AURKB, CHEK1 and NEK2 as the Potential Target Proteins of Scutellaria barbata on Hepatocellular Carcinoma: An Integrated Bioinformatics Analysis

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Objective: We aim to explore the potential anti-HCC mechanism of Scutellaria barbata through integrated bioinformatics analysis.

Methods: We searched active ingredients and related targets of Scutellaria barbata via TCMSP database, PubChem and SwissTargetPrediction database. Then, we identified HCC disease targets from GEO dataset by WGCNA. Next, the intersected targets of disease targets and drug targets were input into STRING database to construct PPI networking in order to obtain potential therapeutic targets of Scutellaria barbata. Cytoscape software was used to carry out network topology analysis of potential targets. We used the R package for GO analysis and KEGG analysis. Finally, we used AutoDock vina and PyMOL software for molecular docking.

Results: Sixteen active components from Scutellaria barbata were lastly selected for further investigation. A total of 442 component targets were identified from 16 active ingredients of Scutellaria barbata after the removal of duplicate targets. GSE45436 was selected for construction of WGCNA and screening of differentially expressed genes. A total of 354 genes were up-regulated in HCC samples and 100 were down-regulated in HCC patients. Twenty-one common genes were obtained by intersection and 10 critical targets were filtered for further investigation. The enrichment analysis showed that cell cycle, DNA replication, p53 signaling pathway were mainly involved. The molecular docking results showed that 4 potential combinations were with the best binding energy and molecular interactions.

Conclusion: AURKB, CHEK1 and NEK2 could be the potential target proteins of Scutellaria barbata in treating HCC. Cell cycle, DNA replication, p53 signaling pathway consist of the fundamental regulation cores in this mechanism.

Keywords: Scutellaria barbata, hepatocellular carcinoma, effector mechanism, network pharmacology, bioinformatics analysis, molecular docking

Introduction

Liver cancer is one of the malignant tumors with the highest incidence rates in the world, of which the incidence rate increases by approximately 700,000 patients worldwide each year, and over half of the global incidence and mortality of liver cancer occurs in China. Hepatocellular carcinoma (HCC) is the most common primary liver cancer subtype that contributes majority incidence of liver cancers worldwide. Epidemiologically, the number of NASH-related HCC cases...
is increasing, while that of HBV- or HCV-related cases is declining and that of alcohol-related cases remaining unchanged. Nevertheless, the burden of HCC would probably increase worldwide in the future as the population growth and aging expected in the next few decades. Hence, it is of great significance to explore new therapeutic targets for the treatment of HCC. Besides surgery, chemotherapy, interventional radiology, immunotherapy, molecular-targeted therapy and liver transplant, herbal products in Traditional Chinese Medicine (TCM) now have been considered as an upcoming therapeutic approach for prevention and treatment of HCC.5

Scutellaria barbata (Ban Zhi Lian in Chinese) is used as an immunomodulatory and antitumor agent in TCM.6 Scutellaria barbata is the whole dried grass of a labiate plant Scutellaria barbata D. Don, and has an extensive application for efficacy enhancement in cancer treatment in China and Asian countries.7 Effective ingredients of Scutellaria barbata have shown growth-inhibitory effects in various types of cancer, including liver cancer.8–10

Network pharmacology, first proposed in 2007, has become an effective assistant to systematically provide a novel strategy to research the mechanism of TCM formula with multiple compounds.11 Weighted gene co-expression network analysis (WGCNA) is a new tool that is adopted to identify co-expressed modules and hub genes.12 WGCNA can extract gene co-expression modules and contact it to the clinical feature.13 At present, WGCNA can be useful to identify candidate biomarkers or therapeutic targets of various diseases, and compare differentially expressed genes (DEGs) in order to help investigating the interactions among genes in different modules.14 Molecular docking technique is adopted to realize the drug-receptor interaction and orientation and approach of drug molecules to the target site.15

In this study, we aim to explore the potential anti-HCC mechanism of Scutellaria barbata through network pharmacology and bioinformatics analysis. Furthermore, molecular docking was also performed to validate the binding interaction of the components in the binding pocket of the target proteins.

Materials and Methods

The flow chart of this study is shown in Figure 1.

