Identification of 13 Key Genes Correlated With Progression and Prognosis in Hepatocellular Carcinoma by Weighted Gene Co-expression Network Analysis

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Hepatocellular carcinoma (HCC) remains hard to diagnose early and cure due to a lack of accurate biomarkers and effective treatments. Hence, it is necessary to explore the tumorigenesis and tumor progression of HCC to discover new biomarkers for clinical treatment. We performed weighted gene co-expression network analysis (WGCNA) to explore hub genes that have high correlation with clinical information. In this study, we found 13 hub genes (GTSE1, PLK1, NCAPH, SKA3, LMNB2, SPC25, HJURP, DEPDC1B, CDCA4, UBE2C, LMNB1, PRR11, and SNRPD2) that have high correlation with histologic grade in HCC by analyzing TCGA LIHC dataset. All of these 13 hub genes could be used to effectively distinguish high histologic grade from low histologic grade of HCC through analysis of the ROC curve. The overall survival and disease-free survival information showed that high expression of these 13 hub genes led to poor prognosis. Meanwhile, these 13 hub genes had significantly different expression in HCC tumor and non-tumor tissues. We downloaded GSE6764, which contains corresponding clinical information, to validate the expression of these 13 hub genes. At the same time, we performed quantitative real-time PCR to validate the differences in the expression tendencies of these 13 hub genes between HCC tumor tissues and non-tumor tissues and high histologic grade and low histologic grade. We also explored mutation and methylation information of these 13 hub genes for further study. In summary, 13 hub genes correlated with the progression and prognosis of HCC were discovered by WGCNA in our study, and these hub genes may contribute to the tumorigenesis and tumor progression of HCC.

Keywords: quantitative real-time PCR, bioinformatics, weighted gene co-expression network analyses, hepatocellular carcinoma, biomarker

Abbreviations: AUC, area under curve; BP, biological process; CC, cellular component; DEGs, differentially expressed genes; dissTOM, dissimilarity topological overlap matrix; FC, fold change; FPKM, fragments per kilobase of transcript per million fragments mapped; GEIPA, Gene Expression Profiling Interactive Analysis; GEO, Gene Expression Omnibus; GO, Gene Ontology; GS, gene significance; HBV, Hepatitis B virus; HCC, hepatocellular carcinoma; IHC, immunohistochemical; KEGG, Kyoto Encyclopedia of Genes and Genomes; MEs, module eigengenes; MF, molecular function; MM, module membership; NCBI, National Center for Biotechnology Information; PCR, polymerase chain reaction; PPI, protein–protein interaction; ROC, receiver operating characteristic; TCGA, The Cancer Genome Atlas; TOM, topological overlap matrix; WGCNA, Weighted Gene Co-Expression Network Analysis.
INTRODUCTION

Hepatocellular carcinoma is one of the most malignant tumors and is the third leading cause of cancer death in the world (Altekruse et al., 2009; GBD 2015 Mortality and Causes of Death Collaborators, 2016; Li et al., 2017). HCC is extremely hard to diagnose at an early stage, and there are currently no effective treatments for patients with advanced-stage HCC. Surgery remains the most common treatment for HCC; nevertheless, the decade survival rate is not increasing obviously (Waghray et al., 2015; Wallace et al., 2015). HBV infection is one of the most common factors in HCC in the world (de Martel et al., 2015). Patients with higher TNM stage and histologic grade have a worse prognosis compared with those for whom they are lower (Ikeda et al., 1993). It is crucial to reach an understanding of the pathological changes from normal liver to HCC. Meanwhile, effective early detection and diagnostic methods are urgently needed for improving the prognosis and treatment of HCC patients.

