A metabolic reprogramming-related prognostic risk model for clear cell renal cell carcinoma: From construction to preliminary application

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Metabolic reprogramming is one of the characteristics of clear cell renal cell carcinoma (ccRCC). Although some treatments associated with the metabolic reprogramming for ccRCC have been identified, remain still lacking. In this study, we identified the differentially expressed genes (DEGs) associated with clinical traits with a total of 965 samples via DEG analysis and weighted correlation network analysis (WGCNA), screened the prognostic metabolism-related genes, and constructed the risk score prognostic models. We took the intersection of DEGs with significant difference coexpression modules and received two groups of intersection genes that were connected with metabolism via functional enrichment analysis. Then we respectively screened prognostic metabolic-related genes from the genes of the two intersection groups and constructed the risk score prognostic models. Compared with the predicted effect of clinical grade and stage for ccRCC patients, finally, we selected the model constructed with genes of ABAT, ALDH6A1, CHDH, EPHX2, ETNK2, and FBP1. The risk scores of the prognostic model were significantly related to overall survival (OS) and could serve as an independent prognostic factor. The Kaplan-Meier analysis and ROC curves revealed that the model efficiently predicts prognosis in the TCGA-KIRC cohort and the validation cohort. Then we investigated the potential underlying mechanism and sensitive drugs between high- and low-risk groups. The six key genes were significantly linked with worse OS and were downregulated in ccRCC, we confirmed the results in clinical samples. These results demonstrated the efficacy and robustness of the risk score prognostic model, based on the characteristics of metabolic reprogramming in ccRCC, and the key genes used in constructing the model also could develop into targets of molecular therapy for ccRCC.

KEYWORDS
clear cell renal cell carcinoma, metabolic reprogramming, weighted correlation network analysis, prognostic model, key gene
Introduction

Renal cell carcinoma (RCC) is a universal cancer, which accounts for 2.2% of the total cancer incidence and 1.8% of the total cancer mortality (1). According to The Cancer Statistics (2), at the end of 2021, about 76,080 new cases of RCC would be diagnosed, and 13,780 patients with RCC would die in the USA. More than 40% of RCC patients will have metastasis after surgical intervention (3). Clear cell renal cell carcinoma (ccRCC), the prevailing and invasive histological subtype of RCC, has become a worldwide issue. Clinicians mainly employed the T, N, and M classification system and Fuhrman nuclear grade to prognosticate patients’ prognosis of ccRCC and guide clinical treatment decision-making. However, these prognostic tools require improvements and novel, robust, and specific prognostic models to acquire more accurate predictions.

A complex biological system cannot be changed by a single part of its components but by the interaction of these components. Bioinformatics, which introduces computational methods and mathematical models, enlarges the magnitude of data accumulated in the genomic, transcriptomic, and proteomic studies, allowing us to simulate the complexity of the biological system and understand these systems (4). Bioinformatics technologies have become increasingly prevalent in finding molecular mechanisms and specific biomarkers of diseases.

Weighted correlation network analysis (WGCNA) and differently expressed gene (DEG) analysis are increasingly being used as the analytical methods of bioinformatics. WGCNA is a system biology method for discovering modules of highly correlated genes and summarizing these modules by using the intramodular hub gene (5). Then, selecting important modules associated with clinical traits for further analysis. DEG analysis can find quantitative changes in gene expression levels and study molecular mechanisms of gene regulation. Using the combination of DEG and WGCNA could improve the accuracy of discriminating highly related candidate biomarker genes. Analyzing the genes that have been screened out, we found that these genes were primarily related to metabolism. Previous studies have shown a strong link between RCC and changes in metabolic pathways (6–8), and abnormally accumulated lipid droplets have been found in the ccRCC cytoplasm (9). Nevertheless, the prognostic effect of these metabolic genes on patients remains unclear.