Screening and Preprocessing of Active Components from Scutellaria barbata

By searching the Traditional Chinese Medicine Systems Pharmacology Database and Analysis Platform (TCMSP, https://tcmspw.com/tcmsp.php) database, the qualified components were collected. “Scutellaria barbata” was searched as Herb name in TCMSP, oral bioavailability (OB) ≥30% and drug-like (DL) ≥0.25 were set in the Ingredients module, the selected molecule names were researched in PubChem database (https://pubchem.ncbi.nlm.nih.gov/) and the chemical components without PubChem CID were removed. The canonical SMILES strings and 3D structures of the related components were obtained from the PubChem database by PubchemCID and all the chemical structures were inputted into Excel as backup.

Screening of Molecular Composition Target

The canonical SMILES strings obtained from the PubChem database were uploaded to the SwissTargetPrediction server (http://www.swisstargetprediction.ch/) for molecular composition target prediction. The platform species is set to “Homo sapiens”, and the predicted results of the platform were collected and classified. All predicted results are stored as CSV format. Furthermore, all of the results were de-reprocessed, transferred to Excel format, and a molecular library was established.

Sources of Clinical Data and Screening of Differentially Expressed Genes

In order to further explore the clinical significance of the predicting target of Scutellaria barbata, the key word “liver cancer” was searched in Gene Expression Omnibus (GEO) datasets (https://www.ncbi.nlm.nih.gov/gds), and the datasets with more than 100 samples were selected. In this study, the whole genome genes were detected, and the DEGs in the dataset were analyzed and screened. $|\log \text{FC}|\geq 1$ was used as the filtering threshold criteria of logFC and adj-P-value <0.05 as the corrected threshold of P-values. LogFC>1 was regarded as up-regulated genes in tumor samples while LogFC<-1 would be considered as down-regulated genes in tumor samples. The heatmap and volcano plot of the screened genes were drawn by R software (Version 3.6.3).
Figure 1 Workflow for exploring the molecular mechanisms of Scutellaria barbata against Hepatocellular carcinoma.
Construction of Gene Co-Expression Networks

The R package “WGCNA” was used for finding clusters (modules) of highly correlated genes, for summarizing such clusters using the module eigengene or an intramodular hub gene. In this study, WGCNA construction and module detection were performed using the “WGCNA” R package and top 25% genes with the largest variance differences were used to construct weight gene co-expression networks in GSE45436. First, the gene expression data were used to obtain a Pearson correlation matrix between the genes. A soft threshold $\beta$ can improve strong correlations and weaken weak correlations. The power of $\beta = 6$ (scale-free $R^2=0.85$) was selected as the soft-thresholding parameter to ensure a scale-free network. Next, the linkage hierarchical clustering was produced by the topological overlap measure to carry out module detection and acquire the cluster dendrogram. Under the condition of setting minimum cluster size as 30, 16 modules were generated via Dynamic Tree Cut algorithm.

Intersection of WGCNA Co-Expression Genes and DEGs

In this study, the positive correlation modules that generated by WGCNA could be considered as up-regulated genes in tumor samples, while the negative correlation modules were regarded as down-regulated genes in tumor samples. The hub module with the highest positive correlation in normal and tumor patients matched with the differentially up-regulated genes, and the hub module with the highest negative correlation was intersected with the differentially down-regulated genes, the result was mapped into Venn diagram, and the overlapping genes obtained were used as HCC-related genes in this study.

Searching for Overlapping Targets and Key Targets of Scutellaria barbata and HCC

The overlapping targets of Scutellaria barbata and HCC were found, and a Venn diagram was drawn. Next, the overlapping targets were inputted into String (http://string-db.org/, version 11) for protein–protein interaction (PPI) analysis. The String database is a biological database for prediction of PPIs, with data from several sources. For further exploration, the minimum required confidence score was medium confidence (0.400), and disconnected nodes in the network would be hidden. The results were exported in tab-separated value (TSV) format once the protein interaction relationship was obtained. The PPI network was constructed via utilizing the network visualization software Cytoscape 3.8.1 (http://www.cytoscape.org/). CytoHubba is a tool used to identify hub objects and subnetworks from a complex interactome. CytoHubba provides 11 topological analysis methods including degree centrality (DC), Edge Percolated Component (EPC), Maximum Neighborhood Component (MNC), Density of Maximum Neighborhood Component (DMNC), Maximal Clique Centrality (MCC) and centralities based on shortest paths, such as Bottleneck (BN), closeness centrality (CC), betweenness centrality (BC), EcCentricity, Radiality, and Stress. Following the construction of a PPI network, hub genes were identified with the MCC algorithm. MCC algorithm can accurately identify hub genes in HCC.