The use of microarray as a high-throughput platform for gene expression analysis has been soaring in recent years (Murakami et al., 2006; Li et al., 2018). Large numbers of changed genes can be detected easily by this technology, and these can be analyzed to identify critical significant genes as biomarkers (Giulietti et al., 2018; Tang et al., 2018). There are plenty of public free databases, such as TCGA and the GEO, from which abundant information can be excavated via various bioinformatics methods. WGCNA, a method of building a multiple gene co-expression model, can be used to cluster similarly expressed genes and analyze the relationships of differential character and phenotype within modules (Langfelder and Horvath, 2008; Giulietti et al., 2016; Wang Y. et al., 2019). WGCNA can divide genes into a model or network based on pairwise correlations between genes based on their similar expression profiles, and these models can correlate to different information on HCC. Compared to conventional methods, WGCNA establishes a more meaningful correlation between gene expression and clinical information. In this study, we identified 13 hub genes that have strong relationships with histologic grade in HCC by WGCNA, using the TCGA LIHC dataset. The ROC curve showed that almost all hub genes could effectively identify high histologic grade and low histologic grade. Meanwhile, the overall survival and disease-free survival of almost hub genes have significance in 10 years. Almost all of these 13 hub genes had higher expression in HCC tumor tissues compared with non-tumor tissues. After downloading GSE6764, we validated that these hub genes had the same expression tendencies as shown by the results of TCGA. Quantitative real-time PCR was also used to validate the differential expression of these hub genes between HCC tumor tissues and non-tumor tissues and high histologic grade and low histologic grade.

MATERIALS AND METHODS

Study Approval
None of the patients had been subjected to any neoadjuvant therapy before surgery. Informed consent was obtained from each patient. The protocols used in this study were approved by the Protection of Human Subjects Committee of Zhongnan Hospital. All research was performed in accordance with relevant guidelines and regulations.

Data Collection and Processing
The brief study design is shown in Figure 1. The RNA-sequencing (RNA-seq) data and corresponding clinical information for HCC were downloaded from TCGA LIHC dataset, which included 371 tumor samples and 50 adjacent tumor samples. The clinical information was comprised of event, TNM stage, histologic grade, new event, diagnosis age, and HBV infection. Due to some clinical information being missing, we used not available (NA) to replace the missing values for WGCNA. After normalization and filtering with the “edgeR” package (Wilson and Miller, 2005; Robinson et al., 2010), p-value < 0.05 and |logFC (FC)| > 1 were used for screening DEGs. The validation dataset GSE6764, which contained 10 normal liver tissues, 8 very early HCC tissues, 10 early HCC tissues, 7 advanced HCC tissues, and 10 very advanced HCC tissues, was downloaded from the GEO.

GO Enrichment and KEGG Pathway Analysis
The GO database, which is comprised of MF, BP, and CC, is used for observing gene annotation (Harris et al., 2004). The KEGG is a database resource for understanding high-level functions and utilities of the biological system (Kanehisa et al., 2016). DAVID1 is an online website that can provide a comprehensive set of functional annotation tools through which investigators can understand the biological meaning behind a large list of genes (Huang da et al., 2009). In this study, DAVID 6.7 was used to analyze the GO enrichment and KEGG pathways of all DEGs. The p-value < 0.05 was considered statistically significant.

Weighted Gene Co-expression Network Construction
We selected the top 25% variance genes, which included 4938 genes, to construct a co-expression network with the “WGCNA” package in R. Elimination of outliers is the first step in this analysis. Next, we select an appropriate soft-thresholding power that would ensure scale-free topology, R² = 0.9. The adjacency was then transformed into a TOM by using TOM similarity and the corresponding dissimilarity (dissTOM). Finally, at least 30 modules (|R| ≥ 0.25) were aggregated into different MEs by the TOM similarities and Pearson distances. The corresponding dissimilarity was then transformed into a TOM by using TOM similarity and the corresponding dissimilarity (dissTOM). Finally, at least 30 modules (|R| ≥ 0.25) were aggregated into different MEs by the TOM similarities and Pearson distances. The adjacency was then transformed into a TOM by using TOM similarity and the corresponding dissimilarity (dissTOM). The network based on pairwise correlations between genes based on their similar expression profiles, and these models can correlate to different information on HCC. Compared to conventional methods, WGCNA establishes a more meaningful correlation between gene expression and clinical information. In this study, we identified 13 hub genes that have strong relationships with histologic grade in HCC by WGCNA, using the TCGA LIHC dataset. The ROC curve showed that almost all hub genes could effectively identify high histologic grade and low histologic grade. Meanwhile, the overall survival and disease-free survival of almost hub genes have significance in 10 years. Almost all of these 13 hub genes had higher expression in HCC tumor tissues compared with non-tumor tissues. After downloading GSE6764, we validated that these hub genes had the same expression tendencies as shown by the results of TCGA. Quantitative real-time PCR was also used to validate the differential expression of these hub genes between HCC tumor tissues and non-tumor tissues and high histologic grade and low histologic grade.

