Identify key genes and pathways in pancreatic ductal adenocarcinoma via bioinformatics analysis

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Abstract
Background: Pancreatic cancer has many pathologic types, among which pancreatic ductal adenocarcinoma (PDAC) is the most common one. Bioinformatics has become a very common tool for the selection of potentially pathogenic genes.

Methods: Three data sets containing the gene expression profiles of PDAC were downloaded from the gene expression omnibus (GEO) database. The limma package of R language was utilized to explore the differentially expressed genes (DEGs). To analyze functions and signaling pathways, the Database Visualization and Integrated Discovery (DAVID) was used. To visualize the protein-protein interaction (PPI) of the DEGs, Cytoscape was performed under the utilization of Search Tool for the Retrieval of Interacting Genes (STRING). With the usage of the plug-in cytoHubba in cytoscape software, the hub genes were found out. To verify the expression levels of hub genes, Gene Expression Profiling Interactive Analysis (GEPIA) was performed. Last but not least, UALCAN analysis online tool was implemented to analyze the overall survival.

Results: The 376 DEGs were highly enriched in biological processes including signal transduction, apoptotic process and several pathways, mainly associated with Protein digestion and absorption and Pancreatic secretion pathway. The expression levels of nucleolar and spindle associated protein 1 (NUSAP1) and SHC binding and spindle associated 1 (SHCBP1) were discovered highly expressed in pancreatic ductal adenocarcinoma tissues. NUSAP1 and SHCBP1 had a high correlation with prognosis.

Conclusions: The findings of this bioinformatics analysis indicate that NUSAP1 and SHCBP1 may be key factors in the prognosis and treatment of pancreatic cancer.

Introduction
Pancreatic cancer is famous for its high malignancy, rapid development and poor prognosis. The five-year survival rate of pancreatic cancer is no more than 6%.\(^1\). It has many pathologic types, among which pancreatic ductal adenocarcinoma (PDAC) is the most common one and has a 5-year survival rate less than 3% .\(^2,3\).

Even to this day, no treatment other than surgical excision such as chemotherapy or radiation has
had a significant effect on pancreatic cancer. The specific mechanism of pancreatic cancer's strong ability to proliferate and invade has not been explored. Pancreatic cancer can easily invade and metastasize to surrounding tissues through blood vessels and lymph nodes. Many patients who have been diagnosed with pancreatic cancer have lost the opportunity for surgery because it is usually advanced.

Up to now, we still need more biomarkers which have high sensitivity and specificity rather than CA19-9, the only biomarker approved by the Food and Drug Administration. CA19-9 has many limitations, especially specificity and effectiveness. Differentially expressed genes play an important role in the proliferation, invasion and progression of pancreatic cancer, including p16, SMAD4, K-ras and p53. But they still couldn't fully explain how pancreatic cancer develops.

Bioinformatics has become a very common tool for the selection of potentially pathogenic genes. Many researchers have used bioinformatics to identify potential oncogenes. Despite the lack of in vivo or in vitro analysis, it is still a clever way to find potential oncogenes. For example, Look for potential oncogenes in ovarian cancer and pancreatic cancer.

Microarray analysis has been a very mature technology and has been widely used to screen differentially expressed genes in cancer. This present study is set out to investigate the DEGs between pancreatic ductal adenocarcinoma tissues and normal pancreatic tissues by determining gene expression profiles from GEO database. Besides, the functional enrichment and interaction network analyses of the DEGs were verified to help us explore the hub genes and the underlying molecular mechanisms of pancreatic ductal adenocarcinoma.

In result, 376 DEGs and 30 hub genes were identified, which may be potential biomarkers for pancreatic ductal adenocarcinoma. Among these genes, SHCBP1 and NUSAP1 were worthy of further experimental verification of their mechanism in pancreatic tumors and were likely to be a key factor in the prognosis and treatment of pancreatic cancer.

Materials And Methods

2.1 Affymetrix Microarray Data
National Center for Biotechnology Information Search database created a gene expression database (http://www.ncbi.nlm.nih.gov/geo), which preserves the high-throughput gene expression data provided by research institutions around the world.

