The expression of macropinocytosis-related genes in ovarian cancer and their relationships with prognosis

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Research Article

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Abstract

**Background:** The mechanism of macropinocytosis has been reported in receptor sorting used by motile cells. Besides, the role of macropinocytosis was previously recognized in cancer progression. We evaluated the prognostic value of macropinocytosis gene expression in ovarian cancer (OC).

**Method:** Ten candidate genes were selected in the intersection between 134 macropinocytosis-related genes from Genecards database and 2925 OC prognostic genes using the Cancer Genome Atlas (TCGA) database. Heat map showed ten candidate genes expression. Gene ontology (GO) and Kyoto encyclopedia of genes and genomes (KEGG) analysis was conducted on the ten candidate genes. Protein–protein interactions were constructed using the STRING database. Hub genes were identified based on PPI networks. The key hub genes were selected both in differential expression analysis and Kaplan-Meier survival analysis. Then we identified transcription factor-gene interaction. The relationships between clinical characteristics and the key hub genes expression were performed with T test. Clinicopathologic factors correlated with overall survival (OS) conducting univariate, multivariate and LASSO Cox regression analyses. Human Protein Atlas (HPA) databases were utilized to verify the results. Furthermore, Gene Set Enrichment Analysis (GSEA) identified the potential key pathways that dominate macropinocytosis in OC.

**Result:** Elevated EZR, HSPG2 and SLC9A1 expression was significantly associated with OC poor survival and clinical features. Transcription factor-gene interaction and GSEA analysis reported many key regulators and signaling pathways that were enriched in OC with varying degrees of macropinocytosis-related genes expression.

**Conclusions:** The three macropinocytosis-related genes might be utilized as new candidate prognostic biomarkers for OC.

Introduction

Ovarian cancer (OC) is the most lethal gynaecological malignancies and possesses a high capacity for metastasis [1]. OC metastasizes either by direct extension from the ovarian/fallopian tumor to neighboring organs (bladder/colon) or when cancer cells detach from the primary tumor [2]. An in-depth understanding of the cellular and molecular mechanisms of OC metastasis are crucial for developing effective preventative measures.

Macropinocytosis is a unique pinocytosis process by which cancer cells internalize extracellular proteins or necrotic cell debris and deliver them to lysosomes for further degradation. During tumor development, macropinocytosis provide not only a survival possibility under nutritional deficiencies for cancer cells, but also the potential for tumors to limitlessly grow in harsh tumor microenvironments [3]. Macropinocytosis is driven by actin cytoskeleton remodeling. Several actin polymerization regulators, such as small GTPases, p21-activated kinase 1 (Pak1), and PI3K, have been related to the formation of plasma membrane protrusions and macropinocytic activity [4]. Recently, one study showed that the
internalization of eATP by macropinocytosis in human lung cancer cells could drive epithelial-mesenchymal transition (EMT) to induce metastasis [5]. Another study also reported that cadherin-6B was removed from premigratory neural crest cells through internalization induced by macropinocytosis, then promoted EMT and migration [6]. These studies revealed the significance of macropinocytosis in the tumor metastasis. Up to now, some studies showed that macropinocytosis had the advantage for the drug delivery in OC therapy [7, 8]. However, a clear potential of macropinocytosis-related genes with prognostic meaning and clinical diagnosis in OC has not yet been more comprehensively studied.

Accordingly, the establishment of new biomarkers related to macropinocytosis is crucial for the early detection and prognosis of OC. The Cancer Genome Atlas (TCGA) database was used to analyze the clinical information of OC and mRNA expression difference in normal and OC tissues. The key hub genes were selected from 134 candidate macropinocytosis-related genes for subsequent correlation analyses of clinical factors, univariate, multivariate and LASSO Cox regression analyses. To identify the contribution of the key hub genes to the signaling pathway related to macropinocysis, Gene Set Enrichment Analysis (GSEA) was worked for functional enrichment analysis. Furthermore, Human Protein Atlas (HPA) was applied to render verication support to our outcomes from TCGA. This study was conducive to the enormous potential of macropinocysis-related genes as a new prognostic marker of OC in the future.