In this study, we used WGCNA and DEG analyses to analyze the mRNA expression data of ccRCC from The Cancer Genome Atlas (TCGA) and Gene Expression Omnibus (GEO) databases, which showed differential co-expression genes, and to explore the relationship between these metabolic genes and prognosis of patients with ccRCC. Screening by WGCNA, we obtained the clinical traits of correlated genes, which may be considered as biomarkers and targets for treatment. Using prognostic metabolism-related genes, we constructed a prognostic prediction model and validated it.

Materials and methods

The workflow of this study is shown in Supplementary Figure S1, and we will elaborate on each step in the following sub-sections.

Acquisition datasets from TCGA and GEO databases

The gene expression dataset of ccRCC was downloaded from TCGA (https://portal.gdc.cancer.gov/) and GEO (https://www.ncbi.nlm.nih.gov/geo) databases. Before analysis of the dataset, patients with missing data of pathological diagnosis and corresponding clinical information were excluded. Afterward, the ccRCC dataset downloaded from the TCGA database included 611 samples and corresponding clinical information. The TCGA-KIRC dataset annotated using the Human hg38 gene standard track contains 72 normal counts, 539 tumor counts, and 19600 genes of RNAseq data.

Four datasets, including GSE36895, GSE46699, GSE53757, and GSE66270, were downloaded from the GEO database. The platform of such datasets is GPL570 [HG-U133_Plus_2] Affymetrix Human Genome U133 Plus 2.0 Array, which was used in gene probe annotation. We combined these four datasets into a single dataset and then normalized and cleaned the merged dataset using the R package affy (version 1.66.0), impute (version 1.62.0), limma (version 3.44.3), and sva (version 3.36.0) (10–12) (https://bioconductor.org/bioclite/R). The merged GEO-ccRCC dataset included 172 normal samples and 182 tumor samples. If one gene corresponded to duplicated probes, then we used the mean value of these probes.

Identification of robust DEGs

TCGA-KIRC and GEO-ccRCC datasets were utilized for analysis. The ccRCC samples of patients were divided into two sets, normal and tumor samples. The R package limma (version 3.44.3) was used in analyzing the data and screening the DEGs with |logFC|>1 and adj. P<0.05. By using R software, the DEGs of TCGA-KIRC and GEO-ccRCC datasets were visualized as a volcano plot, the abscissa and ordinate of which were adj.P and logFC, respectively. The upregulated genes were marked red, and the downregulated genes were marked green. The top 100 DEGs were visualized by a heatmap plot.

WGCNA and Venn diagram

WGCNA was used to identify the key modules of highly correlated genes and explore the relationship between network genes and external sample traits, with the expression data
obtained from TCGA and GEO databases. The R package limma (version 3.44.3) was used in checking these expression data, removing duplicate rows, and replenishing missing values. The R package WGCNA (version 1.70.3) was used in analyzing the data. The samples were clustered (cut line as 20,000), and all of the samples were divided into two groups, namely, normal and tumor. Then, the adjacency matrices were transformed into topological overlap matrix (TOM), and the corresponding dissimilarity was calculated (1-TOM). Here, we set the soft-thresholding power as 2 (TCGA-KIRC) and 16 (CEO-ccRCC), cut height as 0.25, and minimal module size as 50. Based on the 1-TOM, the same gene expressions were grouped into a gene co-expression module. Then, important modules were selected, and the intersection with DEGs of TCGA-KIRC and GEO-ccRCC datasets was used. Further analysis, such as GO and KEGG analyses, was conducted on genes that overlapped with those obtained by the four abovementioned datasets.

Functional annotation and functional enrichment analyses

The R package clusterProfiler (version 3.16.1) (13), org.Hs.eg.db (version 3.11.4), enrichplot (version 1.8.1), and ggplot2 (version 3.3.3) were used in conducting Gene Ontology (GO) enrichment and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analyses, with adjusted P< 0.05, a cut-off criterion indicating statistical significance.