Utilizing the keywords “pancreatic ductal adenocarcinoma”, we searched the GEO database. Three GEO datasets were downloaded, including GSE46234 contributed by Raeder H, GSE46385 contributed by Walters DM, and GSE71989 contributed by Schmittgen T. These gene expression profiles of pancreatic ductal adenocarcinoma were downloaded based on GPL570 Affymetrix Human Genome U133 Plus 2.0 Array. The GSE46234 dataset included 4 pancreatic ductal adenocarcinoma tissues and 4 normal pancreatic tissues. The GSE46385 consisted of 9 pancreatic ductal adenocarcinoma tissues and 3 normal pancreatic tissues. GSE71989 contained 13 pancreatic ductal adenocarcinoma tissues samples and 8 normal pancreatic tissues. We presented the detailed information about the datasets in Table 1.

2.2 Screening for DEGs

By using Affy package of R language, we manipulated the raw data by adjusting background, normalizing the data and log-transforming the raw data values. Using Perl language software, we merge these GEO datasets together and batch normalized the data set by utilizing the sva package in R language. These DEGs in pancreatic ductal adenocarcinoma and normal pancreatic tissues were analyzed by utilizing the limma package. Log2-foldchange(log2FC) > 2 and a corrected $p$ value < 0.05 were set as the cut-off criteria of DEGs samples.

2.3 GO and KEGG Pathway Enrichment Analyses

We utilized the DAVID(version 6.8) online database to explore the Biological functions of the DEGs obtained from the integration of microarray data. In order to verify the enrichment signaling pathways of DEGs, Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway and Gene Ontology (GO) enrichment analyses were performed with DAVID.

2.4 PPI Network Integration, Modules Analysis, and Selection of Hub Genes

We utilized the STRING (version11) online database to build a PPI network, in order to identify the
interaction between protein-protein interaction (PPI). We set the highest confidence of the argument of interactions at >0.7. To visualize and to analyze the PPI network, we used the Cytoscape (version 3.6.1) software. The plug-in cytoHubba was utilized for exploring key nodes and fragile motifs in the PPI network.

2.5 RNA sequencing expression and survival analysis and of core genes

UALCAN is an interactive web resource for analyzing cancer OMICS data (TCGA and MET500). It has high quality graphics using java script and CSS. To identify these DEGs, we applied the GEPIA website to analyze the data of RNA sequencing expression on the basis of thousands of samples from TCGA and Genotype-Tissue Expression (GTEx).

Results

3.1 DEGs in pancreatic ductal adenocarcinoma

In whole study, there were 26 pancreatic ductal adenocarcinoma tissues and 15 normal pancreatic tissues. 376 DEGs were identified, including 191 downregulated and 185 upregulated genes. We show the DEGs in Figures 1, known as the volcano plots of DEGs. We present the Heatmap in Figure 2. The gene names of them are exhibited in table 2.

3.2 GO and KEGG Analysis of DEGs.

As shown in Table 3 and Figure 3, all 376 DEGs were analyzed by DAVID and the results of GO analysis revealed for biological processes (BP), DEGs were highly enriched in signal transduction, apoptotic process, proteolysis, cell adhesion and cell-cell signaling; for molecular function (MF), DEGs were mainly enriched in identical protein binding, calcium ion binding, protein homodimerization activity, serine-type endopeptidase activity, catalytic activity and receptor binding; for GO cell component (CC), DEGs were significantly enriched in extracellular exosome, extracellular space, extracellular region, integral component of plasma membrane and cell surface.

We presented the KEGG analysis results in Table 4, which indicated that DEGs were particularly enriched in Protein digestion and absorption, Pancreatic secretion, ECM-receptor interaction and some other pathways. (P < 0.05).
3.3 Protein–protein interaction network (PPI) and modular analysis

In order to explore the biological characteristics of these DEGs, we created a PPI network with the usage of the STRING database. We presented the result in Figure 4. Meanwhile we utilized the cytoHubba plug-in to screen hub genes and the result is shown in Figure 5 and table 5.

3.4 Survival Analysis

By comparing all the literature on pancreatic cancer to date, we found two genes that had not been explored and studied in pancreatic cancer especially pancreatic ductal adenocarcinoma. In order to validate the expressions of these two hub genes, we utilized the GEPIA website to analyze the data of RNA sequencing expression on the basis of 350 samples (Figure 6.Aand Figure 6.B). Moreover, UALCAN website was used to analyze the hub genes survival in the 177 samples which is derived from the TCGA project (Figure 6.Cand Figure 6.D). The result indicated the expressions of NUSAPI and SHCBP1 were closely related to the survivals in pancreatic adenocarcinoma.

