSMD3 was identified as being higher expressed in the DDR-activated subgroup. Previous



**Figure 12** Analysis of TMB in LUAD. (A) TMB score. (B) Relationship diagram showing the association between TMB and risk counts. (C) Distribution map of the 10 genes showing the greatest variation between groups. (D) Forest plot displaying the variation difference results of the top 10 genes. \*\*\*, P<0.01. TMB, tumor mutational burden; MB, megabase; OR, odds ratio; CI, confidence interval; LUAD, lung adenocarcinoma.



**Figure 13** Box plot illustrating the differences in CTA gene numbers between the low-risk and high-risk groups. \*\*\*, P<0.01. CTA, cancer-testicular antigen.

studies have highlighted the involvement of *CSMD3* gene mutations in immune response regulation and tumor prognosis (34,35).

We further investigated the immune microenvironment in LUAD and discovered that different DDR subtypes were associated with distinct immune profiles. In the DDRsuppressed subgroup, there was a notable increase in plasma cells, T lymphocytes, and activated memory CD4 cells, with mast cells being particularly abundant. Tumor-invading mast cells have been linked to resistance to anti-PD-1 therapy (36). Conversely, the DDR-activated subgroups exhibited heightened expression of M0 macrophages, M1 macrophages, activated memory CD4 T cells, CD8 T cells, follicular helper T cells, resting NK cells, and neutrophils. This immune cell composition could potentially enhance immune responses and improve patient outcomes (37-39). For certain patients with lung cancer, the use of immune checkpoint inhibitors (ICIs) has demonstrated improved



Figure 14 TIDE scores for the low-risk and high-risk groups. \*\*\*, P<0.01. TIDE, tumor immune dysfunction and exclusion.

long-term efficacy. Currently, several drugs are available to delay or prevent resistance to ICIs, and a high expression of PD-L1 serves as a significant indicator for ICI treatment (40-43). Notably, in our study, PD-L1 (CD274) expression was significantly elevated in the DDR-activated subgroup. These findings suggest that DDR subtypes exhibit distinct differences in immune cell infiltration, indicating potential variations in immunotherapy responses between the subtypes.

According to previous research, DDR genes have prognostic potential in various cancer types (44-46). In our study, we selected nine DDR-related genes (CASP14, DKK1, ECT2, FLNC, HMMR, IGFBP1, KRT6A, TYMS, and FCER2) to create a signature for identifying DDR subtypes in patients with LUAD, with the aim of clinical application. Patients were classified into different DDR types based on this signature in both the training and validation cohorts. To assess the accuracy of the model, patients were further divided into high-risk and low-risk groups. KM survival analysis demonstrated that the highrisk group had a poorer prognosis compared to the low-risk group. Moreover, univariate and multivariate Cox regression analyses indicated a significant negative correlation between the risk score derived from the DDR signature and other clinicopathological parameters in predicting survival. Additionally, the high-risk group exhibited lower TIDE scores, higher numbers of CTAs, and higher TMB scores, suggesting that patients with a high-risk score may have a more favorable response to immunotherapy (47,48). However, further randomized trials involving patients with LUAD receiving immunotherapy are necessary to validate the predictive performance of the DDR signature in terms of immunotherapy response. In summary, the risk score derived from the DDR signature can serve as a stable and independent indicator for prognosis and potential immunotherapy effect. It holds important clinical significance as an effective tool for classifying patients with LUAD. However, further research and validation are required to fully elucidate the clinical implications and utility of the risk score in guiding treatment decisions for patients with LUAD.

#### Conclusions

Our study contributes to the understanding of DDR heterogeneity and the identification of DDR subtypes in patients with LUAD. The distinct characteristics of DDR subtypes offer valuable insights into the clinical management and decision-making of LUAD. We have successfully developed a nine-gene risk signature associated with DDR in LUAD, which demonstrated its potential as an effective and stable classification tool for clinical practice. Moreover, our DDR subtype signature holds promise as a biomarker for guiding immunotherapy in patients with LUAD. Overall, these findings have significant implications for LUAD patient care and highlight the importance of considering DDR subtypes in personalized treatment approaches.

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*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (revised in 2013).

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