Screening of the prognostic metabolism-related hub gene signature

We intersected the intersection genes with metabolic genes, which were given by the metabolic pathway based on the KEGG online database. Afterward, we obtained the key metabolic genes and adopted univariate Cox regression analysis to screen hub genes associated with prognosis. We regarded P< 0.05 as a significant difference.

Construction and evaluation of the risk score prognostic model

We obtained the prognostic metabolism-related gene and then performed a Lasso-cox regression analysis to construct a prognostic metabolic-related gene signature. In constructing the risk score prognostic model, we computed the risk score for each patient and divided all the patients into two parts, namely, high-risk patients and low-risk patients, based on the median risk value.

Risk score = \( (\text{Expr}_{\text{gene-1}} \times \text{Coef}_{\text{gene-1}}) + (\text{Expr}_{\text{gene-2}} \times \text{Coef}_{\text{gene-2}}) + ... + (\text{Expr}_{\text{gene-n}} \times \text{Coef}_{\text{gene-n}}) \)

where \( \text{Expr} \) is the expression of the gene in the signature, and \( \text{Coef} \) is the Cox coefficient of the gene.

We investigated the time-dependent prognostic significance of the risk score prognostic model using the R package survivalROC (version 1.0.3) and compared it with the predicted effect of age, T, N, M, grade, and stage.

Verification of the protein level and prognostic values of key genes of the prognostic model

The Human Protein Atlas database (HPA, https://www.proteinatlas.org/), which provides a large amount of transcriptomic and proteomic data in specific human tissues and cells for research, is a valuable database (14). We confirmed the protein level of each key gene between ccRCC and normal tissue using the HPA database, in which immunohistochemistry (IHC) was used to determine protein expression. In addition, we used the UALCAN database (http://ualcan.path.uab.edu/index.html) to confirm the protein level in different stages of ccRCC and normal renal tissues, which provides protein expression analysis option using the data obtained from the Clinical Proteomic Tumor Analysis Consortium (CPTAC) dataset (11). Based on the data obtained from the TCGA database, we used the survival package in R software to explore the prognostic values of key genes and performed Kaplan Meier survival analysis as a box plot graph. Exploring the relationship between disease-free survival (DFS) and the expression of key genes in patients with ccRCC, we used the online tool gene expression profiling interactive analysis (GEPIA, http://gepia.cancer-pku.cn/).

Assessment of the forecast effect on the risk prognostic model

In validating the association between the risk score and patients’ survival time, we used the pheatmap (version 1.0.12) R package to plot the risk plot and assessed the prognostic value between low- and high-risk patients using the Kaplan-Meier survival curve. We used univariate and multivariate Cox regression analysis to assess the associations between the risk score and various clinicopathological parameters in the TCGA-KIRC Cohort using the Forrest plot. Moreover, exploring the relationship between risk score and tumor grade and stage, we plotted the violin figure using the online tool Sangerbox 3.0 (http://vip.sangerbox.com/). To validate the forecast effect, we randomly sampled 70% of the TCGA-KIRC samples by using
the caret (version 6.0.93) R package and formed the validation cohort of this model.

**Prediction of patient’s prognosis and treatment**

We used the R package rms (version 6.2.0) to build a predictive nomogram, including clinicopathological characteristics and risk score. Then we used foreign (version 0.8.80) and survival (version 3.3.1) R packages to calculate the concordance index (C-index) and to plot the calibration curves of the predictive nomogram. We also performed a GSEA analysis to identify enriched terms and selected the top 13 significant pathways visualized as the multiple-GSEA plot.

**Validation of clinical tissue samples by RT-qPCR experiments**

According to the manufacturer’s instructions, we isolated the total RNA of ccRCC tissues and corresponding normal renal tissues using TRIzol (Invitrogen), which was converted into cDNA using PrimeScript™ RT Master Mix (Takara). After cDNA was subjected to reverse transcription PCR using a SYBR-Green master kit (Vazyme, Nanjing, China) on the Applied Biosystems 7500 system, the following cycles were performed: predenaturation at 95°C for 5 min; denaturation at 95°C for 10 s, annealing and extension at 60°C for 34 s; and repeated denaturation, annealing, and extension for 40 cycles. We used β-acting as the housekeeping gene to normalize the relative expression of genes as an endogenous control using the comparative Ct (threshold cycle) method (ΔΔCt). The primers of key genes for the quantitative polymerase chain reaction assay were obtained from Primer Bank, which are shown in Supplementary Table S1.