Discussion

In the past few decades, many causes and potential mechanisms for the formation and development of pancreatic adenocarcinoma have been identified by a lot of experts. However, there was no significant improvement in 5-year survival. Moreover, the identification of reliable molecular markers with high prognostic value has not been found in pancreatic ductal adenocarcinoma. Therefore, we need to explore and discover a specific biomarker to detect early pancreatic adenocarcinoma, which is conducive to the treatment and survival of patients. Last but not least, we need to discover and validate new molecular targets to develop drugs that can effectively treat pancreatic cancer. Many studies concentrate on an independent genetic event, or the result is generated from independent studies which are inconsistent with each other by microarray analysis.16-18

Some GEO samples (GSM) of pancreatic adenocarcinoma were not grouped into pancreatic ductal adenocarcinoma, the result may lose some details or may even be wrong for the mixed disease subtypes. Pancreatic adenocarcinoma pathogenesis in different patients may depend on common
changes of the expression of particular critical genes, and rather on personal particular changes of different genes. Comparing pancreatic adenocarcinoma instead of pancreatic ductal adenocarcinoma tissues with normal tissues may miss some important genes. Our study was done by comparing pancreatic ductal adenocarcinoma tissues with normal pancreatic tissues which may find some potential genes had not found in previous research. Our researcher found, in agreement with previous studies, most of the hub genes plays an important role in the development and progression of pancreatic ductal adenocarcinoma and they were reported and explored by previous research. That proves the reliability of our research in some ways.

*TOP2A* could activates EMT process which was highly associated with tumor metastasis in pancreatic cancer patients and could enhance tumor progression.\(^\text{19}\) PHGDH could down regulate the expression of *CCNB1, CCND1, MMP-2, and MMP-9* in pancreatic cancer cells and inhibited the cell invasion, migration, and proliferation.\(^\text{20}\) The down-regulated expression of *CDC20* in pancreatic cancer cells could do the same work. Meanwhile it induced cell apoptosis in pancreatic cancer cells.\(^\text{21}\) The knock-down of *NDC80* in Panc-1 cells could inhibited cell proliferation and colony formation. Moreover, it infected Panc-1 cells resulted in induction of apoptosis and cell cycle arrest at G2/M phase.\(^\text{22}\) The interference suppression of *KIF20A* resulted in an inhibition of motility, invasion and proliferation of pancreatic cancer cell lines.\(^\text{23}\) The upregulated *PBK/TOPK* in pancreatic cancer and upregulated expression levels was highly correlated with the invasive property of pancreatic cancer cells. By regulating *PBK/TOPK* pathway, the invasion ability of PDAC cells can be regulated.\(^\text{24}\) The knock-down of *IMP3* in pancreatic cancer reduced PDAC cell viability, extracellular matrix adhesion and invasion. *IMP3*-depleted cells presented lower levels of *CD44* protein and *KIF11* mRNA.\(^\text{25}\) Overexpression of *BUB1B, CDK1, CCNA2* and *CDC20* in pancreatic cancer was highly associated with disease-free survival in PDAC patients.\(^\text{26}\)

As for the genes we found, Nucleolar and Spindle Associated Protein 1 (*NUSAP1*) and SHC binding and spindle associated 1 (*SHCBP1*) were both hub genes and interact with the hub genes which we
discussed above, which suggested that *NUSAP1* and *SHCBP1* may also play a very important role in pancreatic cancer.

We found that *NUSAP1* had been reported the expression of *NUSAP1* is up-regulated in non-small cell lung cancer, which is associated with the growth and development of non-small cell lung cancer and prognosis of the patients. 27. In non-small cell lung cancer, the inhibited expression of *NUSAP1* could result in the blocking the cell cycle on G1 phase. 28. In addition, *NUSAP1* Inhibited Cell Proliferation in Invasive Breast Cancer Cells. 29. Last but not least, *NUSAP1* was associated with HCC onset, progression, and prognosis, and exhibited higher expression in HCC compared with normal livers. 30. In all, *NUSAP1* was associated with different types of tumors. Until now, there is no such study in pancreatic cancer. The GEPIA website revealed that *NUSAP1* has a clear trend of increasing gene expression levels in Pancreatic adenocarcinoma compared to normal samples. The report was on the basis of 350 samples. Meanwhile, UALCAN website revealed that *NUSAP1* had a significantly worse survival. Above all, *NUSAP1* may play an important role in pancreatic cancer.

