Identification of hub genes in thyroid carcinoma to predict prognosis by integrated bioinformatics analysis

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ABSTRACT

The aim of this study was to identify hub genes closely related to the pathogenesis and prognosis of thyroid carcinoma (THCA) by integrated bioinformatics analysis. In this study, through differential gene expression analysis, 1916 and 665 differentially expressed genes (DEGs) were obtained from The Cancer Genome Atlas (TCGA) and Gene Expression Omnibus (GEO) database, and 7 and 11 co-expressed modules were identified from the TCGA-THCA and GSE153659 datasets, respectively, by weighted gene co-expression network analysis. We identified 162 overlapping genes between the DEGs and co-expression module genes as candidate hub genes. Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses of the 162 overlapping DEGs identified significant functions and pathways of THCA, such as thyroid hormone generation and metabolic process. A protein–protein interaction (PPI) analysis detected the top 10 hub genes (QSOX1, WFS1, EVA1A, FSTL3, CHRDL1, FABP4, PRDM16, PPARGC1A, PPARG, COL23A1). Finally, survival analysis, clinical correlation analysis, and protein abundance validation confirmed that 3 of the 10 hub genes were associated with survival prognosis of patients with THCA, and 8 of them were associated with the clinical stages of THCA. In summary, we identified hub genes and key modules that were closely related to THCA, and validated these genes by survival analysis, clinical correlation analysis, and Human Protein Atlas image analysis. Our results provide important information that will help to elucidate the pathogenesis of THCA and identify novel candidate prognostic biomarkers and potential therapeutic targets.
Introduction

Thyroid carcinoma (THCA) is the most common cancer involving the endocrine system and the most prevalent malignant tumor of the head and neck, accounting for about 1%–2% of systemic malignancies. The incidence rate is predominant in females and has been increasing year by year [1,2]. The etiology of THCA remains unclear and is currently thought to be the synergistic result of multiple carcinogenic factors, such as nutrition, dietary habits, obesity, and radiation exposure [3,4]. Treatments for THCA include surgery, thyroid hormone, and radiation therapy. The comprehensive and systematic therapies mean that most patients with differentiated THCAs have a relatively good prognosis, but patients with distant metastasis or undifferentiated THCAs often have a less favorable prognosis [5,6]. Therefore, it is urgent to identify the hub genes involved in THCA to help develop new therapeutic targets.

The development of next-generation high-throughput sequencing technologies and genomics has generated large amounts of cancer gene expression data and clinical information that have been stored in many public databases [7,8]. This has laid the foundation for further elucidating the molecular mechanisms of cancer and the biological functions of hub genes by bioinformatics methods and experimental validation. Weighted gene co-expression network analysis (WGCNA) is commonly used to identify co-expression gene modules that are highly correlated with clinical phenotypes [9].

In this study, we aimed to identify potential prognostic biomarkers for distinguishing THCA at different clinical stages. Previous studies [10–14] have used transcriptome and microarray data for gene differential expression analysis, Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway functional enrichment analysis, protein–protein interaction (PPI) network analysis, and/or survival analysis. In this study, we added WGCNA, and validated the identified hub genes by clinical correlation analysis and Human Protein Atlas (HPA) image analysis.

Materials and methods

Data acquisition from the TCGA and GEO databases

THCA gene expression data and survival information were obtained from The Cancer Genome Atlas (TCGA) (https://portal.gdc.cancer.gov) and Gene Expression Omnibus (GEO) (https://www.ncbi.nlm.nih.gov/geo) databases. We downloaded TCGA RNA-Seq raw counts and corresponding clinical data from the TCGA database using the R package TCGAbiolinks [15]. A total of 567 samples were collected, including 509 THCA and 58 normal thyroid tissues. We downloaded the GEO high-throughput sequencing GSE153659 dataset using the R package GEOquery [16]. A total of 24 THCA and 7 adjacent normal thyroid tissues were collected.