Materials and methods

Data extraction and arrangement

The workflow of this study is shown in (Fig. 1). To explore the key genes and pathways which affect OC metastasis, macropinocytosis-related genes were selected from the GeneCards database (protein coding and relevance score > 1). The whole transcriptome sequencing data set of raw read counts and fragments per kilobase per million (FPKM) data and corresponding clinical data of all ovarian OC patients were downloaded from the TCGA database (n = 381) for collecting overall survival data, and then ranked according to $P$-value ($P < 0.05$). To obtain macropinocytosis genes related to OC prognosis, we intersected macropinocytosis-related genes and OC prognostic genes. The candidate genes were found in the intersection. Then we combined TCGA with normal tissues from the Genotype-Tissue Expression (GTEx) data from Xena database (tumor tissues = 427, normal tissues = 88) for further analyses. The candidate genes expression heat map was produced using the ComplexHeatmap R package (2.13.1).

Gene ontology (GO) and Kyoto encyclopedia of genes and genomes (KEGG) pathway enrichment analyses

To comprehensively investigate the biological meaning behind the candidate genes, clusterProfiler 4.4.4 was used to obtain the set of functional annotation. GO is an extensively used tool for annotating genes with potential functions, such as molecular function (MF), biological pathways (BP) and cellular
components (CC). KEGG enrichment analysis is a practical resource for analytical study of gene functions and associated high-level genome functional information. $P<0.05$ was regarded as the threshold value. Results were visualized with ggplot2 3.3.6.

**Protein-protein Interactions (PPIs) network and hub gene analysis of the candidate genes**

All candidate genes were subject to PPI network analysis utilizing STRING database with the median confidence 0.4 [9]. Cytoscape (v3.10.0) was used to visualize the PPI network. Both the color and the size of the nodes indicates the score calculated using degree topological analysis methods.

**Extraction of hub genes**

Cytoscape software was applied to analyse the hub genes, which are important nodes with many interaction partners. We utilized the cytoHubba plugin in Cytoscape to identify the hub genes [10]. Determination of the top 20 genes depends on the BottleNeck algorithm. The ranks of hub genes are represented by a gradient from red to yellow. Finally, the hub genes were ranked for the shortest accessible paths between hub genes, making them easier to observe.

**Expression level and survival analyses of the Hub genes in OC**

The gene expression level and OC patients survival was analyzed in R (4.2.1), and data was visualised using ggplot2[3.3.6]. Briefly, RNA sequencing expression and survival data from TCGA and GTEx project was used to perform the key hub genes expression differences analyses in normal and OC tissues by Wilcoxon rank sum test. Only the samples with both survival status and survival time information could be used for overall survival (OS), progress free interval (PFI) and disease specific survival (DSS). Kaplan-Meier survival curves were evaluated using the long-rank test. Only genes with high expression level and low survival time in tumor tissues ($P$-value < 0.05) could be considered as the key hub genes with significant differences for subsequent analyses.

**Identification of transcription factors associated with the key hub genes**

Transcription factors (TFs) control chromatin and transcription by identifying specific DNA sequences [11]. In addition to controlling genome expression, they provide essential information for molecular understanding. We explored the regulatory effects of the TFs on the key hub genes through the TFs retrieved from the ChipBase database [12], then obtained a set of common TFs by taking the intersection of these TFs. Finally, the Cytoscape software was used to visualize the common TFs of the key hub genes.

**Correlation analysis of clinical factors and univariate, multivariate and LASSO Cox regression analyses from TCGA database**
The clinical data that removed missing information was conducted to visualize the key hub genes expression differences in clinical characteristics respectively. T test was used between two groups. Univariate and multivariate Cox regression analyses were implemented to identify independent prognosis risk factors. Only factors with entire information in age, clinical stage, histologic grade, venous invasion, lymphatic invasion, and gene expression was assessed for prognosis risk factors. All comparisons were considered statistically significant with a \( P \)-value < 0.05. Forest map was used to visualize the univariate Cox regression analyses with the ggplot2 3.3.6. We next used LASSO Cox regression to visualize the risk score and grouping of the prognostic model with ggplot2 3.3.6.

