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ORIGINAL RESEARCH

Development of a Prognostic Nomogram for Acute Myeloid Leukemia on IGHD Gene Family

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Purpose: Acute myeloid leukaemia (AML) is a common haematological disease in adults. The overall survival (OS) remains unsatisfactory. It is critical to identify potential prognostic biomarkers and develop a nomogram that predicts overall survival in patients with AML.

Patients and Methods: We used gene expression dataset and clinical data from The Cancer Genome Atlas (TCGA) and Genotype-Tissue Expression (GTEx) to identify differential expression analysis, survival analysis, and prognostic value of IGHD gene family (IGHDs) in AML patients. A risk score model was built through Lasso analysis and multivariate Cox regression. We also developed a nomogram and evaluated its accuracy with Harrell's Harmony Index (C-index) and calibration curve. Last, the Therapeutically Applicable Research to Generate Effective Treatments (TARGET) database was used for external validation.

Results: IGHD1-20 mRNA expression level was an independent prognostic factor for patients with AML by multivariate analysis. After Lasso analysis and multivariate Cox regression, we constructed a 3-gene model (IGHD1-1, IGHD1-20, IGHD3-16) associated with OS in AML. Risk score and age were validated as independent risk factors for prognosis and were used to build a nomogram. The C index and calibration curve results show that its ability to predict 1-year, 3-year and 5-year overall survival is accurate.

Conclusion: The mRNA level of *IGHDs* was increased in AML patients. *IGHD1-20* was an independent risk factor for OS in AML patients. The IGHDs risk model (IGHD1-1, IGHD1-20, IGHD3-16) relates to the OS of AML patients. The nomogram, including risk score and age, can conveniently and effectively predict the overall survival rate of patients.

Keywords: acute myeloid leukaemia, IGHD gene family, IGHD1-20, prognosis

Introduction

Acute myeloid leukaemia (AML) is a malignant disease in which hematopoietic stem cells differentiate abnormally in the hematopoietic system, which leads to clonal expansion of primordial cells. Then abnormal primordial cells are accumulated at different stages of immaturity, which leads to the decrease of normal erythrocyte, leukocyte, and platelets, and then produces symptoms such as fatigue, dyspnea, infection, bleeding, and so on. AML can occur at different ages, but the elderly are the most common, and studies have shown that the median age at diagnosis is 68 years old.^{1,2} 60% to 85% of adults aged 60 or under can get a complete response after effective treatment. However, in elderly patients over 60, the complete remission rate is only 40% to 60%. Furthermore, most AML patients are prone to relapse within three years after diagnosis.³ Studies have shown that acute myeloid leukaemia is a disease with significant genetic heterogeneity,

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which refers to the accumulation of somatic acquired genetic changes in hematopoietic progenitor cells in AML patients, which will affect the standard mechanisms of self-renewal, proliferation, and differentiation. Moreover, the characteristic of this gene change is the most important feature that affects the prognosis of patients.^{4–6} Therefore, it is crucial to weigh disease-related and patient-related prognostic factors in determining the intensity of treatment. Thus, it is critical to developing practical, accurate biomarkers to evaluate the diagnosis and prognosis of patients with AML.

Studies on the expression frequency and level of Immunoglobulin yHeavy Chain (IgG) in AML showed that IgG was expressed in AML cell lines and primary myeloblasts with high frequency and high level, but not in monocytes and neutrophils of non-hematopoietic tumour patients and healthy controls. In further studies, IgG V_H DJ_H transcripts were detected in AML cell lines and selected primary myeloid cells, verifying that IgG expression was produced by AML cells. AML-derived IgG gene rearrangements demonstrate excessive somatic mutation of variable gene fragments, as well as the use of restricted (acute myeloid leukaemia cell lines) or biased (primary myeloid cell lines) gene fragments. Anti-human IgG can decrease the survival rate of AML cells and induce their apoptosis. Although the function of AML-derived antibodies remains unclear, these findings imply that AMLderived antibodies may be a novel AML-associated gene that is involved in the genesis and development of leukemia.7