Data preprocessing and differential expression analysis

Different intergroup comparisons were conducted using the Rstudio software (version 1.3.959, https://rstudio.com/) and Bioconductor R packages (http://www.bioconductor.org/). First, the gene expression matrices of the TCGA-THCA and GSE153659 datasets were normalized separately, and formatted in the RStudio input file format. Then, volcano plots and heatmaps of the differentially expressed genes (DEGs) were drawn using the limma [17] and pheatmap [18] packages in R. Genes with adjusted P-value <0.05 and |logFC (fold change)| >1 were considered to be statistically differentially expressed.

Weighted gene co-expression network analysis (WGCNA)

WGCNA can identify modules of highly correlated genes and hub genes by constructing scale-free co-expression networks [19]. We used the R package WGCNA [20] in RStudio to perform WGCNA for the TCGA-THCA and GSE153659 datasets separately. The filtered and trimmed raw expression data and clinical information were input, then all the genes...
and samples were clustered by Pearson correlation-based hierarchical clustering. To construct a scale-free gene co-expression network, we set soft threshold power values using the pickSoftThreshold function, and calculated the scale-free fit index and average connectivity of each power. Then, the adjacency matrices were converted into a topological overlap matrix, which was used for gene hierarchical clustering and dynamic shear module recognition, and similar modules were clustered and merged. Finally, the gene list of each module was obtained and gene-module tree and module-trait relationship diagrams were drawn.

Intersection of differential expression analysis and WGCNA

In the module-trait relationship diagrams of the TCGA-THCA and GSE153659 datasets, the top two modules most significantly related to THCA were found. Overlapping genes between DEGs and top two most relevant co-expression module genes (CEMGs) were considered as candidate hub genes of THCA, which were visualized as a Venn diagram using the R package VennDiagram [21].

Functional and pathway enrichment analyses

To explore the biological functions and key pathways closely associated with the candidate hub genes of THCA [22], we used the R packages org.hs.eg.db, clusterProfiler, enrichplot, and ggplot2 [23,24] to perform GO [25] and KEGG [26] enrichment analyses. P-value <0.05 was considered statistically significant.

Construction of PPI network and identification of hub genes

PPI networks can provide interaction information that can help to identify hub genes and key modules involved in the development of THCA. We used the STRING database (http://www.string-db.org/) to construct a PPI network of candidate genes [27]. Genes with a minimum required interaction score ≥0.5 were chosen to build a full network model, which was then visualized in Cytoscape software (http://www.cytoscape.org).

We used the Maximal Clique Centrality (MCC) algorithm of the cytoHubba plugin to select the top 10 genes in the network and then visualized them in Cytoscape [28].

Survival analysis and clinical correlation analysis of hub genes

To confirm the reliability of the identified network hub genes and verify their prognostic value, we performed an overall survival (OS) analysis using the Kaplan Meier method [29] in the R packages survival and survminer [30] based on the clinical data from the TCGA-THCA dataset. We used the online survival analysis tool GEPIA2 (http://geopia2.cancer-pku.cn/) to evaluate the disease-free survival (DFS) [31] of the patients with THCA. Then the R package ggpubr [32] was used to analysis the clinical correlation between gene expression and tumor stage. Only the patients for which complete clinical information was available were included in our analysis. The hub genes were classified into two groups based on their expression levels and the median values. Genes with P < 0.05 were regarded as prognosis-related hub genes.

Validation of protein abundance levels of prognosis-related hub genes

The Human Protein Atlas (HPA) (http://www.proteinatlas.org/) is an antibody-based database that provides a large amount of transcriptome and proteomic data on various disease and normal tissues [33]. We used the HPA to validate protein abundance levels of the prognosis-related hub genes by downloading immunohistochemical (IHC)-stained images and analyzing them.

Results

We identified hub genes and key modules that were highly related to the prognosis of THCA by integrated bioinformatics analysis, including differential expression analysis, WGCNA, GO and KEGG enrichment analyses, PPI network analysis, survival analysis, and clinical correlation analysis, and validated protein abundance by examining
related HPA images. The goal was to further elucidate the pathogenesis of THCA and find novel prognostic biomarkers and potential therapeutic targets.