Another gene we found in this study was *SHCBP1*. In gastric cancer, *SHCBP1* suppressed apoptotic and promoted cell cycle progression which accelerated tumor growth and invasiveness. 31. Meanwhile, *SHCBP1* accelerated cisplatin induced invasion, migration and apoptosis resistance by activating Wnt pathway in lung cancer. 32. Besides, the interference suppression of *SHCBP1* could significantly promote the apoptosis of lung cancer cells. 33. In non-small cell lung carcinoma, the release of *SHCBP1* from SHC adaptor protein 1 (*SHC1*) could result in development of non-small cell lung carcinoma malignant progression and cell stemness. 34. In gliomas, the overexpression of *SHCBP1* promotes migration and invasion by activating the *NF-KB* signaling pathway. 35. Last but not least, the overexpression of *SHCBP1* could significantly promote HCC cell proliferation, survival and colony formation in HCC cell lines and the knockdown of *SHCBP1* resulted in cell cycle delay and suppressed cell proliferation in human hepatocellular carcinoma cells. 36. The GEPIA website reported that *SHCBP1* has a clear trend of increasing gene expression levels in Pancreatic adenocarcinoma.
compared to normal samples. The report was on the basis of 350 samples. Meanwhile, UALCAN website revealed that SHCBP1 had a significantly worse survival. Above all, SHCBP1 may play an important role in pancreatic cancer.

Conclusion
All in all, our data analysis study found SHCBP1 and NUSAP1 which has not been published in pancreatic ductal adenocarcinoma. They were identified on the base of mixed and normalized three different microarray datasets. Just as other researchers have used bioinformatics to explore potential oncogenes, we demonstrate the value of these two molecules for further exploration and verification from different perspectives. Results revealed that NUSAP1 and SHCBP1 were worthy of further experimental verification of their mechanism in pancreatic tumors and are likely to be a key factor in the prognosis and treatment of pancreatic cancer especially pancreatic ductal adenocarcinoma.

Declarations

**Ethics approval and consent to participate** Not applicable. Our manuscript does not report on or involve the use of any animal or human data or tissue, this section is not applicable to our submission.

**Consent for publication:** Individual person’s data contained in our manuscript is shared by GEO which is opened to publication.

**Availability of data and material:** Cytoscape (version 3.6.1) software/ R language(version 3.6.1) is used in this manuscript

**Competing interests:** The authors declare that they have no competing interests.

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### Tables

| Contributor  | Organism       | GEO       | Platform | PDAC tissues | Normal pancreatic tissues |
|--------------|----------------|-----------|----------|-------------|---------------------------|
| Raeder H     | Homo sapiens   | GSE46234  | GPL570   | 4           | 4                         |
| Walters DM   | Homo sapiens   | GSE46385  | GPL570   | 3           | 9                         |
| Schmittgen T. | Homo sapiens   | GSE71989  | GPL570   | 8           | 13                        |

Abbreviation: GEO, Gene Expression Omnibus. PDAC, pancreatic ductal adenocarcinoma