The genes encoding the variable region of the H chain of human immunoglobulin molecule are recombined by variable genes (IGHV), diversity genes (IGHD), and linked genes (IGHJ), and each gene is selected from the ordered cluster of IGHV, IGHD, and IGHJ genes located on chromosome 14 (14q32.33).⁸ To some extent, antibody diversity results from combinatorial diversity, resulting from the many permutations that can occur when these genes recombine, a process that is often thought to involve essentially random rearrangement of the V, D, and D J genes. Some studies suggest that there are both random and genetic mechanisms for generating antibody lineages.^{9–11} Studies have shown that because the IGHDs provide a vital part of the antigen-binding loop in the Immunoglobulin, its role in determining antigen-binding specificity should not be underestimated has shown that its expression data in normal B cells is minimal. Due to the very short IGHDs fragments, their sizes vary from 11 to 37 nucleotides, coupled with exonuclease activity and point mutation, the correct identification of *IGHDs* becomes relatively tricky.¹² The locus of *IGHDs* was fully described and published not long ago, but now the development of high-throughput sequencing technology accelerates the further study of the human Ig gene and makes it possible for us to understand further and explore the role of the IGHDs in diseases.^{8,13}

Studies have shown that because the *IGHDs* supplies a vital section of the antigen-binding loop in the Immunoglobulin, its role in the essential region of determining antigen-binding specificity should not be underestimated, and studies have displayed that its expression data in normal B cells is minimal.^{12,14,15} Although some researchers have noticed the abnormal expression of immunoglobulin gamma heavy chain (IgG) in AML, the studies on human immunoglobulin heavy-chain *IGHD* gene repertoire abnormal expression and prognosis are still few. This study will retrospectively analyze and study the survival prediction value of the expression level of the *IGHDs* in AML based on the considerable sample information from the public database.

Patients and Methods Transcriptome Data and Clinical Information

The workflow of the development of a prognostic nomogram for acute myeloid leukaemia on the *IGHD* gene family is shown in Figure 1. First, we downloaded the GTEx gene expression dataset, the TCGA acute leukaemia gene expression dataset, and the corresponding sample information from the University of California Santa Cruz (UCSC) genome database. Then, we extracted the gene expression data of normal bone marrow samples (70 samples) from the GTEx gene expression dataset and AML samples (151 samples) from the TCGA gene expression dataset. The LIMMA software package was used to merge the expression data of the two gene databases for standardized processing, and the expression of *IGHDs* was extracted for subsequent analysis.

Differential Expression Analysis of IGHDs

Extracted from the transcriptome sequencing data, *IGHDs* quantity data, using R language limma packet analysis, identified the 32 family genes differentially expressed in AML and normal bone marrow. An absolute log2-fold change (|



Figure I Workflow of the study.

FC|) of > 1 and an adjusted P-value of < 0.05 were set as cutoff criteria. Clustering heat maps were drawn using the PHEATMAP software package in R language to show the expression of each gene in the sample.

Survival Analysis

Using R packet "Survminer" and "Survival ", all *IGHDs* expression levels were stratified, and the patients were divided into high RNA expression levels groups and low RNA expression levels groups. Kaplan-Meier survival analysis was used to clarify the association between 5-year OS and *IGHDs* expression.

Build a Prognosis Model with IGHDs

The survival package was used to analyze the survival of *IGHDs* with differentially expressed genes (DEGs) in AML patients, and then the univariate Cox regression analysis was fitted, the P < 0.05 gene as a prognostic significance. The OS prognostic risk model was established by using Lasso-penalized and multivariate Cox regression analysis. A linear combination of regression coefficients obtained from multivariate Cox regression analysis multiplied by their model gene expression produced a prognostic risk score for all three genes.^{18,19} According to the median risk score (M), all patients were

divided into high-risk and low-risk groups, and a risk heat map, risk curve, and survival status were drawn. Kaplan-Meier method was used to plot the survival curve of the *IGHDs* model, and the Log-Rank method was used to analyze the prognostic difference between the high-risk group and the low-risk group. Time-dependent receiver operating characteristic (ROC) curves were used to evaluate the ability of the *IGHDs* model to predict accuracy in 1 - year, 3 - year, and 5-year overall survival.