Identification of DEGs

The TCGA-THCA dataset contained the gene expression matrix and clinical data for 567 samples, including 509 THCA and 58 normal thyroid tissues. The GSE153659 dataset contained the expression profiles from 24 TCGA and 7 paired normal tissues. After data preprocessing and gene differential expression analysis, we identified 1916 DEGs from the TCGA-THCA dataset; 910 were down-regulated and 1006 were up-regulated (Figure 1a). We also identified 665 DEGs from the GSE153659 dataset; 471 were down-regulated and 194 were up-regulated (Figure 1b). The top 50 DEGs from each dataset were visualized in heatmaps (Figure 1c,d).

WGCNA of the whole transcriptome expression matrix

Genes with similar expression patterns tended to have similar biological functions. Therefore, we constructed a weighted gene co-expression network according to the RNA-Seq count data of 14,161 genes in the TCGA-THCA dataset and 58,437 genes in the GSE153659 dataset. We analyzed the soft threshold powers of the network topology with threshold weights from 1 to 20 and determined the scale independence and mean connectivity of the WGCNA. We selected $\beta = 2$ and 6 as the optimal soft-thresholding parameters (Figure 2(g,h)). The hierarchical clustering dendrograms of module eigengenes are shown in Figure 2(a,b). We identified 7 modules from the TCGA-THCA dataset (Figure 2(c)) and 11 modules from the GSE153659 dataset (Figure 2(d)), and plotted module-trait diagrams to assess the relationships between gene modules and clinical characteristics. The two modules with the highest correlation with clinical phenotypes were MEred and MEturquoise in TCGA-THCA and

![Figure 1](image-url)

**Figure 1.** Identification of differentially expressed genes (DEGs) in the TCGA and GSE153659 datasets of thyroid carcinoma (THCA). The cutoff criteria were $|\log\text{FC}| \geq 1.0$ and adjusted $P < 0.05$. (a, b) Volcano plots of DEGs in the TCGA-THCA (a) and GSE153659 (b) datasets. (c, d) Heatmaps of the top 50 DEGs in the TCGA-THCA (c) and GSE153659 (d) datasets. Red, up-regulated DEGs; green, down-regulated DEGs.
MEred and Mepink in GSE153659 as shown in Figure 2(e,f).

Intersection of DEGs and CEMGs

The differential expression analysis and WGCNA identified 1916 DEGs in the TCGA-THCA dataset and 665 DEGs in the GSE153659 dataset, 5459 CEMGs in the MEred and MEturquoise modules of TCGA-THCA, and 1602 CEMGs in the MEred and MEpink modules of GSE153659. A total of 162 overlapping genes were detected between the DEGs and top two most relevant CEMGs. These genes were considered as candidate hub genes of THCA and were used for the subsequent analyses (Figure 3).

Enrichment analyses of the 162 overlapping genes

We performed GO functional and KEGG pathway enrichment analyses of the 162 overlapping genes using the R packages listed in the Methods section. The enriched GO terms (Figure 4(a)) under the biological process category were mainly thyroid hormone metabolic process, fat cell differentiation, regulation of transmembrane receptor protein serine/threonine kinase signaling pathway, and endoplasmic reticulum stress-induced intrinsic apoptotic signaling pathway. Under the cellular component category, the enriched terms were primarily related to apical plasma membrane, endoplasmic reticulum lumen, and collagen-containing extracellular matrix. Under the molecular function category, the enriched
terms were mainly related to oxidoreductase activity, extracellular matrix structural constituent, GTPase regulator activity, and actin filament binding. The KEGG pathway analysis (Figure 4(b)) found that the tyrosine metabolism, thyroid hormone synthesis, and protein digestion and absorption pathways were significantly enriched.

**Identifying hub genes in the PPI network**

We built the PPI network of the 162 overlapping candidate hub genes using the STRING database (Figure 5(a)). The MCC algorithm of the cytoHubba plug-in in Cytoscape was used to identify hub genes from the PPI network (Figure 5(b)). The top 10 network hub genes encoded QSOX1 (quiescin sulphhydryl oxidase 1), WFS1 (wolframin ER transmembrane glycoprotein), EVA1A (Eva-1 homolog A, regulator of programmed cell death), FSTL3 (follistatin like 3), CHRD1L1 (chordin like 1), FABP4 (fatty acid binding protein 4), PRDM16 (PR/SET domain 16), PPARGC1A (PPARG coactivator 1 alpha), PPARG (peroxisome proliferator activated receptor gamma), and COL23A1 (collagen type XXIII alpha 1 chain).

