Identification of potential biomarkers in Pancreatic Adenocarcinoma of micro-array gene expression data

Address

Abstract: We uncover molecular biomarkers using GSE16515 data set publicly reachable at NIH/NCBI Gene Expression Omnibus database. Using Biobase, GEOquery, gplots packages in R software 3.6. Based on gene expression analysis, we detect 278 differentially expressed genes of up regulation, whereas we find 77 down-regulated gene. The gene ontology of pathway enrichments and KEGG enrichment analyses of DEGs were studied. 120 KEGG pathways associated with pancreatic adenocarcinoma (PAAD) were identified, among which the PI3K-Akt signaling pathway was observed to be significant. Other hub genes discussed in this study, may be used as potential targets for PAAD and related diseases diagnosis and treatment. The following 21 hub genes were detected through NetworkAnalyst on the basis of protein-interaction network of DEGs by the online tool STRING: CDK1, CCNB1, CDC20, PPARG, MET, ISG15, LEF1, SFN, DMD, FN1, RUNX2, UBC, TOP2A, ECT2, WNT2, EFNA5, PAK3, PKM, ITGB4, NEK2, and ALB. Similar to previous analysis of differentially expressed genes, the expression levels of PPARG and SFN were examined in TCGA database was significantly upregulated in PAAD tumor tissues compared with normal tissues. Additionally, the module analysis of protein-protein interactions revealed that 'focal adhesion', 'ECM-receptor interaction', 'amoebiasis', and 'protein digestion and absorption' had a close association with PAAD. In addition, PI3K/AKT signaling pathway in PAAD was observed to be significant. Other hub genes discussed in the study, might be utilized as promising targets for PAAD and related diseases diagnosis and drug therapies.

Keywords: biomarker; differentially expressed genes; pancreatic adenocarcinoma; gene ontology pathway enrichment; cancer

Biographical notes:

This paper is a revised and expanded version of a paper entitled **[title]** presented at [**name, location and date** of conference].

1 Introduction

Pancreatic adenocarcinoma (PAAD) is ranked one of the most deadly tumor due to the diagnosis at a distant stage which has a 5-year overall survival rate as 3% (Siegel et al., 2018). There has been significant improvements in terms of treatments such as pancreatectomy, radiotherapy, adjuvant and neo-adjuvant chemotherapies and palliative care in the previous decades (Gillen et al., 2010; Tempero et al., 2014). However, pancreatectomy still stays the most efficient treatment, specifically for initial phase pancreatic cancer (Lambert et al., 2019). Thus, an updated knowledge of simple and basic mechanism of pancreatic cancer is necessary for more useful and curable therapies and the advancement of patient survival.

Microarray has become a significant tool in investigating pancreatic cancer genes and target therapeutic drugs (W. Zhou et al., 1998). Recent studies suggest an extensive gene expression analysis of pancreatic cancer by reviewing gene expression data sets by comparing tumor and normal cells (Alldinger et al., 2005; Jones et al., 2008; W. Zhou et al., 1998).

Moreover, relative analysis of the differentially expressed genes (DEGs) stays moderately constrained, and a dependable biomarker profile would be a need to develop new gene targets (Grützmann et al., 2005). The protein expression alterations in the advancement and growth of pancreatic adenocarcinoma cancer and related diseases require comprehensive analysis. Furthermore, the relations among the detected DEGs, specifically protein-protein interaction (PPI) networks and underlying signaling pathways should be clarified. Prospect biomarkers from such investigations can later be tested utilizing other methods for use in early diagnosis of cancers (Grønborg et al., 2006).

Pei *et al.*, 2009, recently performed microarray experiments to study the differentially expressed levels of FKBP5 gene between the pancreatic tumor and normal samples (data accessible at NCBI GEO database (Edgar et al., 2002), accession GSE16515)."

The DEGs of pancreatic tumor and normal samples were identified by comparing gene expression by fold change and t-test. Afterwards, the DEGs were screened using DAVID Gene Ontology (GO) and the analysis of pathway enrichments (Sherman & Lempicki, 2009). By studying their hub nodes globally and between tumor and normal samples constructing PPI networks, the objective of this project is to study the molecular mechanisms of pancreatic cancer and related diseases growth and to come up with candidate biomarkers for diagnosis, therapeutic targets and predictions.