GO and KEGG Enrichment Analysis Potential Targets of Scutellaria barbata

Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis was performed to examine the enrichment of terms. The R package “ggplot2”, “org.Hs.eg.db”, “clusterProfiler” and “enrichplot” was used for GO and KEGG enrichment analysis. Enriched GO terms and KEGG pathways were identified with the cut-off criterion of $P < 0.05$.

Construction of Active Component-Antihepatoma Target-Pathway Network of Scutellaria barbata

The network of active component of Scutellaria barbata - anti-hepatoma target - pathway was constructed by Cytoscape software. The node in the network represents Scutellaria barbata, active components of Scutellaria barbata, key target genes, gene enrichment pathway. The edge in the network is used to connect Scutellaria barbata and active components, active components and key target genes, key target genes and enrichment pathway. The whole network shows the relationship between Scutellaria barbata, active components, anti-hepatoma target and pathway, and explores the anticancer mechanism of Scutellaria barbata by constructing this network.
Molecular Docking
The core target protein structures were downloaded from Protein Data Bank (PDB) database (http://www.rcsb.org/). The water molecules were removed and the original ligands were separated via PyMol 1.8 software. After preservation, they were transferred into Autodock Tools 1.5.6 software and converted to “pdbqt” format. The mol2 structure of qualified components downloaded from TCMS database, and transferred to Autodock Tools 1.5.6 software in order to convert to “pdbqt” format. The abovementioned two “pdbqt” format files were input into Autodock-vina 1.1.2 software for molecular docking. PyMol software was used to visualize the docking results and establish docking interaction pattern.

Prognostic Value of the Potential Target Genes of Scutellaria barbata
GEPIA2 (Gene Expression Profiling Interactive Analysis, http://gepia2.cancer-pku.cn/) was used to explore the prognostic values of the potential target genes in patients with HCC. We used the “Survival Analysis” module of GEPIA2 to obtain the OS (Overall Survival) and DFS (Disease Free Survival) Kaplan-Meier curve of potential target genes in HCC. The patients were grouped into high-expression and low-expression according to the median expression of the potential target genes. Cutoff-high (50%) and cutoff-low (50%) values were used as the expression thresholds for splitting the high-expression and low-expression cohorts.

Table 1 Detail Information of 16 Compounds of Scutellaria barbata

| No. | Components | PubChemCID | CAS       | Molecular ID   | OB%  | DL   |
|-----|------------|------------|-----------|----------------|------|------|
| 1   | 5,7,4’-trihydroxy-6-methoxyflavanone | 5322074 | 94942-49-1 | MOL012245 | 36.63 | 0.27 |
| 2   | 5-hydroxy-7,8-dimethoxy-2-(4-methoxyphenyl)chromone | 14353376 | 57096-03-4 | MOL012248 | 65.82 | 0.33 |
| 3   | 7-hydroxy-5,8-dimethoxy-2-phenyl-chromone | 14825644 | N/A       | MOL012250 | 43.72 | 0.25 |
| 4   | Baicalin   | 64982      | 21967-41-9 | MOL002776 | 40.12 | 0.75 |
| 5   | CLR        | 5997       | 57-88-5   | MOL000953 | 37.87 | 0.68 |
| 6   | Beta-sitosterol | 222284 | 83-46-5   | MOL000358 | 36.91 | 0.75 |
| 7   | Rivularin  | 13889022   | 70028-59-0 | MOL012266 | 37.94 | 0.37 |
| 8   | Stigmasterol | 5354503 | 915-05-9  | MOL0001973 | 40.39 | 0.85 |
| 9   | Stigmasterol acetate | 5280794 | 83-48-7   | MOL000449 | 43.83 | 0.76 |
| 10  | Dinatin    | 5281628    | 1447-88-7 | MOL001735 | 30.97 | 0.27 |
| 11  | 24-Ethylcholest-4-en-3-one | 5484202 | 67392-96-5 | MOL001755 | 36.08 | 0.76 |
| 12  | Salvigenin | 161271     | 19103-54-9 | MOL002915 | 49.07 | 0.33 |
| 13  | Rhamnazin  | 5320945    | 552-54-5  | MOL000351 | 47.14 | 0.34 |
| 14  | Sitosterol | 12303645   | 83-46-5   | MOL000359 | 36.91 | 0.75 |
| 15  | Moslosoflavone | 188316 | 3570-62-5 | MOL008206 | 44.09 | 0.25 |
| 16  | Quercetin  | 5280343    | 117-39-5  | MOL00098  | 46.43 | 0.28 |