Table II: Screening DEGs in PDAC by integrated microarray
| DEGs          | Gene names                                                                 |
|--------------|-----------------------------------------------------------------------------|
| **Upregulated** | GPX8DHFRMBOAT1TMSB10ITGA2AHRLGALS3BP7MEM154A MTD1TDX6E0GTLINC0342S100A6LOC100506119S100A11PCE1DBOSPL3I1RAPKIF20BMYOFI18GPRC5ANHSS DR16CSHOTAIR1MRACGAPIYCARNDNO1Q2ER2H1UN1BE25BIFKHL2S100PCKS2F2Z2E2H2EFNB2TOP2ANCEH1ARPC1B ABRACLUHRF1CAPGCOOR2A5FNXAF1GNA15RMR2STYK1C19orf23LY75PPARGBCAR3OLFML28TYMPRL39LGB2ISG15TIPMP1SERPH15HCBP1TMEM133SLC16A4ANX1APSC5K5KF20AVILLCRIP1COL5A1C12orf5MAP3K8TRIM59LAMB3 |
| **Downregulated** | DTLCDH3RAD51APIITGB1PMAIP1SAMD9INHBABUB1BV NNI1CENPWOAS1SEMAMSCCTAPRC1CAPO85TEAPISTYL2A MIGO2GAS2L3AHNAK2GIN51Z2INTNDC80NUSAP1FOXF2CD55CE5PD5KK1CENPKLAMC2TMCC5THC1R5SD2CC2L0SUGTPLAC8SLC6A14MLNCEACAM6PKLGAL51KIF11CD H11ROBO1RAB31P3PLA2G7TSPAN15ULFDGAP5POSTNC TSEUEB2TUF2L2HPDNUMT2PCCN1BCXL5GBP3COL5A2ANLNC15orf48LIPGXC18AKR1B10CDC2EGR3CD5T7MEM45BCOMPCXRC4CDC42EPSMP12IF44L1D1COL11A1COL3AITNFRSF11BSPIAO1C5FTA2TMPS54COL1A2CCL18IF27GCNT3NGFB3COL10A1PTGS2S100A2AGR3CXL3SFRP4CTGFPFLAUST6GALNAC1LUMGREML1CN2MMPI5PCACLDN18DUOX2C8orf4VCANRK76AGABRPTFF1LYZ5SFRP2 |

Abbreviation: DEGs, differentially expressed genes

Table III. Top 10 GO analysis of genes associated with pancreatic ductal adenocarcinoma
|_BP| GO:000716| signal transduction| 35| 9.43396| 0.01199| 18.57c| 6|
|BP| GO:000691| apoptotic process| 27| 7.27762| 5.27E-05| 0.089c| 3|
|BP| GO:000650| proteolysis| 26| 7.00808| 1.79E-05| 0.030c| 1|
|BP| GO:000715| cell adhesion| 21| 5.66037| 7.34E-04| 1.24205| 1|
|BP| GO:000726| cell-cell signaling| 19| 5.12129| 2.71E-06| 0.004c| 8|
|BP| GO:004249| response to drug| 19| 5.12129| 3.15E-05| 0.053c| 1|
|BP| GO:000695| inflammatory response| 18| 4.85175| 0.00130| 2.2012| 9|
|BP| GO:004306| negative regulation of apoptotic process| 18| 4.85175| 0.00835| 13.3021| 2|
|BP| GO:003019| extracellular matrix organization| 17| 4.58221| 1.57E-06| 8.39E-06| 5|
|BP| GO:000828| positive regulation of cell proliferation| 17| 4.58221| 0.00135| 30.74c| 2|
|CC| GO:007006| extracellular exosome| 96| 25.8760| 6.34E-09| 8.89E-24| 1|
|CC| GO:000561| extracellular space| 91| 24.5283| 6.77E-27| 8.97E-24| 2|
|CC| GO:000557| extracellular region| 88| 23.7196| 9.07E-20| 1.20E-16| 3|
|CC| GO:000588| integral component of plasma membrane| 36| 9.70350| 0.06699| 60.0868| 4|
|CC| GO:000998| cell surface| 22| 5.92991| 0.00174| 2.28161| 5|
|CC| GO:000557| proteinaceous extracellular matrix| 20| 5.39083| 9.94E-07| 0.0011| 6|
|CC| GO:004847| perinuclear region of cytoplasm| 19| 5.12129| 0.05199| 50.6976| 7|
|CC| GO:000578| endoplasmic reticulum lumen| 16| 4.31266| 4.26E-06| 0.0035| 8|
|CC| GO:003101| extracellular matrix| 16| 4.31266| 5.81E-04| 0.7672| 9|
|CC| GO:000558| collagen trimer| 11| 2.96496| 9.94E-06| 0.0013| 10|
|MF| GO:004280| identical protein binding| 27| 7.27762| 0.00316| 4.54090| 1|
|MF| GO:000550| calcium ion binding| 26| 7.00808| 0.00355| 5.0847| 2|
|MF| GO:004280| protein homodimerization activity| 24| 6.46900| 0.01591| 20.95c| 3|
|MF| GO:000425| serine-type endopeptidase activity| 19| 5.12129| 2.66E-06| 0.0038| 4|
|MF| GO:000382| catalytic activity| 13| 3.50404| 3.17E-04| 0.4634| 5|
|MF| GO:000510| receptor binding| 13| 3.50404| 0.04366| 48.03c| 6|
|MF| GO:000808| growth factor activity| 12| 3.23450| 3.32E-04| 0.4860| 7|
|MF| GO:000517| hormone activity| 11| 2.96496| 1.43E-05| 0.020c| 8|
|MF| GO:000486| serine-type endopeptidase inhibitor activity| 10| 2.69541| 1.13E-04| 0.1652| 9|
|MF| GO:000820| heparin binding| 10| 2.69541| 0.00411| 5.8632| 10|