Earlier studies tackling pancreatic cancer and related diseases underlying biomarkers, due to the implicit evaluation of source and progenitor populations, need to support experimental studies with numerical analysis and statistical methods in addition to previous experiments on mice (Dineen et al., 2010)

Traditional therapeutic alternatives, including chemotherapy, are not efficient enough fight pancreatic cancer, and notwithstanding advancements in the last decade, the survival rate has not increased and becoming the second cause of cancer-related deaths by 2030 (Rahib et al., 2014). Thus, constant efforts of the advancement of novel therapeutic alternatives is a need (Kamimura et al., 2018).

Much work has been done with microarray gene expression technology to reveal the central mechanism of pancreatic formation and progression and focus these methods for therapeutic approach. It still persists a request for more efficient treatments or methods that can improve curative responses to pancreatic cancer medication. In this project, we used microarray data sets of public transcriptome datasets of performed microarray experiments to detect the differentially expressed genes between the pancreatic tumor and

normal samples. Although experimental studies are needed to confirm our findings, our results will reveal potential biomarkers and bright therapeutic objectives for timely screening of pancreatic cancer. The present study also focused on the phosphoinositide-3/Akt (PI3K/Akt) signaling pathway and triggering receptor tyrosine kinases (RTKs) that plays a key task in regulating downstream responses, involving cell viability, growth cycle, proliferation, cell migration and vascularization, by phosphorylating a variety of intracellular proteins (Franke, 2008; Hemmings & Restuccia, 2012). The pathway exists in all cells of major eukaryotes and is extremely maintained (Manning & Cantley, 2007). The current and future impersonal studies of suppressants fighting PI3K/AKT pathway in cancers should be clarified further. Our goal is to characterize PI3K/AKT pathway in pancreatic cancer to the practice and put forward the new propitious to patients for targeted therapies.

2 Materials and Methods

2.1. Preprocessing of the data set

The publicly accessible gene expression data set from pancreatic tumor and normal samples were pull out from the GEO database with GSE16515 (Pei et al., 2009). Genomic information ranging from gene sequences to protein structure predictions were obtained. As described by Pei et al., 2009, the data set contains a total of expression of 54,676 genes of 36 tumor and 16 normal specimens; a total of 52 specimens.

Using the GEOquery package in Bioconductor following conventional procedures in R studio, the GSE16515 data set is studied (Durinck et al., 2009; Tarca et al., 2006). The other packages we used in R studio are as the following; Biobase, biomaRT and gplots packages (Davis & Meltzer, 2007; Durinck et al., 2005; Warnes et al., 2009). To estimate the adjusted *p value* and avoid Type I errors, we used Bejamini-Hochberg Procedure to correct multiple testing. In order to adapt the statistical tests locally, hypergeometric model was performed for both of the down-regulated and up-regulated DEGs in the functional GO and pathway enrichment analysis, and false discovery rate (FDR) were computed (Benjamini & Hochberg, 1995; Dudoit et al., 2003; Hochberg & Tamhane, 1987).

2.2. Experimental data and analysis codes

Analysis were conducted in the R statistical environment. Sample codes and analysis of GSE16515 data can be found https://github.com/mathbioGVN/GSE16515.HSC.microarray.project repository. We compared samples into two groups provided that pancreatic tumor and normal cells. The data set was normalized by computing the means of the samples of each group in R programming language. The process on separated samples which is grouped by categories was performed as computing fold-change (biological significance) difference between the means of the categories. A broadly performed statistical model is the t-distribution and its versions. A t-test compares the discrepancy of the average gene expression levels between the two samples or subgroups, given the noisiness of the data i.e. the difference in means between samples divided by the standard deviation. The genes are filtered in accordance with both fold change and p value criterion. Despite the fact that, methods to correct for multiple comparisons have been applicable for a long time such as Bonferroni correction, most of these methods are not appropriate to analyze gene expression data sets (Tarca et al., 2006). We highlight statistical significance performing t-test by taking p value cutoff 0.01 and fold cut-off > 2 to identify down and up-regulated DEGs between each category understudy.