Independent Analysis of Risk Scores

After controlling for other confounding factors, gender, age, bone marrow blasts, haemoglobin, leukocyte, monocytes, and risk score of AML patients were included in univariate and multivariate Cox regression analysis to explore the effect of gender, age, bone marrow blasts, haemoglobin, leukocyte, monocytes, and risk score on prognosis.

Establishment and Evaluation of the Nomogram in the TCGA Dataset

We constructed the Nomograph based on the multivariate Cox regression analysis of the factors related to the significant prognosis of AML. We used the Hosmer-Lemeshow test to draw the calibration curve of the Nomograph to evaluate the prediction ability of the Nomograph in the queue. Harrell's concordance index (C-index) was used to evaluate the discrimination performance of the Normogram quantitatively. The C-index is between 0. 5 and 1. The accuracy of the C-index between 0.50 and 0.70 is low, between 0.71 and 0.90 is medium accuracy, while higher than 0.90 is high accuracy.

Gene Ontology (GO) and the Kyoto Encyclopedia of Genes and Genomes (KEGG) Analysis

We performed gene ontology (GO) analysis of differentially expressed genes (DEGs) in patients with high-risk scores to evaluate the richness of biological process (BP), cellular component (CC), and molecular functional (MF) annotation. The Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis is also used to annotate the signalling pathways associated with these genes. The "cluster profile" R software package was used for GO and KEGG enrichment, and the adjusted p-value less than 0.05 was regarded as the cutoff value. Use the "ggplot2" R software package to visualize the results of GO and KEGG analysis.

Correlation Analysis of Family Genes

We use R's corrplot package to draw the correlation coefficient matrix of *IGHDs* obtained by calculating the correlation coefficient of family genes.

External Validation

We validate the nomogram via using expression data of the TARGET-AML gene expression dataset and the corresponding clinical information from the University of California Santa Cruz (UCSC) genome database.

Statistical Analyses

Data analysis and drawing are done by R3.6.1, GraphPadPrism 8.0.1, and SPSS software (version 23.0).

Results

Differential Expression of IGHDs in AML and Normal Bone Marrow

Download AML clinical information and calibrated AML gene expression data from the TCGA database. Moreover, normal bone marrow samples were from the GTEx. The expressions of *IGHDs* were extracted for differential analysis. The results indicated that all the 31 *IGHDs* were differentially expressed, and the expression was upregulated in AML. Differential expression levels were visualized via a heatmap (pValue ≤ 0.05 , Figure 2).

The Effect of Abnormal IGHDs Expression on Survival in AML

All *IGHDs* expression levels were stratified, and the patients were divided into high RNA expression levels groups and low RNA expression levels groups according to the median value of *IGHDs* transcript expression. The median values of each IGHD family gene expression were shown as follows: *IGHD1-1* 4.99,*IGHD1-20* 4.36,*IGHD2-2* 4.12,*IGHD2-15* 3.98,*IGHD3-3* 5.64,*IGHD3-10* 4.39, *IGHD3-16* 5.49,*IGHD6-19* 5.03,*IGHD2-8* 3.75,*IGHD2-21* 2.84,*IGHD3-9* 4.99,*IGHD3-22* 3.45,*IGHD6-25* 3.28, *IGHD4-17* 4.51,*IGHD5-12* 5.17,*IGHD6-13* 4.75.

We next evaluate the prognostic effect of *IGHDs* expression on survival. By Kaplan-Meier analysis, patients with high *IGHDs* (*IGHD1-1,IGHD1-20,IGHD2-2*, *IGHD2-15,IGHD3-3,IGHD3-10,IGHD3-16,IGHD6-19, IGHD2-8,IGHD2-21,IGHD3-9,IGHD3-22,IGHD6-25,*



Figure 2 Heatmap of differential IGHDs. The blue, white, and red colors of the subject correspond to low, medium, and high expression. ****pValue <0.001; *pValue <0.05.

IGHD4-17, IGHD5-12, IGHD6-13) expression showed high expression of IGHDs presented shorter OS than those with low expression of IGHDs among AML patients (pValue ≤ 0.05 , Figure 3). For other IGHDs transcripts, there was no difference in survival time between the highexpression and low-expression groups.