Identification of Clinical Significant Modules
The correlations between clinical information and MEs were calculated by WGCNA in R. We selected the most significant and correlated modules with HCC. In this study, the genes in blue and turquoise MEs with the histologic grade of HCC

1https://david-d.ncifcrf.gov/
were our next research objects. Meanwhile, two concepts of GS and MM were proposed in our study. Genes that have high significance for clinical information and MM were confirmed by these two parameters.

**Hub Gene Identification and Verification**

The connectivity of genes was measured by the absolute value of the Pearson’s correlation. Genes with high within-module connectivity were considered to be hub genes of the modules ($\text{cor.geneModuleMembership} > 0.85$). Hub genes in given modules tended to have a strong correlation with certain clinical information, which was measured by the absolute value of the Pearson’s correlation ($\text{cor.geneTraitSignificance} > 0.2$). Next, we intersected hub genes in MEs with all DEGs via the edgeR package. At the same time, we eliminated some hub genes that have been researched multiple times in HCC. We definitely considered these hub genes significant by this filter. To validate the functions of these hub genes, the GSE6764 microarray was downloaded from the GEO database in NCBI.

**Survival Analysis and ROC Analysis of Hub Genes**

To identify whether these hub genes have deeper value, we searched the GEPIA website\(^2\) for overall survival and disease-free survival analysis of all of the hub genes (Tang et al., 2017). The log-rank test was used to show statistical significance. Meanwhile, the AUC, calculated by R, was used to differentiate different histologic grades (high histologic and low histologic) of HCC.

**Mutation and Methylation Information of Hub Genes**

We searched the cBioPortal website\(^3\) to identify whether these hub genes have epigenetics information about mutation and methylation (Cerami et al., 2012). The corresponding figures were downloaded from that website.

**Immunohistochemical Information of Hub Genes**

To explore the expression of these hub genes in HCC and normal tissues, we searched the Human Protein Atlas public database\(^4\) for IHC information.

**Tissue Collection and Quantitative Real-Time PCR**

A total of 16 pairs of HCC and corresponding adjacent tissues were obtained from 16 HCC patients after surgery in Zhongnan Hospital, Wuhan University. The histologic grades of all HCC tissues were identified by the pathology department in our

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\(^{2}\)http://gepia.cancer-pku.cn/

\(^{3}\)http://www.cbioportal.org/

\(^{4}\)https://www.proteinatlas.org/
hospital. All samples were stored with RNA protective reagent at 
−80°C. Total RNA was extracted from HCC tissues by TRIzol 
reagent (Invitrogen, Carlsbad, CA, United States). Extracted 
total RNA was quantified using a NanoDrop spectrophotometer 
(Termo Scientific Inc.) at 260 and 280 nm. RNA was used 
for reverse transcription when \( A_{260}/A_{280} = 2.0 \). The reverse 
transcription was processed with the PrimeScript RT reagent Kit 
with gDNAEraser (Takara, Tokyo, Japan). Quantitative real-time 
PCR was performed with the SYBR Green PCR kit (Toyobo, 
Osaka, Japan) using the CFX Connect Real-Time PCR Detection 
System (Bio-Rad, United States) after setting up the appropriate 
protocol. The GAPDH was used as an internal control. The 
\( 2^{-\Delta\Delta Ct} \) method was used to analyze the results of quantitative 
real-time PCR. All the primers were designed by the website 
PrimerBank\(^5\) and are listed in Supplementary Table 1.

RESULTS

Identification of Differentially Expressed Genes

After data processing, a total number of 2356 significant DEGs, 
including 789 down-regulated and 1567 up-regulated genes, were 
identified between HCC tissue and adjacent tumor tissue by the 
“edgeR” package in R. Volcano and heatmap plots were used to 
show this result (Figures 2A,B).