2.3. Differentially expressed genes and clustering analysis

Using GEOquery package in Bioconductor, gene expression values were pull out for each sample and converted to base-2 logarithmic scale using R language. We used gplots package of R to create heatmaps of DEGs with heatmap.2 function. Clustering analysis of DEGs was performed to compare the expression pattern of DEGs in tumor and normal cells.

2.4. GO terms and analysis of the pathway enrichments

Expression measurements annotations for up-regulated and down regulated DEGs for each group probes mapped to gene names using Ensemble Biomart package in R. All of the DEGs were characterized by their biological processes, molecular functions, and cellular components of gene ontology (GO) enrichment of the database for Annotation and DAVID which stands for Visualization and Integrated Discovery (Huang et al., 2007; Sherman & Lempicki, 2009). All classified genes were cautiously examined and further parts like the Universal Protein resource, and physical properties Gene Ontology (GO) and annotation types were taken using DAVID and KEGG Kyoto Encyclopedia of Genes and Genomes (Eden et al., 2009; Supek et al., 2011). We then compared the results of DAVID with NetworkAnalyst enrichments performed with KEGG (G. Zhou et al., 2019).

2.5. Core genes filtered through the protein-protein interaction (PPI) network

NetworkAnalyst, publicly accessible on the web, provides analysis of PPI networks for single gene lists using STRING Interactome (Szklarczyk et al., 2015). To comprehensively decipher the regulatory mechanisms in pancreatic ad and related diseases, DEGs from pancreatic tumors and normal cells samples were analyzed to form a PPI network with previously reported GO classification and enrichment.

2.6. Expression levels and survival analysis of core genes in PAAD

Based on the information of TCGA database, Ualcan (http://ualcan.path.uab. edu/index.html) (Chandrashekar et al., 2017) was utilized to conduct survival analysis. Kaplan-Meirer survival analysis was carried out using 21 identified hub genes relying on the hub gene expression values in PAAD. In contrast to normal cells, gene expression levels presents important individual differences in tumor cells. Low expression level shows the TPM value is equal or below the upper quartile value whereas high expression level shows the TPM value is above the upper quartile value.

3 Results

3.1. Experimental data analysis

With gene expression result of the GSE16515 data set, we detect differentially expressed genes (**DEGs**) in total 355 genes from **pancreatic tumor** and **normal cells** which was demonstrated in volcano plot (Figure 1). We find the down-regulated and up-regulated DEGs of pancreatic tumor and normal cell comparison. The expression values were pull out, and a heatmap was created to show the tumor and normal cell discrepancy (Figure 2). DEGs were selected with common t test, and labelled with fold change cut-off t 2 and t 2 and t 4 down-regulated gene.

(A) The boxplot shows gene expression of each sample of the raw data without normalization.
(B) Plots displaying the gene expression discrepancy in pancreatic tumor and normal cells comparison. Black illustrates no change (NO), red illustrates down-regulated (Down), and blue illustrates up-regulated (Up) DEGs, FC, fold change.

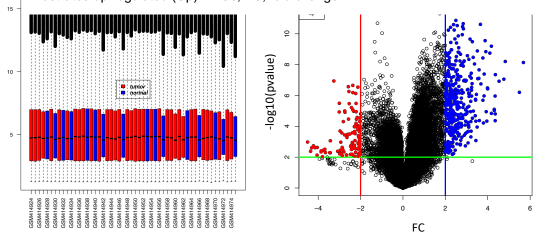
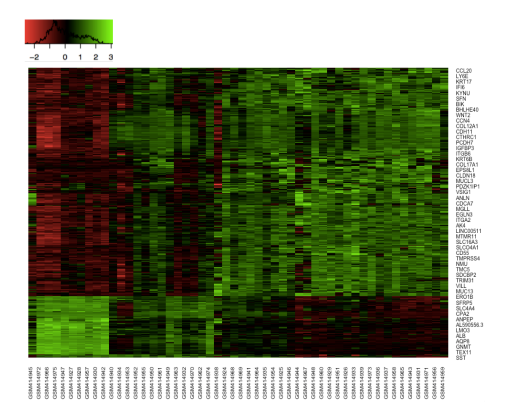


Figure 2 Heat map demonstrating DEGs in pancreatic tumor and normal cells aged samples. Each columns present samples, and rows present genes. Base-2 logarithmic values of the gene expression data are calculated. The progressive color changing from red to green represents the ranging from down to up-regulated DEGs.