**Results**

**Identification of DEG and WGCNA**

We divided the samples of the TCGA-KIRC dataset into two groups, namely, normal group and tumor group, and identified the DEGs with \(|\log FC|>1\) and ad. just \(P<0.05\). We screened 3,747 DEGs (1,924 up-regulated and 1,823 down-regulated genes, Figure 1D) from 14,684 genes (Supplementary Table S2). Then, we selected the top 100 DEGs (50 upregulated and 50 downregulated genes) for visualization by heatmap plot (Figure 1A). We used the WGCNA R package to construct weighted gene co-expression modules where each module is assigned with color; a total of 11 modules were included in the TCGA-KIRC (Figure 1B). In visualizing the relationship between each module and two clinical traits (normal and tumor), we plotted the heatmap of the module-trait relationship (Figure 1C). The genes of each module membership are listed in Supplementary Table S3. We found that the turquoise and purple modules were the top two association with clinical traits (MEturquoise module: \(r=0.82, P=7e-153\); MEpurple module: \(r=0.61, P=3e-64\)), and the genes of these two modules were downregulated in ccRCC. The relationships between module membership and gene significance are presented in Figures 1E, F (turquoise module: \(cor=0.93, P<1e-200\); purple module: \(cor=0.72, P=1.8e-19\)).

The GEO-ccRCC dataset consisted of four datasets, and all the samples of the database were divided into normal and tumor groups. We identified 1,344 DEGs from 21,653 genes (650 upregulated and 694 downregulated genes, Figure 2B) with \(|\log FC|>1\) and adjust \(P<0.05\) (Supplementary Table S4). We plotted the heatmap plot for the top 50 upregulated and 50 downregulated genes (Figure 2A). Based on the GEO-ccRCC database, we built a total of four weighted gene co-expression modules used in WGCNA analysis (Figure 2C), and the heatmap of the module-trait relationship visualized the correlation between each module and clinical traits, namely, normal and tumor (Figure 2D). The genes of each module membership are listed in Supplementary Table S5. The blue module was the highest relation with clinical traits (\(r=0.92, P=7e-144\)), and the relationship between module membership and gene significance for the tumor is visualized and shown in Figure 2E.

**Acquisition of overlapping genes and functional enrichment analysis**

We obtained 3,747 DEGs in the TCGA-KIRC dataset, 1,344 DEGs in the GEO-ccRCC dataset, 10,601 and 114 co-expression genes in the turquoise and purple modules of the TCGA dataset, and 1,717 co-expression genes in the blue module of the GEO dataset. We respectively recorded the intersection of the turquoise and purple modules with DEGs in the TCGA-KIRC dataset, DEGs in the GEO-ccRCC dataset, and the blue module in the GEO dataset. Then, we obtained overlapping 1 (a total of 550 genes, Figure 3A) and overlapping 2 (a total of 77 genes, Figure 3D) and performed enrichment GO analysis (Figures 3B, E) and KEGG analysis (Figures 3C, F) on overlapping 1 and overlapping 2. GO enrichment analysis of genes in overlapping 1 showed that the biological process (BP) was primarily enriched in kidney epithelium development, kidney development, and renal system development. In addition, the cellular component (CC) was primarily enriched in the apical part of cell and basolateral plasma membrane, and the molecular function...
(MF) was primarily gathered in the active ion transmembrane transporter activity and active transmembrane transporter activity (Supplementary Table S6). Based on GO analysis of overlapping 2 genes, the BP showed that these genes were primarily enriched in the small-molecule catabolic process, organic acid catabolic process, and carboxylic acid catabolic process. The enrichment of the CC primarily occurred in the peroxisomal matrix and microbody lumen. Moreover, coenzyme binding and aldehyde-lyase activity were more related to these genes in the MF (Supplementary Table S7). For KEGG pathway analysis, carbon metabolism and the HIF-1 signaling pathway were associated with overlapping 1 genes (Supplementary Table