**Identification of relevant signaling pathways of the key hub genes by Gene Set Enrichment Analysis (GSEA)**

GSEA was performed using the R package clusterProfiler [4.4.4] for functional enrichment analysis from the TCGA database to investigate possible biological functions of the key hub genes, including GO Enrichment, DO Enrichment, KEGG Enrichment and Reactome Enrichment. The following parameters were set to run the enrichment test. The name of each gene was selected as the expression dataset, and “c2.cp.all.v2022.1.Hs.symbols.gmt” was chosen as the gene sets database. The max size excluded larger sets and the min size excluded smaller sets were set in default values of 500 and 15, respectively. Signaling pathways with a false discovery rate (FDR) q-value < 0.25 was used to select the enriched signaling pathways when adjusted \( P \)-value < 0.05.

**The tissue-level of the key hub genes expression in Human protein atlas**

The location of the key hub genes in the body was further determined by searching for its expression in the HPA (https://www.proteinatlas.org/search/HAMP).

**Results**

**Data extraction and arrangement**

134 genes that relate to macropinocytosis were selected from the GeneCards database. The intersection between macropinocytosis-related genes and OC prognostic genes (2925 genes, \( P < 0.05 \)) from TCGA were regarded as candidate genes. In total, we identified 10 candidate genes as follows: ARHGEF26, SNX33, PAK1, SLC9A1, LRP1, HSPG2, UBAP2, CXCR4, SLC38A5 and IL32 (Fig. 2A). Then we combined TCGA with normal tissues from GTEx database to constructed a gene expression heat map to visualize the expression levels of the 10 candidate genes (Fig. 2B).

**GO and KEGG pathway enrichment analysis**

Function annotation analysis of the 10 candidate genes was performed. We present the top three items for each category in a bar chart and a bubble chart. In GO enrichment analysis, the 10 candidate genes...
are significantly enriched in the regulation of blood–brain barrier, vascular transport and regulation of actin cytoskeleton organization in BP subsets. Cell–substrate junction, focal adhesion and cell leading edge are in CC subsets. Lipoprotein particle receptor binding, amyloid–beta binding and aromatic amino acid transmembrane transporter activity are in MF subsets. To explore the biological functions and enriched pathways of the 10 candidate genes, KEGG enrichment analysis was performed. The top three pathways identified by KEGG pathway analysis are as follows: salmonella infection, regulation of actin cytoskeleton and proteoglycans in cancer (Fig. 2C and 2D).

**Identification of the Hub Genes and selection of the key hub genes**

The ten candidate genes were uploaded to STRING to explore the network of PPI. PPI network map was visualized through Cytoscape software to identify common gene interaction (Fig. 2E). Subsequently, we identified the most significant hub genes through BottleNeck method using the CytoHubba plug-in in Cytoscape, including CCL5, DAG1, CDC42, CXCR4, FGF2, EZR, CD4, NCK1, NID1, HSPG2, CCL2, CXCL1, CCL19, CXCL12, SLC9A1, CCL21, CXCL13, PAK1, CALR and LRP1 (Fig. 2F). Then we performed differential expression analyses and survival analyses to select the key hub genes with high expression in tumor tissues and low survival from the twenty hub genes to perform further analyses. We finally analysed three key hub genes including EZR, HSPG2 and SLC9A1 (Fig. 3).