Result
Identification of Targets of Scutellaria barbata
A total of 94 components from Scutellaria barbata were found from the TCMS database. 22 active components were available under the condition of OB≥30% and DL≥0.25. Active components without PubChem CID were excluded due to the absence of canonical SMILES strings and 3D structures, which were significant for predicting the component targets. Sixteen active components were lastly selected for further investigation (Table 1). They were 7-hydroxy-5,8-dimethoxy-2-phenyl-chromone (MOL012250), 5-hydroxy-7,8-dimethoxy-2-(4-methoxyphenyl)chromone (MOL012248), 5,7,4’-trihydroxy-6-methoxyflavanone (5,7,4’-trihydroxy-6-methoxyflavanone, MOL012245), Baicalin (MOL002776), CLR (MOL000953), beta-sitosterol (MOL000358), rivularin (MOL012266), Sitosterylacetate (MOL001973), Stigmasterol (MOL000449), Dinatin (MOL001735), 24-Ethylcholest-4-en-3-one (MOL001755), Salvigenin (MOL002915), Rhamnazin (MOL0001973), sitosterol (MOL000359), Moslosoflavone (MOL008206), quercetin (MOL000098). Four hundred and forty-two component targets were identified from 16 active ingredients of Scutellaria barbata via SwissTargetPrediction database after the removal of duplicated targets. According to the prediction result of SwissTargetPrediction, quercetin may be highly combined with 100 targets such as NOX4, AVPR2, AKR1B1, XDH, MAOA, IGF1R, FLT3,
CYP19A1, EGFR, etc. Rhamnazin may have a high degree of binding with 74 targets such as XDH, AKR1B1, ABCG2, CA7, CA12, etc. Beta-sitosterol may have a high degree of binding with 19 targets such as AR, HMGCR, CYP51A1, etc. Sitosterol may have high binding degree with 46 targets such as AR, HMGCR, CYP51A1 and NPC1L1, while Stigmasterol may have high binding degree with 45 targets such as AR, NPC1L1, HMGCR, CYP51A1 and CYP19A1. Further details can be viewed in Supplementary File 1.

Identification of Hub Targets of HCC

GSE45436 was selected for this study, which was published on July 31, 2014, and the final data was updated on September 11, 2019. The GSE45436 dataset contains 134 samples, including 93 tumor samples and 41 normal control non-tumor samples. The expression profiles of 23,518 genes in normal samples (n=41) and HCC samples (n=93) were selected to identify the DEGs and construct gene co-expression networks. Four hundred and fifteen up-regulated genes and 797 down-regulated genes were detected (Figure 2A). The top 50 up-regulated and down-regulated DEGs were shown in the heatmap (Figure 2B). The power of $\beta=6$ (scale-free $R^2=0.85$) was selected as the soft-thresholding parameter to ensure a scale-free network (Figure 2C). Dynamic Tree Cut algorithm can identify the module, and the gene expression values in the module are in similarity. Highly similar modules were merged, and a total of 16 co-expression modules were identified. The modules contained at least 54 genes and 2056 genes at most, each module was marked with one color (Figure 2D). Module-trait relationships are shown in Figure 2E. Each row corresponds to a module eigengene, while each column corresponds to a clinical trait. The correlation coefficient (upper number) and corresponding P-value (lower number) in each cell resulted in the correlation between the gene module and the clinical trait. Black module ($r=-0.83$, $p=5e-35$) shows the highest negative correlation between the normal samples and the HCC samples. Turquoise module ($r=0.78$, $p=3e-28$) shows the highest positive correlation between the normal samples and the HCC samples. Hence, the genes in black module and turquoise module can be regarded as the down-regulated and up-regulated hub genes, respectively. The overlapping genes obtained by merging the DEGs in normal and tumor patients with turquoise and black module-related target genes could be recognized as the key disease-related genes. As shown in Figure 3A and B, a total of 454 key disease-related genes have been identified. Among all, 354 genes were up-regulated in HCC samples and 100 were down-regulated in HCC patients.