GO and KEGG Pathway Enrichment Analysis of DEGs

To obtain a deeper understanding of the annotation and function 
of all of the DEGs, we put all of these DEGs into DAVID to 
analyze significant GO and KEGG pathways. The up-regulated 
DEGs were remarkably enriched in cell cycle, M phase, M 
phase of mitotic cell cycle, mitotic cell cycle, and other BP 
(Figure 3A). The KEGG enrichment analysis identified cell 
cycle, DNA replication, pathways in cancer, and other pathways 
(Figure 3B). Meanwhile, the down-regulated DEGs were mainly 
enriched in response to wounding, acute inflammatory response, 
oxidation–reduction, and other BP (Figure 3C). The KEGG 
enrichment results of down-regulated DEGs are complement 
and coagulation cascades, fatty acid metabolism, PPAR signaling 
pathway, and other pathways (Figure 3D). It seems that the 
disorder of these pathways possibly reflected the intricate 
pathological mechanism of HCC.

Weight Gene Co-expression Network 
Construction and Key Module 
Identification

After downloading the FPKM value expression matrix of all 
HCC samples, we selected the top 25% variance genes, including 
4938 for WGCNA. To eliminate outliers, we chose 130 for the 
cut tree height for the samples (Figure 4A). The number of 
HCC samples under the red line was 352 after clustering. The 
sample dendrogram and trait heatmap of 352 samples in our 
study are shown in Figure 4B. We chose the power of \( \beta = 5 \) 
(scale-free \( R^2 = 0.92 \), slope = −1.52) as the soft-thresholding

\(^5\)https://pga.mgh.harvard.edu/primerbank/
to construct a scale-free network (Figures 4C,D). By gathering similarly expressed genes, a total of nine modules were identified, namely black module [947], blue module [748], brown module [735], gray module [160], magenta module [35], pink module [160], red module [460], turquoise module [1023], and yellow module [670] (Figure 5A). The MEs in the blue and turquoise modules showed a higher correlation with histologic grade of HCC ($R^2 = 0.33$, $p = 3e^{-10}$; $R^2 = 0.34$, $p = 3e^{-11}$) (Figure 5B). We therefore chose these two modules for further research. The module membership and gene significance for LIHC stage in turquoise and blue module were showed in Figures 6A,B. The relationship between connectivity in modules and gene significance was showed in Figure 6C.

### Identification of Hub Genes

By setting up cor.geneModuleMembership > 0.85 and cor.geneTraitSignificance > 0.2, there are 9 hub genes in the blue module and 46 hub genes in the turquoise module (Supplementary Table 2). The PPI networks of all genes in the two modules were, respectively, constructed by Cytoscape. We used the cytoHubba app to calculate the degree of each node. The nine hub genes were included in the top 20 hub genes in the blue module according to cytoHubba (Figure 6D). The 46 hub genes were also included in the top 50 hub genes in the turquoise module according to cytoHubba (Figure 6E). Next, we took the 44 intersecting genes from the two module hub genes and all DEGs in HCC (Figure 7A). Considering that some of the hub genes have already been reported on and researched in HCC, such as NCAPG, TTK, TOP2A, CDC20, CDK1, and so on (Wong et al., 2009; Li et al., 2014; Liu et al., 2015; Wu et al., 2018; Zhang et al., 2018), we finally chose 13 hub genes (GTSE1, PLK1, NCAPH, SKA3, LMNB2, SPC25, HJURP, DEPDC1B, CDCA4, UBE2C, LMNB1, PRR11, and SNRPD2) on which little research had been done regarding HCC to continue our deeper exploration.

### Validation of Hub Genes

To validate the expression of these 13 hub genes in tumor tissue and adjacent tissue, we downloaded GSE6764, in which there are 11 hub genes that have the same tendency and statistical significance compared with the TCGA database (Figures 7C,D). All of these hub genes belonged to the set of adverse factors in HCC (Figure 7B). To validate different histologic grade expression, four groups (very early HCC, early HCC, advanced HCC, and very advanced HCC) in GSE6764 were considered to approximate histologic grades I–IV, and most of the hub genes had significance in a one-way ANOVA test (Figures 8A,B).