**The gene expression, survival analysis of the key hub genes and the relations to clinicopathological parameters from TCGA**

The gene expressions of EZR (\(P<0.05\), Fig. 3A), HSPG2 (\(P<0.05\), Fig. 3E) and SLC9A1 (\(P<0.05\), Fig. 3I) in OC tumor tissues were higher than those in normal tissues, and the differences were all statistically significant (\(P<0.05\), Fig. 3). Kaplan-Meier survival analysis showed that high expression of EZR (OS, \(P = 0.028\), Fig. 3B) (DSS, \(P = 0.016\), Fig. 3D), HSPG2 (OS, \(P = 0.018\), Fig. 3F) (DSS, \(P = 0.013\), Fig. 3H) and SLC9A1 (OS, \(P = 0.009\), Fig. 3J) (DSS, \(P = 0.014\), Fig. 3L) all possessed worse prognosis with a poor OS and a poor DSS. Only EZR overexpression possessed a poor PFI (\(P = 0.006\), Fig. 3C). SLC9A1 expression was notably associated with histologic grade (\(P<0.05\), Fig. 4M). EZR and SLC9A1 expression was significantly related to age (\(P<0.05\), Fig. 4A and 4K).

**Establishment of TF regulatory networks**

In order to elucidate the key molecules that bridge OC and macropinocytosis, we conducted a framework to indicate gene regulatory networks involving TFs from the key hub genes. By analyzing the interaction network of TFs, we found 128 potential common TFs that regulate the three key hub genes (Fig. 5). Our results indicated the potential connections between the key hub genes with common TFs.

**Hazard factors affecting patient’s survival**
The univariate Cox regression analysis uncovered that the up-regulation of EZR, SLC9A1 and HSPG2 expression was related to poor OS in OC (Table 1 and Fig. 6A). The other clinical variable associated with bad survival was age (hazard ratio [HR]: 1.352; 95% confidence interval [CI]: 1.045–1.749; \( P = 0.022 \)) (Fig. 6A and Table 1). In multivariate Cox analysis, age and high SLC9A1 expression maintained an independent risk factor for OS among OC (Table 1). Group with high-risk score tend to have higher levels of SLC9A1, EZR and HSPG2 expression and less survival compared with the low-risk group (Fig. 6B).

Table 1

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Total(N)</th>
<th>Univariate analysis</th>
<th>( P ) value</th>
<th>Multivariate analysis</th>
<th>( P ) value</th>
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<td></td>
<td></td>
<td>HR(95% CI)</td>
<td></td>
<td>HR(95% CI)</td>
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<tr>
<td>Age</td>
<td>379</td>
<td>1.352 (1.045–1.749)</td>
<td>0.022*</td>
<td>1.390 (1.066–1.812)</td>
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<td>Histologic grade</td>
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<td>Lymphatic invasion</td>
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<td>EZR</td>
<td>379</td>
<td>1.329 (1.026–1.720)</td>
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<td>1.146 (0.872–1.505)</td>
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<td>HSPG2</td>
<td>379</td>
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<td>1.153 (0.856–1.553)</td>
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<td>SLC9A1</td>
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<td>1.416 (1.092–1.835)</td>
<td>0.009**</td>
<td>1.363 (1.010–1.839)</td>
<td>0.043*</td>
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</table>

* \( P<0.05 \); ** \( P<0.01 \); EZR, Ezrin; HSPG2, Heparan Sulfate Proteoglycan 2; SLC9A2, Solute Carrier Family 9

Member A1.

Results of GSEA

We further performed GSEA on the three key genes to discover signaling pathways that were distinguishingly activated in macropinocytosis. As shown in Fig. 6, different degrees of SLC9A1, EZR and HSPG2 expression were related to phagocytosis, focal adhesion, endocytosis and glycolysis gluconeogenesis pathways in cancer. It showed that the three key hub genes may be closely related to macropinocytosis in OC.

The tissue-level of the key hub genes expression in HPA

SLC9A1, HSPG2 and EZR was mostly found in the cytoplasm or membranous of OC cells, as shown through searching their expressions in the HPA database (Fig. 6).

Discussion
Macropinocytosis is a form of fluid-phase endocytosis in which cells use ruffles of the plasma membrane to take up medium into primary endocytic vesicles [13]. It is a nonspecific internalization process and has a vital role in the uptake of extracellular substances and antigen presentation. During tumor development, it helps cancer cells survive in nutrient-deficient environment to resistant to anticancer drugs, and promote invasion and metastasis [14]. This process depends on the activation of the RAS gene, growth factor receptors (GFRs), and other signal pathways [15, 16].