Construction of the Potential Anti-HCC Target Genes of Scutellaria barbata

By merging the target genes corresponding to the active components of Scutellaria barbata and the key disease-related genes, we found 21 common genes (CETP, CYP26A1, TYMS, NEK2, PARP1, AKR1C3, SQLE, NUAK1, AURKB, AURKA, POLA1, FEN1, KIF11, CHEK1, CDK1, NQO1, FABP5, IMPDH, TOP2A, AKR1B10, CDC25B) which were highly possible the key targets for Scutellaria barbata to anti-HCC (Figure 3C). However, only two genes were down-regulated, other 19 genes were up-regulated in HCC patients. The 19 up-regulated targets were inputted into the STRING database to build a PPI network (Figure 4A). According to the MCC and DC, 10 critical nodes (TYMS, NEK2, AURKB, AURKA, POLA1, FEN1, KIF11, CHEK1, CDK1, TOP2A) were filtered as key targets for further molecular docking (Figure 4B). Active component-anti-hepatoma target-pathway network of Scutellaria barbata was built to elucidate intuitive interactions in Scutellaria barbata (Figure 5). As we cross verified these 10 critical target genes with the 16 approved components of Scutellaria barbata (Table 2), we found that AURKB and TOP2A were targeted by half of the components. MOL000098, MOL000351 and MOL012250 were the components of most target genes, while MOL001755 was predicted not to target any of the 10 critical target genes.

GO and KEGG Enrichment Analysis

The last 10 critical genes that were filtered as the key targets were selected to perform GO and KEGG enrichment analysis. According to P value <0.05, the top ten items of biological process (BP) of GO enrichment were mitotic centrosome separation, centrosome separation, centrosome cycle, histone phosphorylation, microtubule organizing center organization, regulation of mitotic nuclear division, regulation of centrosome cycle, nuclear division, regulation of nuclear division, organelle fission. And the top ten items of cellular component (CC) of GO analysis were chromosomal region, spindle microtubule, condensed chromosome, condensed nuclear chromosome, mitotic spindle, spindle, spindle pole, microtubule,
Figure 2 Identification of hub targets of HCC. (A) Volcano plot of DEGs, (B) Heatmap of the top 50 up-regulated and down-regulated DEGs, (C) Analysis of the scale-free network coefficient R-squared for the soft threshold, (D) A cluster dendrogram was built based on the dissimilarity of the topological overlap, (E) Module−trait relationships.
midbody, spindle pole centrosome. The top ten items of molecular function (MF) of GO analysis were histone kinase activity, protein serine/threonine kinase activity, catalytic activity, acting on DNA, protein serine/threonine/tyrosine kinase activity, magnesium ion binding, RNA–DNA hybrid ribonuclease activity, sequence–specific mRNA binding, translation repressor activity, mRNA regulatory element binding, endodeoxyribonuclease activity, producing 5′–phospho-monoesters (Figure 6A and B). The results of the GO analysis elucidated that *Scutellaria barbata* treatment of HCC was strongly associated with cell division pathway. According to the results of R package, a total of 13 pathways were screened by KEGG analysis, involving DNA
Figure 4 PPI analysis. (A) PPI network of the 19 up-regulated targets without disconnected nodes. (B) 10 up-regulated targets that were screened according to the MCC and DC.
replication, p53 signaling pathway, Progesterone-mediated oocyte maturation, Cell cycle, Oocyte meiosis, Cellular senescence, Non-homologous end-joining, Viral carcinogenesis, Human immunodeficiency virus 1 infection, One carbon pool by folate, Antifolate resistance, Base excision repair, Pyrimidine metabolism (Figure 7A and B). The KEGG analysis via DAVID showed 5 pathways were involved, including Cell cycle, DNA replication, p53 signaling pathway, Pyrimidine metabolism and Oocyte meiosis.