Analysis of Prognostic Value and Prognostic Model of IGHDs

The univariate Cox analysis of *IGHDs* was performed for these patients. The univariate analysis results showed that the effects of 13 differentially expressed *IGHDs* on the prognosis of patients with AML were statistically significant (pValue ≤ 0.05), and these 13 genes were further analyzed as candidate genes. The multivariate analysis results displayed that *IGHD1-20* was an independent influencing factor for patients with AML, and the prognosis of patients with high expression was worse (HR=1.47, 95% CI 1.03– 2.11, pValue=0.035, Table 1). The 13 candidate genes selected by univariate Cox regression analysis were analyzed using the Lasso Cox selection method and multiple. Cox regression analysis (Figure 4A and B). As a result, the model was composed of three genes: *IGHD1-1, IGHD1-20*, and *IGHD3-16*. According to the regression coefficient of the prognostic model, the risk score formula of the IGHD gene family was obtained as follows: risk score = (0.565220138× IGHD1-20 expression) + (0.401795251× IGHD1-1 expression)- $(0.695077172 \times IGHD3-16 \text{ expres})$ sion), as shown in Table 2. The risk score of each AML patient was computed in line with the risk score formula. All patients were separated into high-risk group (risk score \geq M) and low-risk group (risk score \leq M) with the median risk value (M= 1.160621935) as a cutoff value. The results of Log-Rank displayed that the prognosis of the high-risk group was worse than that of the low-risk group (pValue ≤ 0.05), as shown in Figure 4C, There was significant variance in the survival curve between the highrisk group and low-risk group, and survival was significantly higher in the low-risk group than in the high-risk group. From the Area Under Curve (AUC) value of the ROC curve, it can be seen that the prediction model has high prediction accuracy (Figure 4D–F). In the risk heat map, the expression of each model gene increases with the increase of patient risk score, indicating that all IGHDs model genes are high-risk genes (Figure 4G). Figure 4H and I reflect patients' risk score distribution and survival status distribution in high and low-risk groups. In general, the risk score curve does an excellent job of distinguishing between high-risk and low-risk groups (Figure 4H). On the



Figure 3 Single-gene survival analysis of IGHD family members in AML patients.

risk score curve, the patient's risk score increased from left to right. Survival time decreased, and mortality increased with increased patient risk scores, as displayed in Figure 4I. Finally, the sex, age, bone marrow blast cell, haemoglobin, leukocyte, monocyte, and risk score of AML patients were involved for univariate and multivariate analysis. Univariate and Multivariate analysis displayed that age and risk score were independent risk factors for

Table I Univariate and Multivariate Cox Regression Analysis Results of IGHDs

ld	Unicox			p value	Multicox			pvalue
	HR	HR.95L	HR.95H		HR	HR.95L	HR.95H	
IGHD1-14	1.09982264	1.008361348	1.1995797	0.031719172	0.8222074	0.59262476	1.140730286	0.24127283
IGHD3-10	1.10578376	1.011842555	1.2084466	0.026427744	1.15927344	0.78517634	1.711609007	0.45722217
IGHD2-2	1.11378333	1.009833466	1.2284335	0.031105958	0.86646661	0.60391372	1.243164977	0.43644977
IGHD6-19	1.10447969	1.014729443	1.2021681	0.021555552	0.99097821	0.72447493	1.355516627	0.95478006
IGHD2-15	1.10373212	1.006146892	1.210782	0.036643541	1.02895318	0.77390763	1.368050407	0.84430339
IGHD5-18	1.10519805	1.014099985	1.2044796	0.02266845	0.90237015	0.6728094	1.210256415	0.49279043
IGHD3-16	1.09467733	1.006049818	1.1911125	0.035730162	0.78631733	0.52620783	1.175001419	0.2407827
IGHD1-7	1.1070905	1.009365324	1.2142773	0.030953886	1.01407497	0.71722322	1.433790808	0.93695718
IGHD5-5	1.08361329	0.995988307	1.1789473	0.061967505	0.98708302	0.72760093	1.339103426	0.93341716
IGHD1-I	1.13837185	1.037486037	1.2490679	0.006196191	1.31742343	0.96286721	1.802537762	0.08481484
IGHD3-3	1.09862233	1.002714648	1.2037034	0.043577528	0.9524548	0.69409	1.306992097	0.76286936
IGHD1-20	1.1303205	1.036228713	1.232956	0.005736078	1.4703998	1.0256788	2.107946098	0.0359109
IGHD7-27	1.12613933	1.023226431	1.2394029	0.015117656	1.04666363	0.9224253	1.1876352	0.47929488

Notes: P ≤0.05 in univariate and multivariate analysis of IGHD I-20. The multivariate analysis column of IGHD I-20 was bolded to highlight the significance of IGHD I-20.