Epithelial-mesenchymal transition (EMT) is the key mechanisms that facilitates the cancer metastasis. It is often defined by the loss of the epithelial marker E-cadherin and the gain of the expression of the mesenchymal marker vimentin [17]. Extracellular ATP (eATP) can be internalized into the lung cancer cells to trigger EMT via purinoceptor signals activated by P2X7. It can also activate production of matrix metallopeptidase (MMPs) and promote lung cancer cell shedding, EMT, migration, and invasion. During this process, Snail and Slug (and probably others), which is known as EMT TFs, was enhanced [5]. During the development of the neural crest cells migration, Cad6B is removed from premigratory neural crest cells through cell surface internalization events that include clathrin-mediated endocytosis and macropinocytosis. Both of these processes are dependent upon the function of dynamin, and inhibition of Cad6B internalization abrogates neural crest cell EMT and migration [6]. Accordingly, tumor cells utilize macropinocytosis to support energy, and may also internalize the adhesion receptor directly to reduce intercellular adhesion and promote migration capacity.

In the present study, we performed differential expression analyses of the ten candidate genes involved in macropinocytosis and prognostic of OC, then selected three key hub genes through PPI and a series of differential expression analyses and survival analyses between tumor and normal OC tissues. They were all overexpressed and significantly correlated with poor survival in OCs. Previous studies have found that EZR [18], HSPG2 [19] and SLC9A1 [20] were potential targets for the treatment of OC, which are consistent with our analysis results. Subsequently, patients with elevated SLC9A1 expression had an advanced histologic grade, however, the EZR and HSPG2 expression levels were not notably associated with the clinicopathological factors. We constructed a prognostic risk model using the three key hub genes through univariate Cox, multivariate Cox and Lasso Cox regression analysis. Increased EZR, HSPG2 and SLC9A1 expression was related to poor OS by univariate Cox regression analysis. In multivariate Cox analysis, increased SLC9A1 expression maintained an independent risk factor for OS. Group with high-risk score tended to have higher levels of SLC9A1, EZR and HSPG2 expression and less survival compared with the low-risk group. Furthermore, the protein expression of the three key hub genes in OC patients and normal ovarian samples was testied by the HPA database.

Next, we conducted a framework to elucidate gene regulatory networks involving the common TFs from the three key hub genes. It has been reported that pancreatic ductal adenocarcinoma (PDAC) cells could escape autophagy inhibition via NRF2-induced macropinocytosis to upregulate and utilize nutrient [21]. In hepatocellular carcinoma cells, hypoxia-inducible factor (HIF)-1 activated the transcription of a membrane ruffling protein, EH domain-containing protein 2 (EHD2), to initiate macropinocytosis to harness extracellular protein as a nutrient to survive [22]. However, its function in OC remains to be
addressed. These key hub genes may have the potential to be the promising biomarkers and new targets in therapeutic approaches for OC.

The functions of EZR, HSPG2 and SLC9A1 were investigated in OC from TCGA data by GSEA. Some signaling pathways such as Ecm Receptor Interaction, Integrin Cell Surface Interactions, Integrin1 Pathway, Focal Adhesion, Endocytosis, Fc Gamma R Mediated Phagocytosis, Fcgamma Receptor Fcgr Dependent Phagocytosis, as well as Glycolysis Gluconeogenesis that differentially enriched in high or low expression of the key hub genes were more comprehensively reported in the present study. These results suggested that EZR, HSPG2 and SLC9A1 may serve as prognostic markers and potential therapeutic targets in OC, which were not summarized in other bioinformatic analyses research of OC.