### Survival and ROC Analysis

The overall survival and disease-free survival of these 13 hub genes were analyzed by using the GEIPA online database. Nearly all of them had a poor prognosis when highly expressed on the
FIGURE 4 | Sample clustering and determination of thresholding power in weighted gene co-expression network analysis (WGCNA). (A) Sample clustering to detect outliers. (B) Sample dendrogram and trait heatmap. (C) Analysis of the scale-free fit index for various soft-thresholding powers ($\beta$). (D) Analysis of the mean connectivity for various soft-thresholding powers. (E) Checking the scale-free topology when $\beta = 5$. 
basis of log-rank test analysis (Supplementary Figures 1–5). The ROC curve was used to measure early HCC (Grades I and II) and advanced HCC (Grades III and IV). The AUC of almost all hub genes exceed 0.65, which meant that these hub genes could effectively differentiate early HCC and advanced HCC (Supplementary Figures 6–8. The AUCs of hub genes are listed in Figure 9A).

Mutation and Methylation Information
To find deeper information on these hub genes, we searched the cBioPortal website to identify whether these hub genes have epigenetics information about mutation and methylation available. Unfortunately, all hub genes had no obvious mutation events (Figure 9B). Meanwhile, PRR11 had more amplifications compared with the other hub genes, which could explain its high expression in HCC. The mRNA expressions (Z-scores) of more than half of the 13 hub genes had negative correlations with methylation (HM450) (Supplementary Figures 9, 10). Among these results, the correlation of DEPDC1B even reached −0.52, which revealed that methylation of the promoter region of DEPDC1B probably regulated expression of the corresponding mRNA.

Immunohistochemical Information
To obtain precise results, we chose the same antibody for each hub gene between HCC and normal tissue. The results
FIGURE 6 | Module membership and corresponding gene significance and protein–protein interaction (PPI) network construction of two key modules. (A) Module membership in the turquoise module. (B) Module membership in the blue module. (C) Relationship between the connectivity of all modules and gene significance. (D) Top 20 hub genes in the blue module according to cytoHubba analysis. (E) Top 50 hub genes in the turquoise module according to cytoHubba analysis.
FIGURE 7 | Confirmation and validation of 13 hub genes differentially expressed in tumor tissues and non-tumor tissues in the TCGA database and GSE6764.

(A) The 44 common genes between all DEGs according to the edgeR package and hub genes according to WGCNA. (B) Forest plot of 13 hub genes. (C) The boxplot of 13 hub genes differentially expressed in tumor and non-tumor tissues of HCC in the TCGA database. P-value was calculated by t-test. *p < 0.05, ***p < 0.001. (D) Boxplot of 13 hub genes differentially expressed in tumor and non-tumor tissues of HCC in GSE6764. P-value was calculated by t-test. *p < 0.05, ***p < 0.001; NS, no significance.
FIGURE 8 | Hub gene expression in different histologic grades in The Cancer Genome Atlas (TCGA) database and GSE6764. (A) Boxplot of 13 hub genes in histologic grades I–IV in the TCGA database. P-value was calculated by one-way ANOVA. ***p < 0.001. (B) Boxplot of 13 hub genes in histologic grades I–IV in GSE6764. P-value was calculated by one-way ANOVA. *p < 0.05, **p < 0.01, ***p < 0.001; NS, no significance.

The results of quantitative real-time PCR showed that 12 hub genes had significantly different expressions in HCC tissues and adjacent tissues on the basis of paired t-test. However, PRR11 showed no significant differential expression between HCC tissues and adjacent tissues (Supplementary Figures 13, 14). Meanwhile, the expression of 13 hub genes in high histologic grade was higher than that in low histologic grade, except SKA3, which demonstrated that the results of WGCNA made sense (Supplementary Figure 14).
FIGURE 9 | The basal information and mutation information for 13 hub genes. (A) Information on 13 hub genes. ROC, receiver operating characteristic; AUC, area under curve; FC, fold change; FDR, false discovery rate. (B) Mutation information on 13 hub genes in the cBioPortal website.

DISCUSSION

Hepatocellular carcinoma seriously impacts human health, and it is likely to recur after surgery combined with other therapy (Worns and Galle, 2010). The 10-year survival rate of patients will observably decrease when metastasis is found, even if optimal treatments are used. It is crucial to explore the etiological and biological mechanisms involved in the occurrence and development of HCC. Better biomarkers for predictive and prognostic molecules
for HCC are urgently required. The RNA-seq-expression and clinical information of HCC from the TCGA database are correlated by WGCNA.