Note: BP, biological process; CC, cell component; MF, molecular function; GO, gene ontology;

Table IV. KEGG pathway enrichment analyses of DEGs associated with pancreatic ductal adenocarcinoma
| Category                        | Description                                           | Count | %       | P Value  | Genes                                                                 |
|--------------------------------|-------------------------------------------------------|-------|---------|----------|------------------------------------------------------------------------|
| KEGG_PATHWAY                   | Protein digestion and absorption                      | 18    | 4.851752| 1.19E-11 | FXPD2, CELA3A, CELA3B, SLC16A10, COL3A1, SLC3A1, COL5A2, COL5A1, PRSS2, PRSS3, COL1A2, CPA2, CELA2B, CPA1, CPB1, COL11A1, CTRL, COL10A1 |
| KEGG_PATHWAY                   | Pancreatic secretion                                   | 16    | 4.312668| 2.90E-09 | PNLIP, FXPD2, CELA3A, PNLIPRP1, COL3A1, COL1A2, ITGA2, LAMC2, COL11A1, COL5A2, COL5A1 |
| KEGG_PATHWAY                   | ECM-receptor interaction                               | 9     | 2.425876| 9.21E-04 | LAMB2, COMP, COL3A1, COL1A2, ITGA2, LAMC2, COL11A1, COL5A2, COL5A1 |
| KEGG_PATHWAY                   | Fat digestion and absorption                           | 6     | 1.617251| 0.001997 | PNLIP, CEL, CLPS, PNLIPRP1, PNLIPRP2, LAMC2, ITGA2, COL5A1 |
| KEGG_PATHWAY                   | Glycerolipid metabolism                               | 7     | 1.886792| 0.002217 | PNLIP, CEL, PNLIPRP1, PNLIPRP2, LAMC2, ITGA2, COL5A1 |
| KEGG_PATHWAY                   | Maturity onset diabetes of the young                   | 5     | 1.347709| 0.002891 | IAPP, INS, NEUROD1, NR5A2, NKX2-2 |
| KEGG_PATHWAY                   | Amoebiasis                                            | 9     | 2.425876| 0.003269 | GNA15, LAMB3, COL3A1, COL1A2, CXCL8, LAMC2, COL11A1, COL5A2, COL5A1 |
| KEGG_PATHWAY                   | Mineral absorption                                     | 6     | 1.617251| 0.003429 | FXPD2, MT1M, MT1H, STEAPI, MT1G, MT1F |
| KEGG_PATHWAY                   | Proximal tubule bicarbonate reclamation               | 4     | 1.078167| 0.015803 | FXPD2, CA4, SLC4A4, AQP1 |
| KEGG_PATHWAY                   | Focal adhesion                                         | 11    | 2.96496 | 0.022393 | LAMB3, PAK3, COMP, COL3A1, COL1A2, ITGA2, LAMC2, EGF, COL11A1, COL5A2, COL5A1 |
| KEGG_PATHWAY                   | Legionellosis                                          | 5     | 1.347709| 0.036982 | IL18, CXCL3, PYCARD, CXCL8, BNIP3 |
| KEGG_PATHWAY                   | Cytokine-cytokine receptor interaction                | 11    | 2.96496 | 0.058396 | TNFRSF11B, IL22RA1, CXCL5, CCL20, CXCR4, IL18, CXCL3, IL1RAP, CXCL8, CCL18 |
| KEGG_PATHWAY                   | Bladder cancer                                         | 4     | 1.078167| 0.070274 | TYMP, CXCL8, EGF, MMP1 |
| KEGG_PATHWAY                   | p53 signaling pathway                                 | 5     | 1.347709| 0.071255 | CCNB1, RRM2, PMAIP1, SFN, IGFBP3 |
| KEGG_PATHWAY                   | Axon guidance                                          | 7     | 1.886792| 0.076909 | PAK3, CXCR4, ROBO1, PLXNA2, EFNB2, PLXNB3, SEMA3C |
| KEGG_PATHWAY                   | Bile secretion                                         | 5     | 1.347709| 0.077599 | FXPD2, NCEH1, AQP8, SLC4A4, AQP1 |
| KEGG_PATHWAY                   | Complement and coagulation cascades                    | 5     | 1.347709| 0.07922  | CD55, SERPINA5, C6, C5, PLAU |
| KEGG_PATHWAY                   | Platelet activation                                    | 7     | 1.886792| 0.083922 | P2RX1, COL3A1, COL1A2, ITGA2, COL11A1, COL5A2, COL5A1 |