During the process of macropinocytosis formation, excessive activation of Ras or Rac GTPases can drive actin cytoskeleton remodelling causing plasma membrane ruffling and bulk engulfment of extracellular material [23]. Actin can drive protrusion at the plasma membrane, and plays a key role in sculpting membranes that drive endocytic uptake, trafficking, and recycling. Recycling of receptors such as integrins and receptor tyrosine kinases (RTKs) regulates adhesion to the ECM as well as actin organisation for migration towards nutrient gradients [24]. This mechanism is essential in recycling of proteins involved in invasion and metastasis including various integrins [25]. In non-small-cell lung cancer, upregulation of proteins involved in recycling such as clathrin light chain b (CLCb) and dynamin-1 (Dyn1) can enhance EGFR recycling to plasma membrane through clathrin-mediated endocytosis, leading to increased invasion and metastasis in vivo [26]. These processes depend on actin dynamics and are regulated by signals from membrane receptors and contact with the matrix. Accordingly, targeting macropinocytosis via the actin cytoskeleton or its regulators could be a potential therapeutic target.

During our analyses, EZR, HSPG2 and SLC9A1 was mainly enriched on integrin, adhesion and endocytosis pathway in OC, but the underlying relationship in the diagnosis and prognosis of OC remains unclear. Considering the combination of targeting macropinocytosis and OC metastasis, may provide new possibilities for OC diagnosis, prognosis, and treatment. They would be new potential targets of therapeutic drugs for OC.

**Conclusions**

Our research found the affinitive relationship between three macropinocytosis-related genes expression (EZR, HSPG2 and SLC9A1) and the prognosis of OC. Moreover, the Ecm Receptor Interaction, Integrin Cell Surface Interactions, Integrin1 Pathway, Focal Adhesion, Endocytosis, Fc Gamma R Mediated Phagocytosis, Fcgamma Receptor Fcgr Dependent Phagocytosis, and Glycolysis Gluconeogenesis may be the key pathways controlled by macropinocytosis-related genes in OC. Therefore, macropinocytosis-related genes may become important markers and new targets for early diagnosis, precise treatment, and prognostic assessment of OC.
Declarations

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Competing Interests

The authors declare no conflict of interest.

Author contributions

Formal analysis, data curation, writing - original draft, validation and data analysis was performed by Ying shao. Visualization was performed by Shuai Huang. Yingshao and Zhaochun He conceived the idea, revised the manuscript and approved the final submission. All authors read and approved the final manuscript.

Date Availability

Any data and R script in this study can be obtained from the corresponding author upon reasonable request. In this study, TCGA and GTEx expression data set is available on the TCGA and GTEx data portal.

Ethics approval

Not applicable.

Consent to participate

Not applicable.

Consent to publish

Not applicable.