![Figure 1](image_url)

**FIGURE 1**
DEG and WGCNA analysis of TCGA-KIRC dataset. (A) Heatmap of top 100 DEGs between ccRCC and normal renal tissue from TCGA-KIRC dataset. (B) Clustering dendrograms of all the genes of TCGA-KIRC, based on the difference in topological overlap, assigned modules with different colors. (C) The eleven co-expression modules and the module trait between ccRCC and normal renal tissue. (D) Volcano Plot of 14,684 genes of TCGA-KIRC dataset. Green assigned downregulated genes and red assigned up. (E) Module membership in the turquoise module. (F) Module membership in the purple module.
Meanwhile, carbon metabolism, glycolysis/gluconeogenesis, and peroxisome were primarily related to overlapping 2 genes (Supplementary Table S9).

Screening of prognostic metabolism-related hub genes and construction of the risk score prognostic model

As a result of functional enrichment analysis, metabolism and genes in overlapping 1 and 2 have a strong connection. We used the intersection of overlapping 1 and 2 with metabolism-related genes (Figures 4A, D) and adopted univariate Cox regression analysis to screen prognostic metabolism-related genes. In overlapping 1, we screened 13 prognostic metabolism-related genes (ACADSB, ALAD, DEGS1, ECI2, GPT2, GSTM3, HADH, HK2, LDHD, OAT, PFKP, PSAT1, and UPP2; Figure 4B). Then we constructed the risk score prognostic model 1 (AUC=0.694, Figure 4C) with Lasso-cox regression analysis. Meanwhile, we screened six prognostic metabolism-related genes (ABAT, ALDH6A1, CHDH, EPHX2, ETNK2, and FBP1; Figure 4E) in overlapping 2 and performed
Lasso-cox regression analysis to construct the risk score prognostic model 2 (AUC=0.795, Figure 4F). Therefore, we selected the risk score prognostic model 2 and the genes used to construct the model for follow-up analysis.

Verification of prognostic metabolism-related hub genes

Based on the immunohistochemical results from the HPA database (Supplementary Table S10), the protein level of the six prognostic metabolism-related genes in tumors was generally lower than that in normal tissue (Figure 5A). We further verified the protein level of each hub gene in every tumor stage based on the CPTAC dataset (Figure 5B). The results of the boxplot showed that the protein level of the six prognostic metabolism-related genes was significantly downregulated in different stages of ccRCC, compared with normal renal tissues. Kaplan-Meier analyses (Figure 6A) indicated that the low expression level of each of the six hub genes was significantly associated with poor overall survival (OS) of patients with ccRCC (P<0.05). Moreover, the low expression level of ALDH6A1, CHDH, and ETNK2 was related to worse OS (P<0.001). Meanwhile, based on the GEPIA2 database, we
observed that the expression level of ALDH6A1 and FBP1 had a major relationship with worse DFS (P<0.05, Figure 6B), in patients with ccRCC.