Figure 4 (A, B) Creation of the prognostic model based on LASSO algorithm; (C) the survival curve of patients in high and low risk groups; (D–F) the Area Under Curve (AUC) value of the ROC curve corresponding to 1 year, 3 years, and 5 years of *IGHDs* model; (G) risk heatmaps; (H) Risk score curves; (I) survival status figures.

the prognosis of patients with AML, and the patients with high scores had a poor prognosis (pValue ≤ 0.05 , Figure 5A and B).

Establishment and Evaluation of a Nomogram for OS Prediction in AML

Age and risk level were significant as critical predictive factors in univariate and multivariate regression analyses (Figure 5A

Table	2	Multiple	Cox	Regression	Analysis	of	the	Model	Genes
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Id	Coef	Exp(Coef)	Se(Coef)	z
IGHD3-16	-0.69508	0.499036	0.359876	-1.93143
IGHD1-1	0.401795	1.494505	0.255663	1.571582
IGHD1-20	0.56522	1.759835	0.317023	1.782898

and B). Considering all the essential predictors above, we have generated a comprehensive nomogram (Figure 5C). The C-index was 0.737. The calibration curve of the *IGHDs* model for the possibility of OS at 1 year, 3 years, and 5 years exhibited accurate predictive ability (Figure 5D–F).

Expression of Model Genes in Different Karyotypes

The European Leukemia Network divides patients into favourable, moderate, and risk groups in the stratification of acute leukaemia. We also analyzed the relationship between the *IGHD1-1*, *IGHD1-20*, *IGHD3-16* expression and cytogenetic karyotype. The results displayed that patients in the favourable group had the



Figure 5 (A) Univariate prognostic analysis of TCGA-AML clinical indicators and risk scores (B) Multivariate prognostic analysis of TCGA-AML clinical indicators and risk scores. (C) Normogram for predicting 1-year, 3-year, and 5-year OS (D–F) Calibration curves of 1-year, 3-year and 5-year OS Black broken lines symbolizes the perfect predicted model, and the blue full line symbolizes the noticed model. (G–I) The expressions of model genes according to FAB classification in AML. (J–L) The expressions of IGHD1-1, IGHD1-20, IGHD3-16 between AML karyotype.

lowest *IGHD1-1, IGHD1-20, IGHD3-16* expression (pValue ≤ 0.05), and those in the poor group had the highest *IGHD1-1, IGHD1-20, IGHD3-16* transcript expression (pValue ≤ 0.05 , Figure 5G–I).

Expression of Model Genes in French-American and British (FAB) Subtypes of AML

We assayed the expression of model genes in the M0 to M5 FAB subtypes of 130 patients with acute myeloid

leukaemia; only two patients with AML-M6 and only one patient with AML-M7 were not included in the analysis. We found that *IGHD1-1*, *IGHD1-20*, *IGHD3-16* have higher expression in the AML-M0 subtype than in other subtypes.*IGHD1-1* revealed that the lowest transcript expression was detected in AML-M3. *IGHD1-20* transcripts showed lower expression of AML-M5 than other subtypes.AML-M3/M5 patients displayed a lower transcript expression of *IGHD3-16* in contrast to that in other subtypes (Figure 5J–L).