In this study, we constructed protein–protein networks to show the top 20 and 50 genes in the blue and turquoise modules by calculating degree in Cytoscape. Through identification of DEGs, WGCNA, and filtering out some of the genes, we finally screened out 13 hub genes on which few studies had been conducted in HCC. These hub genes were not only expressed differently in HCC and adjacent tissue but also had significance with overall survival and disease-free survival. Meanwhile, almost all of the hub genes could effectively distinguish early grade HCC from advanced HCC, which implied that these hub genes play an important role in the progress of HCC. Genome-wide DNA methylation is well known, but its emergence and development mechanisms remain mysterious (Meissner et al., 2008; Lister et al., 2009). By searching cBioPortal, we found that the correlation between DEPDC1B expression and corresponding methylation reached −0.52 in HCC. DEPDC1B, as a protein that accumulates in G2, coordinates de-adhesion events and cell-cycle progression at mitosis (Marchesi et al., 2014). A previous study showed that DEPDC1B was a guanine nucleotide exchange factor and induced both cell invasion and migration in oral cell lines (Su et al., 2014). However, the function of DEPDC1B in HCC has not been reported yet.

When the expressions of the subset of the hub genes were validated in GSE6764, we found that GTSE1 and PRR11 showed no obvious differences between HCC and normal tissues. At the same time, GTSE1, SKA3, LMNB2, and PRR11 had no obvious tendency between different histologic grades. GTSE1 and PRR11 have been reported in various types of tumor. High expression of GTSE1 can promote breast cancer growth and metastasis. Moreover, GTSE1 can cause multidrug resistance in breast cancer cells (Lin et al., 2019). In Tongue Squamous Cell Carcinoma (TSCC) cells, PRR11 can promote proliferation and invasion by regulating p21, p27, CDK2, and Cyclin A to facilitate S phase progression (Wang C. et al., 2019). Recently, a study showed that down-regulation of SKA3 could inhibit HCC proliferation and invasion in vivo and in vitro. On the basis of gene enrichment analysis (GSEA), SKA3 was enriched in the cell cycle and P53 signaling pathways and these results were verified (Hou et al., 2019). Moreover, it has been reported that SKA3 can facilitate PI3K/AKT signaling pathway to promote cell cycle and progression in cervical cancer (Hu et al., 2018).

Compared with the HCC samples in the TCGA database, GSE6764 had few samples in each group, which may lead to this incomplete result. Besides, we could not find paired patient samples of HCC and normal tissue for immunohistochemistry due to the limitation of the Human Protein Atlas database. By quantitative real-time PCR in 16 paired tissues, we found that all hub genes had significance between tumor tissues and non-tumor tissues except PRR11. Meanwhile, all hub genes except SKA3 had significance for distinguishing high histologic grade from low histologic grade in HCC.

It is necessary for us to enlarge our sample set to validate the expression levels of these hub genes. However, we could not proceed further with the research because of various limitations. We hope that other researchers will be able to uncover the specific functions and mechanisms of these 13 hub genes in HCC, especially DEPDC1B.

**CONCLUSION**

In conclusion, this study aimed to identify hub genes involved in HCC by WGCNA of data from the TCGA database and concluded that 13 functional genes (GTSE1, PLK1, NCAPH, SKA3, LMNB2, SPC25, HJURP, DEPDC1B, CDCA4, UBE2C, LMNB1, PRR11, and SNRPD2) are highly differentially expressed in tumor and non-tumor tissues. Further bioinformatic analysis showed that these hub genes can act as biomarkers of HCC progression and prognosis. At the same time, experimental assay was performed to validate these hub genes. However, further molecular experiments on these hub genes in HCC, in vivo and in vitro, need to be carried out.

**DATA AVAILABILITY STATEMENT**

The RNA-seq data and corresponding clinical information of HCC were downloaded from TCGA. The validation dataset accession GSE6764 microarray was downloaded from GEO database in NCBI. All remaining raw data supporting the conclusions of this manuscript will be made available by the authors, without undue reservation, to any qualified researcher.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by the Protection of Human Subjects Committee of Zhongnan Hospital. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

YG, QL, and ZL designed this study. YG, JL, DG, and KY analyzed the data and wrote the manuscript. YG, BC, PL, and YX conducted the quantitative real-time PCR. All authors reviewed the 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/fgene.2020.00153/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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