Assessment of the predicted effect on the risk score prognostic model

In visualizing the correlation between the survival status and risk score in ccRCC patients, we plotted a risk curve (Figure 7A), on the basis of the TCGA-KIRC dataset. As shown in Figure 7A, patients were divided into high-risk and low-risk groups. The heatmap for the expression of the abovementioned six genes showed that their expression level decreased gradually from the high-risk group to the low-risk group. As the risk score increased, more patients died. Using Kaplan-Meier analysis, we found that the high-risk score was significantly connected related to worse OS (P<0.001, Figure 7B). We used univariate Cox regression and multivariate Cox regression analyses to assess the independent role of the risk score prognostic model. Univariate Cox regression analysis (Figure 7C) indicated that age, grade, stage, T, M, N, and risk score were correlated with OS, and multivariate Cox regression analysis showed that the risk score could serve as an independent prognostic factor (Figure 7D, P<0.001, hazard ratio: 2.033-9.787). Figures 7E, F were violin plots for the risk score of different grades and stages of ccRCC tumor. We observed significant differences in risk scores among different ccRCC tumor grades and stages (P<0.001). The prognostic nomogram for the prediction of 1-, 3-, and 5-year survival in ccRCC is shown in Figure 8A. By using foreign and survival R packages, we calculated the C-index of the TCGA-KIRC dataset (C-index = 0.796) and the validation cohort (C-index = 0.76). Figures 8B, C were calibration curves of the nomogram for predicting patient survival at 3 years and 5 years. We validated the forecast effect using the Kaplan-Meier analysis (P<0.001, Figure 8D), ROC curves (risk score AUC = 0.805, Figure 8E), and the 5-year survival prediction calibration curve (Figure 8F) in the validation cohort.

GSEA analysis and drug sensitivity test

Based on the TCGA-KIRC dataset, we performed a GSEA analysis to investigate the potential underlying mechanism between the high-risk group and the low-risk group. We enriched 178 upregulated pathways (51 in the high-risk group and 127 in the low-risk group). Of the 127 upregulated pathways in the low-risk group, 43 upregulated pathways were significantly different. The top 13 enriched pathways (three in the high-risk group and ten in the low-risk group) are shown in Figure 8H. We used a drug sensitivity test to investigate the sensitive drugs for high-risk patients. The top four major sensitive drugs included Vinblastine sensitivity, ZM.447439, AP.24534, and CGP.60474 (Figure 8I).
RT-qPCR validation of the expression level of the six hub genes

RT-qPCR on 20 paired ccRCC and normal tissue samples (Supplementary Table S11) showed that the expression level of the six hub genes in tumor tissues was generally lower than that of normal renal tissues (P<0.05, Figure 8G). The six hub genes were significantly downregulated in tumor samples compared with normal samples. This result was consistent with the validated expression level of the six genes that constructed our risk score prognostic model based on an online database.

Discussion

With the accumulation of cancer research, the link between cancer and various metabolic changes has been revealed. Goldblatt and Cameron obtained transplantable cancer cells from heart fibroblasts through oxygen deficiency experiments (16). Warburg described that the origin of cancer cells from normal tissue cells has two phases: the first phase is an irreversible injury of respiration, and in the second phase the injured cells maintain their structure and energy supply by fermentation energy. Finally, the highly differentiated body cells are converted into undifferentiated cells and grown wildly (17). The fermentation energy that Warburg described in cancer is the earliest mention of metabolic reprogramming, which is not only the beginning but also the propelling of cancers. In addition, he did not fully comprehend the discovery at that time. Approximately 85% of RCC arise from tubular epithelial cells (18). One of the characteristics of RCC is the mutation of genes that are involved in metabolic pathways, including aerobic glycolysis; fatty acid metabolism; and the metabolism of tryptophan, glutamine, and arginine (19). Therefore, RCC is generally regarded as a metabolic disease, and the major risk factors include aerobic glycolysis and the mutation of metabolic genes (20, 21). In this study, we characterized ccRCC to explore the prognostic prediction model and treatment of ccRCC. We performed bioinformatics analysis to screen out prognostic metabolism-related key genes and construct the risk score prognostic model using these key genes.