Molecular Docking for Computer Virtual Verification

The 10 key targets in Figures 4B and 16 core components screened from Scutellaria barbata were performed by molecular docking, the basic information of ligands and proteins are shown in Table 3. When the lowest binding energy is less than -5.0 kJ ·mol-1, it indicates that the docking effect between molecular ligand and receptor protein is good, and the lower the binding energy is, the better the binding capacity between molecule and protein is. The results showed that the majority of the binding energy were less than −5.0 kJ ·mol-1, which revealed benign docking effect between drug components and target proteins, except for the binding energy between MOL000953, MOL000358, MOL001973, MOL000449 and MOL000359 and AURKA, as well as the binding energy between MOL000953, MOL000358, MOL001973, MOL000449, MOL000359, MOL001755 and FEN1 (Figure 8A).

Among those with better binding energy, the binding energy of AURKB to MOL000953, MOL000358, MOL000449, MOL001755 and MOL000359 were less than that of the original ligand. The binding energy of TOP2A to MOL002776, MOL000953, MOL000358, MOL000449, MOL001735, MOL000351, MOL000359 and MOL000098 were less than that of the original ligand. The binding energy of CHEK1 to MOL002776 and MOL001755 were less than that of the original ligand. The binding energy of TYMS to MOL012250, MOL002776, MOL000953, MOL000358, MOL000449, MOL018107 and MOL000359 were less than that of the original ligand. The binding energy of NEK2 to MOL002776, MOL000358, MOL000098, MOL000449, MOL008206, MOL001755, MOL000351 and MOL000359 were less than that of the original ligand. The binding energy of POLA1 to all of the drug components except MOL001973 were less than that of the original ligand. The results above all showed that these target proteins had better ability to combine with the components of Scutellaria barbata, thus these target proteins may be the potential anti-HCC mechanism of Scutellaria barbata. Further, based on
binding energy and molecular interactions such as hydrogen bonds, we selected the top 4 potential combinations with the best binding energy and molecular interactions. Finally, we found the following combinations, including MOL000449 and AURKB (Figure 8B), MOL002776 and CHEK1 (Figure 8C), MOL002776 and NEK2 (Figure 8D), MOL000098 and CHEK1 (Figure 8E). AURKB, CHEK1 and NEK2 could be the top 3 potential target proteins of Scutellaria barbata in treating HCC.

Table 2 Cross Verification of the 10 Critical Target Genes and Compounds of Scutellaria barbata

| No. | Components in Scutellaria barbata | Critical Targets Predicted in Swiss Target Prediction Database |
|-----|----------------------------------|---------------------------------------------------------------|
| 1   | 5,7,4′-trihydroxy-6-methoxyflavanone | AURKA, CHEK1, CHEK1 |
| 2   | 5-hydroxy-7,8-dimethoxy-2-(4-methoxyphenyl) chromone | AURKB, CDK1 |
| 3   | 7-hydroxy-5,8-dimethoxy-2-phenyl-chromone | NEK2, AURKB, FEN1, CDK1, TOP2A |
| 4   | Baicalin | TYMS, POLA1 |
| 5   | CLR | TOP2A |
| 6   | Beta-sitosterol | TOP2A |
| 7   | Rivularin | NEK2, TOP2A |
| 8   | Sitosteryl acetate | AURKB, POLA1 |
| 9   | Stigmasterol | TOP2A |
| 10  | Dinatin | NEK2, AURKB, CDK1 |
| 11  | 24-Ethylcholest-4-en-3-one | / |
| 12  | Salvigenin | AURKB, CDK1 |
| 13  | Rhamnazin | NEK2, AURKB, CDK1, TOP2A |
| 14  | Sitosterol | TOP2A |
| 15  | Moslosoflavone | AURKB, CDK1 |
| 16  | Quercetin | NEK2, AURKB, CDK1, TOP2A |

Prognostic Value of AURKB, CHEK1 and NEK2

As we considered that AURKB, CHEK1 and NEK2 could be the top 3 potential target proteins of Scutellaria barbata in treating HCC. We further explored the prognostic value of AURKB, CHEK1 and NEK2 in HCC. As shown in Figure 9, high expression of AURKB, CHEK1 and NEK2 were consistently linked to poor prognosis of OS and DFS. High expression of the AURKB was related to poor prognosis for HCC in the aspect of OS (Figure 9A, P=0.029) and DFS (Figure 9B, P=0.048). High expression of the CHEK1 was related to poor prognosis for HCC in the aspect of OS (Figure 9C, P<0.001) and DFS (Figure 9D, P<0.001). High expression of the NEK2 was related to poor prognosis for HCC in the aspect of OS (Figure 9E, P<0.001) and DFS (Figure 9F, P<0.001).