Clinical Characteristics of IGHD1-20 in the High Expression Group and Low Expression Group

IGHD1-20 gene expression in AML patients can be regarded as an independent prognostic factor according to our multivariate regression analysis results. In order to compare the clinical characteristics of the IGHD1-20 high expression group and IGHD1-20 low expression group, All the clinical indicators, such as the age, bone marrow blast cell, peripheral leukocyte, peripheral monocyte, and haemoglobin of AML patients in the TCGA database, were segregated into IGHD1-20 high expression group and IGHD1-20 low expression group and analyzed by Nonparametric test. Fisher's exact test was used to compare the differences in sex, FAB type, and karyotype between the high expression and low expression groups. In these patients, the age, haemoglobin, FAB type, and karyotype between the IGHD1-20 high expression group and IGHD1-20 low expression group were considerably different (pValue <0.05, Supplementary Table 1). However, the sex, bone marrow blast cell, peripheral leukocyte, or peripheral monocyte is not observed differences (pValue>0.05) of these patients with AML statistically.

Mutation Status of IGHDs and Their Correlation with Each Other

In order to evaluate the correlation between the *IGHDs*,we use R packet " corrplot" to analyse. The result dispalyed that *IGHD1-20* expression was related with *IGHD1-14* (r = 0.91), *IGHD1-1*(r = 0.89). The expression of *IGHD3-16* was discovered to be closely associated to that of *IGHD1-1*(r = 0.9), *IGHD1-14*(r = 0.92), *IGHD1-20*(r = 0.95), *IGHD1-26*(r = 0.82), *IGHD1-7*(r = 0.9), *IGHD2-15*(r = 0.92), *IGHD2-2*(r = 0.93), *IGHD2-21*(r = 0.86), *IGHD2-8* (r = 0.89) and *IGHD3-10*(r = 0.94) (Figure 6A).

GO Analysis of DEGs in AML Patients with the High-Risk Score

GO analysis was used to investigate the functional characteristics of these DEGs in patients with high-risk scores, and they were sorted into the biological process (BP), cellular component (CC), and molecular functional (MF) annotation. As shown in Table 2, in the BP group, up-regulated genes were mainly enriched in phagocytosis (GO:0006909), immune response-activating cell surface receptor signalling pathway (GO:0002429), leukocyte (GO:0050900), migration complement activation (GO:0006956), and positive regulation of lymphocyte activation (GO:0051251). In the CC group, upregulated genes were mainly enriched in immunoglobulin complex (GO:0019814, GO:0042571), external side of plasma membrane (GO:0009897), mitochondrial inner membrane (GO:0005743), and MHC protein complex (GO:0042611). In the MF group, up-regulated genes mainly were enriched in antigen binding (GO:0003823), immunoglobulin receptor binding (GO:0034987), electron transfer activity (GO:0009055), and peptide antigen binding (GO:0042605). These data imply that the identified DEGs in AML patients with the high-risk score are mainly enriched in immune response (Supplementary Table 2, Figure 6B).

Signaling Pathway Enrichment Analysis

We conducted a pathway enrichment analysis of AML patients with high-risk scores by KEGG further to understand the biological changes in the disease course. KEGG pathways enrichment analysis indicated that those DEGs were enriched in Hematopoietic cell lineage, Phagosome, Human T-cell leukaemia virus 1 infection, and Cell adhesion molecules (Supplementary Table 3, Figure 6C).

Multiple Validations of Model Genes

Boxplots of *IGHD1-1*, *IGHD1-20*, and *IGHD3-16* were drawn to verify the expression differences of *IGHD1-1*, *IGHD1-20*, and *IGHD3-16* between the high-risk score and low-risk score groups. Figure 6D–F reflect that the expression levels of *IGHD1-1*, *IGHD1-20*, and *IGHD3-16* in the high-risk score group are higher than those in the low-risk score group. *IGHD1-1*, *IGHD1-20*, and *IGHD3-16* are considerably highly expressed in the high-risk score group, which proves the inference that they are all high-risk genes to a certain extent.

The prognostic value of *IGHDs* and the *IGHDs* risk score model were confirmed using the TARGET database. The 278 patients with AML were selected from the TARGET database to explore the prognostic value of *IGHDs* and evaluate the *IGHDs* risk score model. By Kaplan-Meier analysis, patients with high *IGHD1-20* expression had shorter OS than those with low expression of *IGHD1-20* among AML patients (pValue ≤ 0.05 , Figure 6G). Univariate COX analysis showed that *IGHD1-20* was a prognostic factor in patients with AML (pValue=0.002).We used the *IGHDs* risk score model (risk



Figure 6 (A) The correlations of *IGHDs* with each other in AML. blue suggests positive correlation, Red suggests negative correlation, and the depth of color suggests the degree of correlation. Gene functional enrichment of differentially expressed in patients with high-risk scores. (B) Gene ontology analysis; (C) The top 15 most enriched KEGG pathways. (D–F) Boxplots for the difference confirmation of model genes. (G) Kaplan-Meier survival curve of *IGHD1-20* in validation cohort using TARGET database.