**Identification of prognosis-related hub genes and validation of protein abundance levels**

To detect the prognosis-related hub genes among the top 10 network hub genes (QSOX1, WFS1, EVA1A, FSTL3, CHRD1L1, FABP4, PRDM16, PPARGC1A, PPARG, COL23A1), we performed OS and DFS analyses using the R packages survival and survminer [30] and the GEPIA2 tool, respectively. We verified the expression levels of the genes in THCA tissues at different stages. The OS analysis showed that low expression of EVA1A was significantly correlation with unfavorable prognosis in patients with THCA (P < 0.05) (Figure 6(a)). The DFS analysis showed that low expression of CHRD1L1 and FABP4 was markedly associated with poor prognosis (P < 0.05) (Figure 6(b)). Our evaluation of the hub genes using the HPA images indicated that the abundance levels of the EVA1A and CHRD1L1 proteins were lower in tumor tissues than they were normal tissues.
Furthermore, the clinical correlation analysis indicated that CHRDL1, COL23A1, EVA1A, FABP4, FSTL3, PPARGC1A, PRDM16, and QSOX1 were strongly associated with the clinical stages of THCA (Figure 8).

**Discussion**

THCA is a malignant tumor that originates from thyroid follicular epithelium or parafollicular epithelial cells, and it is also the most common head and neck cancer [34]. Currently, THCA is mainly targeted using a comprehensive treatment strategy of surgery combined with thyroid hormone therapy and radiotherapy. However, because of the lack of accurate molecular targets, the prognosis is still poor, especially for patients with advanced clinical stage THCA [35]. Therefore, there is an urgent need to find new prognostic biomarkers and therapeutic targets for THCA. In this study, we used integrated bioinformatics methods such as WGCNA and identified 162 overlapping candidate hub genes in the TCGA-THCA and GSE153659 datasets. The biological functions of these genes were enriched mainly in thyroid hormone generation and metabolic process, regulation of serine/threonine kinase signaling pathway, apical plasma membrane,
endoplasmic reticulum lumen and extracellular matrix, GTPase regulator activity, SH3 domain binding, and actin filament binding. The enriched pathways were mainly thyroid hormone synthesis, protein digestion and absorption, tyrosine, and sulfur and seleno-compound metabolism. The top 10 hub genes (QSOX1, WFS1, EVA1A, FSTL3, CHRDL1, FABP4, PRDM16, PPARGC1A, PPARG, COL23A1) were identified by analyzing the constructed PPI network. Among them, low expression of EVA1A, CHRDL1, and FABP4 significantly affected the survival prognosis of patients with THCA, and the expression levels of CHRDL1, COL23A1, EVA1A, FABP4, FSTL3, PPARGC1A, PRDM16, and QSOX1 were highly correlated with the clinical stages of THCA. The identified hub genes were confirmed by survival and clinical correlation analysis and by examining related HPA images.

EVA1A (Eva-1 homolog A, regulator of programmed cell death) is a novel lysosomal and endoplasmic reticulum-associated protein involved in autophagy and cell division [36]. Several studies have found that high levels of EVA1A inhibit the growth of tumor cells by activating autophagy and inducing apoptosis [36]. Furthermore, EVA1A is a target of microRNA-125b, which plays a role in the response of liver
cancer to chemotherapy by decreasing EVA1A-mediated autophagy [37]. In our study, the downregulated expression of EVA1A in THCA tissues was closely related to poor survival prognosis. CHRDL1 (chordin like 1) is a secreted glycoprotein with a repetitive cysteine-rich domain that binds to the bone morphogenetic protein (BMP) family of ligands [38]. It has been reported that CHRDL1 expression was dramatically downregulated in gastric cancer tissues and was linked to poor survival [39]. Cyr-Depauw C, et al [40]. found that CHRDL1 acts as a negative regulator of breast cancer phenotypes by blocking the ability of BMP ligands to induce increased migration and invasion of breast cancer cells. FABP4 (fatty acid binding protein 4) is an intracellular lipid chaperone that is involved not only in lipid transport but also in inflammation and insulin resistance [41]. Differential expression of FABP4 has been reported in obesity-associated breast cancer, colorectal cancer, hepatocellular carcinoma, and FABP4 is closely correlated with tumor invasion and metastasis [42–44]. Together, these results show that WGCNA and other integrated bioinformatics analyses can detect new disease-causing genes and provide rich resources for future experiments.