Discussion

Scutellaria barbata is a plant of Scutellaria in Labiatae, which has the functions of clearing heat and detoxification, promoting blood circulation and removing blood stasis, reducing swelling and relieving pain in TCM. A large number of modern pharmacological studies have confirmed that Scutellaria barbata had anti-tumor effect. Zhang et al showed that Scutellaria barbata could inhibit the proliferation of A2780 cells and induce apoptosis through mitochondrial pathway, thus playing the role of anti-ovarian cancer.20 Dai et al revealed that Scutellaria barbata inhibited invasion of HCC via MMP/TIMP in vitro.21 Scutellaria barbata polysaccharides inhibited the growth of HCC in mice and affected serum proteomic profiling.6 Extraction of Scutellaria barbata inhibited the tumor growth through down-regulation of Treg cells and manipulating Th1/Th17 immune response in hepatoma H22-bearing mice.22

Although Gong et al have performed a network pharmacology to explore the mechanism of Scutellaria barbata for treating HCC, and they concluded that Scutellaria barbata exerted its anti-hepatoma effect by targeting

Table 3 Information of Proteins Involved in Molecular Docking

| No. | Target Proteins | PDB-ID | Ligand-ID |
|-----|-----------------|--------|-----------|
| 1   | CDK1            | 3LFS   | A07       |
| 2   | AURKB           | 4AF3   | VX6       |
| 3   | TOP2A           | 1ZKX   | ADP       |
| 4   | CHEK1           | 4QYE   | 3DL       |
| 5   | AURKA           | 5DR6   | 5E1       |
| 6   | KIF11           | 3L9H   | EMQ       |
| 7   | FEN1            | 6CVR   | AMP       |
| 8   | TYMS            | 5X66   | MTX       |
| 9   | NEK2            | 2XK8   | 5R1       |
| 10  | POLA1           | 4Q5V   | 2ZE       |
**Figure 6** The results of GO enrichment analysis. (A) The barplot of GO enrichment analysis, (B) The bubble plot of GO enrichment analysis. “Gene ratio” means the percentage of the genes that enriched in corresponding pathway in all the genes that annotated in GO dataset.
Figure 7 The results of KEGG pathways enrichment analysis. (A) The barplot of KEGG analysis, (B) The bubble plot of KEGG analysis. "Gene ratio" means the percentage of the genes that enriched in corresponding pathway in all the genes that annotated in GO dataset.
Figure 8 The results of molecular docking. (A) Heatmap of binding energy between active components and target proteins, (B) Visualization of molecular docking between MOL000449 and AURKB, (C) Visualization of molecular docking between MOL002776 and CHEK1, (D) Visualization of molecular docking between MOL002776 and NEK2, (E) Visualization of molecular docking between MOL000098 and CHEK1.
Figure 9 The results of Prognostic value of AURKB, CHEK1 and NEK2. (A) Overall Survival Kaplan-Meier curve of AURKB, (B) Disease Free Survival Kaplan-Meier curve of AURKB, (C) Overall Survival Kaplan-Meier curve of CHEK1, (D) Disease Free Survival Kaplan-Meier curve of CHEK1, (E) Overall Survival Kaplan-Meier curve of NEK2, (F) Disease Free Survival Kaplan-Meier curve of NEK2.
TTR through neutrophil degranulation pathway, our study combined network pharmacology and bioinformatics analysis in order to provide a more specific anti-HCC mechanism of *Scutellaria barbata*. Besides, we further investigated the reliability of our target prediction through computer virtual verification.