To avoid errors caused by insufficient sample size, the ccRCC samples in this study were obtained from TCGA and GEO databases, a total of 965 samples. We used DEG analysis and WGCNA to screen the clinical traits related to DEGs. Based on the correlation coefficient of genes, WGCNA, a network
analysis method, can identify biologically relevant modules and key genes (5), which are correlated with clinical traits. Such modules and key genes may serve as biomarkers for detection or treatment (22). Therefore, WGCNA has unique advantages in exploring the relationship of clinical traits related to modules. We found that the coexpression modules closely related to clinical traits were downregulated in ccRCC, and the functional enrichment analysis showed that the genes in those modules were related to metabolism. This result is in line with our expected results. Finally, the genes of the risk score prognostic model that we constructed included ABAT, ALDH6A1, CHDH, EPHX2, ETNK2, and FBPI, which were downregulated in ccRCC. Multivariate Cox regression analysis showed that the risk score of the prognostic model could serve as an independent prognostic factor. To compare the predicted effect of the risk score prognostic model with that of age, T, N, M, grade, and stage, we plotted ROC curves by the TCGA-KIRC cohort and the validation cohort. The results of ROC curves showed that the predicted effect of the risk score prognostic model was similar to that of clinical grade and stage. The prognostic nomogram combined abundant factors, including age, T, N, M, grade, stage, and risk score. By using the nomogram, we can make a more accurate prediction of 1-, 3-, and 5-year survival of ccRCC patients.

The six key genes are involved in various metabolic reactions, including amino acid metabolism, choline metabolism, and glucose metabolism. Most of the genes are involved in cancers. The 4-aminobutyrate aminotransferase (ABAT) encodes γ-aminobutyric acid (GABA) transaminase, which is a key enzyme for catabolism GABA, a major inhibitory neurotransmitter, within the mitochondrial matrix. ABAT plays an important role in neurometabolic disorders (23). The deficiency of ABAT mediates the destruction of the GABAergic system, and patients present corresponding clinical manifestations of elevated GABA levels (24). For breast cancer, the loss of ABAT expression could promote the potency of tumorigenesis and metastasis (25), which could be a predictive biomarker for endocrine therapy resistance (26). The results of our study show that the expression of ABAT was downregulated in ccRCC and related with poor DFS of patients. GSEA analysis shows that the beta alanine metabolism pathway associated with ABAT was upregulated in the low-risk group. A previous study indicated that ABAT and aldehyde dehydrogenase 6 family member A1 (ALDH6A1) worked as a tumor suppressor (27) in ccRCC, thereby suppressing tumorigenic capability. In this study, the downregulation of the expression of ALDH6A1, an amino acid metabolism-related gene, was significantly linked with worse OS and DFS in patients. Meanwhile, ALDH6A1 was identified as a potential molecular signature for hepatocellular carcinoma (28), prostate cancer (29), and muscle insulin resistance in type 2 diabetes mellitus (30). Therefore, ALDH6A1 may be a potential key target for regulating ccRCC.
metabolism. Based on the results of the GSEA analysis, ABAT and ALDH6A1 function in the valine leucine and isoleucine degradation pathways, which are upregulated in the low-risk group. In addition, ABAT plays a role in the butanoate metabolism and alanine aspartate and glutamate metabolism pathways. The physiological role of Human choline dehydrogenase (CHDH) is to regulate the concentration of choline and glycine betaine, and CHDH is primarily located in the liver and kidney (31). Moreover, CHDH plays a pivotal role in mitophagy (32). Based on our results, genes of glycine serine and threonine metabolism pathway, including CHDH, and the downregulation of CDHD in ccRCC, were related to worse OS and DFS in patients. Soluble epoxide hydrolase (EPHX2, sEH) serves as a principal enzyme for the metabolism of epoxyeicosatrienoic acids (33), and it is related to cell apoptosis (34). For metabolic diseases, EPHX2 may be a potential therapeutic target (35). In prostate cancer and hepatocellular carcinoma, EPHX2 was downregulated, which was significantly correlated with the progression of tumors (36, 37). Based on the enrichment results of GSEA, EPHX2 is intimate with the peroxisome pathway. Furthermore, the mechanism of CHDH and EPHX2 in ccRCC was deficient. Ethanolamine kinase 2 (ETNK2) was also reported in tumors. The upregulation of ETNK2 enhances hepatic metastasis such as gastric cancer (38). However, ETNK2 was downregulated in our research for ccRCC with a poor OS of patients. Fructose-1, 6-bisphosphatase 1 (FBP1), a rate-limiting enzyme for gluconeogenesis (39), plays a critical role in tumor initiation and progression of ccRCC. FBP1 has two major mechanisms that inhibit ccRCC progression: first, FBP1 can inhibit a potential Warburg effect; second, FBP1 can interact with the HIF inhibitory domain and inhibit nuclear HIF function (40). Targeting FBP1 has been an emerging therapeutical target for cancers (39). Thus, this study aimed to explore more therapeutical targets for ccRCC via screening the prognostic metabolism-related genes.