0.565220138095178* IGHD1-20 score +0.401795250677606* IGHD1-1 0.695077172034113*IGHD3-16) created to compute the risk score of each AML patient. According to the median value, the risk score model could classify the patients in the TARGET database into high-risk groups and low-risk groups. Patients with high-risk groups had shorter OS than those with low-risk groups among AML patients (pValue≤0.05, Figure 7A). The nomogram calibration curves for the possibility of 1-year, 3-year, and 5-year OS exhibited apparent concordance between the forecast outcomes and observations in the TARGET database (Figure 7B–D). The validation results are similar to those in the TCGA queue.

Discussion

In the past few decades, the high heterogeneity of AML has posed considerable challenges to risk stratification.²⁰ The next-generation sequencing technology advanced in recent years adopts the whole genome sequencing method, which carries enormous benefits to multi-population data mining.²¹ As the second generation sequencing technology, such as the application of the molecular targets of

AML, is becoming more and more and further lead to the speedy advancement of targeted drugs, not only that, the second-generation sequencing technology can be used to excavate potential biomarkers, and is used to predict the prognosis of patients with AML results, and select the appropriate treatment strategy, reduce the financial burden on patients and their families.²² Many studies using bioinformatics analysis have suggested the relationship between gene transcriptome characteristics and the prognosis of AML.^{23–30}

There is growing evidence that Immunoglobulin (Ig) is expressed at a high level in cancer cells, and this type of Ig has been named cancer-derived Immunoglobulin. Cancerderived immunoglobulin plays a profound pro-tumour role by promoting the malignant behaviour of cancer cells, mediating tumour immune escape, inducing inflammation, and other mechanisms. Importantly, cancer-derived immunoglobulins show potential applications as diagnostic and therapeutic targets for cancer patients.³¹ Studies have shown that AML-derived IgG may be related to the occurrence of leukaemia and the progression of AML.

We have proved that IgH rearrangement (V_H -D, D-J_H) is the core of antibody diversity. The recombination site



Figure 7 Validation of the prediction model. (A) the survival curve of patients in high and low risk groups. (B–D) Calibration curve for the risk score model using TARGET database.

largely determines the specificity of antibodies called the third complementary determining region (CDR3), Because the IGHD gene is involved in complementary determination region 3 (CDR3), CDR3 covers the connection between V, D and J genes.^{10,32,33} Most of the diversity seen in the third complementary determination region (CDR3) is caused by IGHD fragments and joints formed by changes injunctions and N fragments during D-J_H and V_H-D rearrangements. The mechanism of regulating rearrangement is considered to follow the rule of 12/23, in which D-D or V_H-J_H rearrangement is excluded. However, The V (DD) J recombination conjecture put forward in the study of Kurosawa and Tonegawa in 1982.34 Meek KD's research reveals for the first time that D-D fusions do take place, either a direct rearrangement or an inverted rearrangement. The resulting fusion D segment may be fully capable of being carried out in subsequent $D-J_H$ and V_H -D rearrangements. V (DD) J recombination integrates two IGHD genes into a super-long CDR3 (or tandem CDR3) with tandem fusion *IGHD* gene. Briney BS also proved that tandem CDR3s do exist.³⁶ Our analysis showed that, except *for IGHD4OR15-4A*, and *IGHD4OR15-4B*, all *IGHDs* were up-regulated in patients with Acute Myeloid Leukemia compared with normal controls. The existence of the V (DD) J recombination phenomenon proves the rationality of the up-regulation of all *IGHDs* in our patients with acute myeloid leukaemia. We also found out that the expression of *IGHD1-20* was considerably increased in AML. Compared with the low expression of *IGHD1-20* was significantly worse, and *IGHD1-20* was also an independent prognostic factor.