Table V. Top 30 hub genes
| Gene | Description |
|------|-------------|
| CC1  | cyclin B1 [Homo sapiens (human)] |
| NB1  | cell division cycle 20 [Homo sapiens (human)] |
| C20  | NDC80 kinetochore complex component [Homo sapiens (human)] |
| C80  | DNA topoisomerase II alpha [Homo sapiens (human)] |
| TOP2A| kinesin family member 20A [Homo sapiens (human)] |
| PBK  | PDZ binding kinase [Homo sapiens (human)] |
| RRM2 | ribonucleotide reductase regulatory subunit M2 [Homo sapiens (human)] |
| KIF11| kinesin family member 11 [Homo sapiens (human)] |
| BUB1B| BUB1 mitotic checkpoint serine/threonine kinase B [Homo sapiens (human)] |
| TPX2 | TPX2 microtubule nucleation factor [Homo sapiens (human)] |
| PRC1 | protein regulator of cytokinesis 1 [Homo sapiens (human)] |
| NU1 | nucleolar and spindle associated protein 1 [Homo sapiens (human)] |
| DLG5 | DLG associated protein 5 [Homo sapiens (human)] |
| RA1 | Rac GTPase activating protein 1 [Homo sapiens (human)] |
| CGAP1| centrosomal protein 55 [Homo sapiens (human)] |
| DTL55| denticleless E3 ubiquitin protein ligase homolog [Homo sapiens (human)] |
| RAD51| RAD51 associated protein 1 [Homo sapiens (human)] |
| ZW10| ZW10 interacting kinetochore protein [Homo sapiens (human)] |
| SH1 | SHC binding and spindle associated 1 [Homo sapiens (human)] |
| AN1 | anillin actin binding protein [Homo sapiens (human)] |
| CKS2 | CDC28 protein kinase regulatory subunit 2 [Homo sapiens (human)] |
| URF1| ubiquitin like with PHD and ring finger domains 1 [Homo sapiens (human)] |
| PPY | pancreatic polypeptide [Homo sapiens (human)] |
| CXCL8| C-X-C motif chemokine ligand 8 [Homo sapiens (human)] |
| SST1 | somatostatin [Homo sapiens (human)] |
| ANXA1| annexin A1 [Homo sapiens (human)] |
| NMUU| neuromedin U [Homo sapiens (human)] |
| CXCR4| C-X-C motif chemokine receptor 4 [Homo sapiens (human)] |
| C5L20| C-C motif chemokine ligand 20 [Homo sapiens (human)] |
Volcano plot of gene expression profile data in pancreatic ductal adenocarcinoma tissues and the normal pancreatic tissues. The red, green, and gray points represent upregulated genes, downregulated genes, and none differentially expressed genes, respectively.
Heat map. Heat map showing top 50 up-regulated and down-regulated differentially expressed genes in pancreatic ductal adenocarcinoma tissues compared to the normal tissues. They are screened on the basis of log2FC>2 and a corrected p <0.05.
Figure 3

GO enrichment analyses of DEGs in pancreatic ductal adenocarcinoma.
Figure 4

PPI network. Common DEGs PPI network constructed by STRING online database and Module analysis. The nodes meant proteins; the edges meant the interaction of proteins; green meant down-regulated DEGs and red meant up-regulated DEGs.
Figure 5

The hub genes. The nodes meant proteins; the edges meant the interaction of proteins; green meant down-regulated DEGs and red meant up-regulated DEGs.
Figure 6

Figure 6 A. RNA sequencing expression of SHCBP1  
Figure 6 B. RNA sequencing expression of NUSAP1