Acknowledgments

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References


Figures
Figure 1

The flowchart of our research process
Figure 2

The data extraction, expression levels and function annotation analysis of the macropinocytosis-related candidate genes in OC. Identification of the Hub Genes and selection of the key hub genes. (A). The intersection between macropinocytosis-related genes and OC prognostic genes from the TCGA was regarded as the candidate genes; (B). The heat map of the ten screened macropinocytosis-related candidate genes in normal (88 tissues) and OC tissues (427 OC samples) from the TCGA and GTEx database; (C-D) GO and KEGG pathway enrichment of the ten candidate genes; (E) The PPI network was used to identify common gene interaction shared between the ten candidate genes. Cytoscape was used to visualize the PPI network; (F) The identification of the most significant hub genes through BottleNeck method using the CytoHubba plug-in in Cytoscape, including CCL5, DAG1, CDC42, CXCR4, FGF2, EZR, CD4, NCK1, NID1, HSPG2, CCL2, CXCL1, CCL19, CXCL12, SLC9A1, CCL21, CXCL13, PAK1, CALR and LRP1. OC, ovarian cancer; TCGA, the Cancer Genome Atlas; GO, Gene ontology; KEGG, Kyoto encyclopedia of genes and genomes; BP, biological process; CC, cellular components; MF, molecular function; CCL5, C-C Motif Chemokine Ligand 5; DAG1, Dystroglycan 1; CDC42, Cell Division Cycle 42; CXCR4, C-X-C Motif Chemokine Receptor 4; FGF2, Fibroblast Growth Factor 2; EZR, Ezrin; CD4, CD4 Molecule; NCK1, NCK Adaptor Protein 1; NID1, Nidogen 1; HSPG2, Heparan Sulfate Proteoglycan 2;CCL2, C-C Motif Chemokine Ligand 2; CXCL1, C-X-C Motif Chemokine Ligand 1; CCL19, C-C Motif Chemokine Ligand 19; CXCL12, C-X-C Motif Chemokine Ligand 12; SLC9A1, Solute Carrier Family 9 Member A1; CCL21, C-C Motif Chemokine Ligand 21; CXCL13, C-X-C Motif Chemokine Ligand 13; PAK1, P21 (RAC1) Activated Kinase 1; CALR, Calreticulin; LRP1, LDL Receptor Related Protein 1.
The boxplots and Kaplan-Meier survival curves with mRNA expression of the key hub genes in OC from TCGA database. Different expression of the key hub genes between OC tumor and normal tissues, including EZR (A), HSPG2 (E) and SLC9A1 (F). The higher expression of EZR (OS, P=0.028, Figure 3B) (DSS, P=0.016, Figure 3D), HSPG2 (OS, P=0.018, Figure 3F) (DSS, P=0.013, Figure 3H) and SLC9A1 (OS, P=0.009, Figure 3J) (DSS, P=0.014, Figure 3L) all possessed worse prognosis with a poor OS and a poor DSS. Only EZR overexpression possessed a poor PFI (P=0.006, Figure 3C). OC, ovarian cancer; OS, overall survival rate; PFI, progress free interval; DSS, disease specific survival; EZR, Ezrin; HSPG2, Heparan Sulfate Proteoglycan 2; SLC9A2, Solute Carrier Family 9 Member A1. TCGA, the Cancer Genome Atlas;
Figure 4

The correlation between the key hub genes (EZR, HSPG2 and SLC9A1) expression levels and various clinicopathological features in OC patients using Wilcoxon signed-rank test from the TCGA database. EZR (A) and SLC9A1 (K) expression level was significantly related to age ($P < 0.05$). Higher SLC9A1 expression level was notably associated with high histologic grade (M, $P < 0.05$). OC, ovarian cancer; EZR, Ezrin; HSPG2, Heparan Sulfate Proteoglycan 2; SLC9A2, Solute Carrier Family 9 Member A1; TCGA, the Cancer Genome Atlas.
**Figure 5**

Identification of the common TFs of the key hub genes (EZR, HSPG2 and SLC9A1). (A). Intersection of all TFs of the key hub genes; (B). The common TFs related to the key hub genes were used as nodes in an interconnected regulatory network. Light purple oval nodes represent the key hub genes, and TFs interact with genes as yellow oval nodes. TF, transcription factors; EZR, Ezrin; HSPG2, Heparan Sulfate Proteoglycan 2; SLC9A2, Solute Carrier Family 9 Member A1.
Construction of a risk prognostic model and enrichment plots in OC patients from GSEA based on the key hub genes (EZR, HSPG2 and SLC9A1). (A). Univariate Cox regression analysis was performed for the key hub genes. A value of $P < 0.05$ was considered statistically significant; (B) LASSO regression of the key hub genes; (C-E) Functional enrichment analysis of the key hub genes expression in OCs from TCGA, including EZR (C), HSPG2 (D) and SLC9A1 (E). (F-K). Expression of the key hub genes at the tissue level in the HPA database: (F-G) High level of EZR expression in OC tissue and negative expression in normal ovarian tissue; (H-I) Medium level of HSPG2 expression in OC tissue and negative expression in normal ovarian tissue; (J-K) High level of SLC9A1 expression in OC tissue and negative expression in normal ovarian tissue. OC, ovarian cancer; GSEA, Gene Set Enrichment Analysis; EZR, Ezrin; HSPG2, Heparan Sulfate Proteoglycan 2; SLC9A2, Solute Carrier Family 9 Member A1; TCGA, the Cancer Genome Atlas; HPA, Human Protein Atlas.