In this study, we recognized 10 critical target genes as the potential anti-HCC mechanism of *Scutellaria barbata* through bioinformatics analysis, namely TYMS, NEK2, AURKB, AURKA, POLA1, FEN1, KIF11, CHEK1, CDK1, TOP2A. These genes were all significantly over-expressed in HCC patients and altogether involved in cell cycle, DNA replication, p53 signaling pathway and Oocyte meiosis. To be more accurate, we discovered that AURKB, CHEK1 and NEK2 could be the top 3 potential target proteins of *Scutellaria barbata* in treating HCC based on binding energy and molecular interactions such as hydrogen bonds. AURKB is considered to be an abnormally expressed gene in many cancer and tumor cell lines. AURKB overexpression is an independent molecular marker predicting tumor invasiveness and poor prognosis of HCC. AURKB has several functions during mitosis, including spindle assembly, chromosome segregation, and cytokinesis. As cells enter mitosis, survivin recruits AURKB to the inner centromere. It has been reported that AURKB together with passengers inner centromere protein (INCENP) and surviving constituted chromosomal passengers, which binds to the internal centromere region in the early stage. The CHEK1 kinase is mainly responding to single-strand breaks, and overexpression of CHEK1 in germlinal vesicle oocytes could arrest oocytes at the germlinal vesicle stage. CHEK1 is responsible for controlling G1/S, S, and G2/M checkpoints. One study demonstrated the LIN01224/mir-330-5p/CHEK1 axis was involved in the pathology of HCC. Bao et al highlighted that PLK-4/ATR/CHEK1 pathway might be a potential therapeutic target for HCC. NEK2 is a serine/threonine centrosomal kinase, which is highly expressed and activated in S and G2 phases of the cell cycle. NEK2 plays a crucial role in cell cycle progression in the inter- and M-phases. Zhang et al have found that NEK2 could promote the metastasis of HCC. Numerous studies have shown that NEK2 could mediate tumor growth, metastasis and angiogenesis through mitosis. Wu et al suggested that NEK2 modulated the growth, drug resistance, metastasis and angiogenesis of hepatoma cell via pAKT/NF-kB signaling and matrix metalloproteinase activation. According to the aforementioned content, the results that AURKB, CHEK1 and NEK2 were abnormally escalated in HCC patients showed in our study were consistent with the previous publication. In the result of molecular docking and the binding energy value, we speculate that *Scutellaria barbata* exhibits anti-HCC role by inhibiting the function of these protein via protein–protein interaction.

**Conclusion**

In summary, our study suggests that *Scutellaria barbata* plays a key role in the treatment of HCC with multiple targets and pathways. AURKB, CHEK1 and NEK2 could be the potential target proteins of *Scutellaria barbata* in treating HCC. Cell cycle, DNA replication, p53 signaling pathway and Oocyte meiosis consist of the fundamental regulation cores in this mechanism.

**Abbreviations**

HCC, Hepatocellular carcinoma; TCMSP, Traditional Chinese Medicine Systems Pharmacology; STRING, search tool for recurring instances of neighbouring genes; DL, Drug-likeness; OB, Oral bioavailability; GEO, Gene Expression Omnibus; WGCNA, Weighted gene co-expression network analysis; PPI, protein–protein interaction network; DAVID, Database for Annotation, Visualization and Integrated Discovery; GO, Gene ontology; KEGG, Kyoto Encyclopedia of Genes and Genomes; HBV, Hepatitis B virus; TCM, Traditional Chinese medicine; DEGs, differentially expressed genes; CSV, comma-separated value; FC, fold change; TSV, tab-separated value; DC, degree centrality; EPC, Edge Percolated Component; MNC, Maximum Neighborhood Component; DMNC, Density of Maximum Neighborhood Component; MCC, Maximal Clique Centrality; BN, Bottleneck; CC, closeness centrality; BC, betweenness centrality; PDB, Protein Data Bank; BP, Biological process; CC, cellular component; MF, molecular function; INCENP, passengers inner centromere protein; AURKB, Serine/threonine-protein kinase Aurora-B; CDK1, Cyclin-dependent kinase 1; NEK2, Serine/threonine-protein kinase NEK2; FEN1, Flap endonuclease 1; TOP2A, DNA topoisomerase II alpha; AURKA, Serine/threonine-protein kinase Aurora-A; CHEK1, Serine/threonine-protein kinase Chk1; TYMS, thymidylate synthetase; KIF11, kinesin family member 11; POLA1, DNA polymerase alpha 1, catalytic subunit.
Data Sharing Statement
All data are available in the manuscript and they are shown in figures and tables.

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Author Contributions
All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

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