A recent study found that the KEGG pathways Mineral absorption, Thyroid hormone synthesis, and Protein processing in the endoplasmic reticulum pathways and the GO molecular function
terms protein tyrosine kinase activity, enzyme binding, protein binding, and integrin binding were enriched in follicular thyroid carcinoma [45]. Interestingly, emerging evidence has indicated possible molecular links between THCA and other diseases, such as Parkinson’s disease, osteoporosis, chronic kidney disease, chronic heart failure, and Type 1 and Type 2 diabetes [46]. The Thyroid hormone synthesis, Adherens junction, and Thyroid cancer pathways were found to be common between THCA and chronic kidney disease [47]. A study that used a network-based approach [48] identified seven significant pathways, namely, Thyroid hormone synthesis, Rap1 signaling, Pathways in cancer, PI3K-Akt signaling, Proteoglycans in cancer, Bladder cancer, and Cytokine-cytokine receptor interaction, that were highly correlated with 79 hub genes of THCA that were common to other diseases. They included cancer (24 genes), neurological (20 genes), cardiovascular (14 genes), metabolic (13 genes), endocrine (13 genes), and hematological (12 genes) [48]. The results of our functional and pathway enrichment analysis are consistent with those of many other previously reported studies.

The integrated bioinformatics analyses allowed us to identify hub genes and key modules closely associated with the initiation and progression of THCA. The results were validated by the survival analysis, clinical correlation analysis, and protein abundance data. Our results will help to elucidate the pathogenesis of THCA and provide novel candidate biomarkers and potential therapeutic targets for future studies. A limitation of this study is the lack of focus on the different subtypes of THCA. Clearly, the hub genes associated with the pathogenesis and outcome of THCA still need to be tested experimentally.

**Conclusion**

In this study, we identified three genes (EVA1A, CHRDL1, FABP4) that were associated with survival prognosis of patients with THCA, and eight (CHRDL1, COL23A1, EVA1A, FABP4, FSTL3, PPARGC1A, PRDM16, QSOX1) that were
associated with clinical stages. These genes are candidate novel prognostic biomarkers and potential therapeutic targets.

Research highlights
- We obtained top 10 hub genes (QSOX1, WFS1, EVA1A, FSTL3, CHRDL1, FABP4, PRDM16, PPARGC1A, PPARG, COL23A1) in thyroid carcinoma
- 3 of the 10 hub genes were associated with survival prognosis of thyroid carcinoma
- 8 of the 10 hub genes were associated with clinical stages of thyroid carcinoma
- The hub genes may be novel prognostic biomarkers and potential therapeutic targets

Acknowledgements
The authors would like to thank the TCGA and GEO database for providing valuable gene expression matrix and clinical survival data of THCA. We would also like to thank Margaret Biswas, PhD, from Liwen Bianji (Edanz) (http://www.liwenbianji.cn/ac), for English language editing.

Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.
Authors’ contributions

HRZ and YWP designed the study. YWP wrote the manuscript. JW collected raw data. YWP, L JW and SH performed statistical and bioinformatics analyses. TW and HRZ supervised the study. All authors read and approved the final version of the manuscript.

Ethics approval and consent to participate

TCGA and GEO belong to public databases. The patients involved in the database have obtained ethics approval. Users can download relevant data for free for research and publish relevant articles. Our study was based on open-source data, so there were no ethical issues and other conflicts of interest.

Disclosure of potential conflicts of interest

The authors declare that they have no competing interests.

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