We found that the transcript expression of *IGHD1-1*, *IGHD1-20*, *IGHD3-16* was considerably distinct in different AML subtypes. The expression of these three genes was the highest in AML-M0. The mechanism of this correlation needs further study. Besides, we evaluated the *IGHDs* model as a predictor of the risk of acute myeloid leukaemia. Risk stratification for age and genetic characteristics of AML was considered. The expression of *IGHD1-1, IGHD1-20*, and *IGHD3-16* transcripts in the high-risk group was significantly higher than that in the low-risk group. Multivariate cox analysis showed that *IGHD1-20*, risk stratification were independent risk factors. Despite these results, few reports of associations exist between other *IGHD*s and AML prognostic genes, so this further study is warranted.

MohamedKhass studies have found that specific amino acids in core *IGHD* gene fragments greatly influence the absolute number of developing and mature B cell subsets, antibody production, epitope recognition, protection against pathogen attack, and sensitivity to the production of autoreactive antibodies.³⁷ Our study also performed KEGG pathway enrichment analysis and GO analysis on the genes expressed in patients with the high-risk score. Some of the functions in the results of these analyses are consistent with the known functions of the *IGHD*s gene family, and their significant enrichment in high-risk patients coincides with the high expression of *IGHD*s genes in high-risk patients.

Research by Vale shows that murine *IGHD*s deletions increased susceptibility to infection.³⁸ Some studies have found that changes in the *IGHDs* gene sequence also increase the likelihood of self-reactive antibodies.³⁹ Concerning the clinical significance of *IGHD3-3*, most studies have focused on its non-mutation as a factor in the poor prognosis of chronic lymphocytic leukaemia.

Tschumperrc found that in a large sample of patients with low/moderate RAI risk and non-mutated chronic B-lymphoblaemia, patients expressing *IGHD3-3* required treatment at an earlier time than other non-mutated chronic B-lymphoblaemia and showed that their prognosis was independent of the use of the *IGHV* and *IGHJ* genes.¹⁴ Boyd found deletions of *IGHD1-7*, *IGHD2-8*, *IGHD3-3*, *IGHD4-4/4-11*, and *IGHD6-6* gene sequences in patients with aplastic anaemia whether the deletions of the role of these genes in pathogenesis remains to be studied.⁴⁰

Considering its cost-effectiveness, our signature based on *IGHDs* is easy to exam routinely. There are also some shortcomings to this study. First, the prognostic model's construction was based on a non-randomized, retrospective observational analysis using registry data, which would allow for the introduction of bias. Although its good performance in external validation indicates its potential, more extensive studies may be needed to validate our results further. Although the nomogram successfully predicted OS in AML patients by combining age and risk level, clinical characteristics were viewed as inadequate due to the restricted information in the study cohort. In the future, it will be essential to create a better prognosis nomogram from more centres, with complete clinical information and sequencing data. Secondly, the role of *IGHDs* in the appearance and advancement of AML and its complex mechanism needs to be further studied.

Conclusion

We first studied the role of *IGHDs* in the prognosis of acute myeloid leukaemia. We found that *IGHD1-20* is an independent prognostic factor of acute myeloid leukaemia. We established a prognostic signature of *IGHDs*, constructed a nomogram based on the risk model, and evaluated the nomogram by C-index and calibration curve. Our study provides a new way further to evaluate the prognosis of patients with acute myeloid leukaemia.

Abbreviations

AML, acute myeloid leukemia; OS, overall survival; TCGA, The Cancer Genome Atlas; GTEx, Genotype-Tissue Expression; IGHDs, IGHD gene family; C-index, Harrell's concordance index; TARGET, the Therapeutically Applicable Research to Generate Effective Treatments; DEGs, differentially expressed genes; IgG, Immunoglobulin yHeavy Chain; ROC curves, the time-dependent receiver operating characteristic curves; GO, gene ontology; KEGG, The Kyoto Encyclopedia of Genes and Genomes; AUC, the area under curve; Ig, immunoglobulin; FAB subtypes, French-American and British subtypes.

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Disclosure

The authors report no conflicts of interest in this work.

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