Using the six prognostic metabolism-related genes, we constructed a risk score prognostic model and divided patients with ccRCC into high- and low-risk groups based on the risk score of each patient. We found that the patients in the high-risk group have poor OS, and lass survival time. Based on the violin plots, we discovered that the risk scores on each grade and stage
of ccRCC were significantly different, and the risk scores displayed a significantly positive correlation with the degree of malignancy of ccRCC. The results of multivariate Cox regression analysis showed that the risk score of the prognostic model could serve as an independent prognostic factor. Based on the prediction of clinical chemotherapeutic response analysis, we screened four drugs, namely, Vinblastine, ZM.447439, AP.24534, and CGP.60474, which may be more sensitive for...
patients in the high-risk group. Vinblastine is a dimeric alkaloid isolated from the Madagascar periwinkle plant, which exhibits significant cytotoxic activity, and it is used as an antineoplastic agent in antitumor therapy (41). For our analysis, Vinblastine is the principal sensitive drug for high-risk group patients. Mitotic Aurora kinases are essential for accurate chromosomal segregation during cell division. As an Aurora-selective ATP-competitive inhibitor, ZM.447439 can interfere with the spindle integrity checkpoint and chromosomal segregation (42). In addition, Vinblastine and ZM.447439 function by interfering with cancer cell proliferation. AP.24534 and CGP.60474 are kinase inhibitors. Ponatinib, AP.24534, is a third-generation tyrosine kinase receptor inhibitor (43), and CGP.60474 is an inhibitor of cyclin-dependent kinase (44). Tyrosine kinase inhibitors are novel therapies for ccRCC treatment (45–50). Tyrosine kinases are signaling molecules, and tyrosine kinase inhibitors have become a successful class of drugs in the treatment of ccRCC. Thus, we might consider that these four identified sensitive drugs could be potential treatments for ccRCC, and we believe that novel drugs worked by regulating the pathway of cellular metabolism will appear increasingly in the near future.

This research also has some limitations. First, this is a retrospective study, we need more clinical samples to improve our findings and the predicted effect of the risk score prognostic model. Second, the molecular mechanisms of the six key genes need to be further elucidated in vivo and in vitro experiments for ccRCC clinical applications.

Conclusions

In conclusion, based on the metabolic reprogramming characteristics in ccRCC and combined with WGCNA analysis, we identified six metabolism-related genes, which could be potential treatment targets for ccRCC. Furthermore, we constructed a risk score prognostic model, the risk score of which constitutes an effective independent prognostic factor. By including the risk score, the nomogram can help us make a more accurate prediction of patient survival. The improvement of the prognostic model may improve the outcome prediction for ccRCC patients in the future.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

Author contributions

Conception and design: JL, CL, QZ, and LD. Administrative support: JL and CL. Provision of study materials or patients: CL, QZ, and LD. Collection and assembly of data: QDZ and CN. Data analysis and interpretation: CL, QZ, TZ, and LD. Manuscript writing: All authors. All authors contributed to the article and approved the submitted version.

Funding

This study was supported by the National Natural Science Foundation of China (Grant no. 82002718); and the Jiangsu Natural Science Foundation (Grant no. BK20191077).

Acknowledgments

The authors highly appreciate all the members of Biomathematics for improvement of this manuscript.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.982426/full#supplementary-material
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