3.2 Gene ontology and enrichment analysis

Table 2 shows the significant enrichment of DEGs using biological processes (BP) involving collagen catabolic process, extracellular matrix organization, collagen fibril organization and type I interferon signaling pathway. The significant enrichment of DEGs in cellular component (CC) contains extracellular space, extracellular exosome, and extracellular region. Finally, the significant enrichments GO terms in molecular function (MF) is revealed metalloendopeptidase activity, extracellular matrix structural constituent, and heparin binding.

KEGG signaling pathway study outcomes demonstrated that the DEGs were considerably enriched in ECM-receptor interaction, focal adhesin, PI3K-Akt signaling pathway, and central carbon metabolism in cancer. Further, PI3K-Akt signaling pathway has vital influence on multiple cellular process which also has a role in cancer pathways almost in all cells (Jiang et al., 2020).

 $\textbf{Table 2.} \quad \text{Gene expression data set retrieved with top significant pathways GO enrichment analysis of DEGs in pancreatic cancer. } \\$

Category	Term	Count	<i>p</i> -value	Genes
BP	GO:0030574~collagen catabolic process	14	1.38E-10	MMP7, MMP1, COL11A1, COL12A1, KLK6, MMP12, COL1A1, MMP11, ADAMTS2, COL1A2, COL5A1, COL5A2, COL8A1, COL10A1
BP	GO:0030198~extracellular matrix organization	19	2.75E-08	ERO1A, ERO1B, POSTN, LAMB3, ITGB4, COL11A1, ITGA2, LAMA3, FN1, LAMC2, COL1A1, COMP, VCAN, COL1A2, COL5A1, COL5A2, COL8A1, COL10A1, ITGB6
ВР	GO:0030199~collagen fibril organization	10	3.05E-08	GREM1, COL1A1, MMP11, ADAMTS2, COL1A2, COL5A1, COL11A1, COL12A1, COL5A2, LOXL2
BP	GO:0060337~type I interferon signaling pathway	10	2.64E-06	ISG20, IFI27, RSAD2, OAS1, OAS2, MX1, IFI6, ISG15, XAF1, OASL
CC	GO:0005615~extracellular space	77	6.52E-20	
СС	G0:0070062~extracellular exosome	105	4.85E-14	
СС	GO:0005576~extracellular region	72	6.01E-13	
СС	GO:0005578~proteinaceous extracellular matrix	22	1.67E-08	COL17A1, ECM1, POSTN, MMP7, MMP1, COL11A1, FN1, ADAMTS12, MMP12, COMP, MMP11, ADAMTS2, VCAN, COL1A2, COL5A1, COL5A2, MMP28, COL10A1, PI3, MUC4, WNT2, CTHRC1

MF	GO:0004222~metalloendopeptidase activity	12	6.95E-06	MMP12, MMP11, ADAMTS2, ADAM28, MMP7, FAP, MMP1, ADAM12, MMP28, ADAM9, ADAMTS12, KLK7
MF	GO:0005201~extracellular matrix structural constituent	9	2.87E-05	COL1A1, COMP, VCAN, COL1A2, COL5A1, COL11A1, COL5A2, MUC5B, MUC5AC
MF	GO:0008201~heparin binding	13	3.81E-05	POSTN, AOC1, MMP7, REG4, F11, FN1, LAMC2, CEL, THBS2, COMP, CXCL10, COL5A1, LIPH
KEGG	hsa04512:ECM-receptor interaction	15	3.15E-09	LAMB3, ITGB4, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, COL1A1, COMP, COL1A2, COL5A1, COL5A2, SDC1, ITGB6
KEGG	hsa04510:Focal adhesin	17	3.15E-09	LAMB3, ITGB4, EGF, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, COL1A1, COMP, COL1A2, COL5A1, COL5A2, ITGB6, PAK3, MET
KEGG	hsa04511:PI3K-Akt signaling pathway	17	02E-2	LAMB3, ITGB4, EGF, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, EFNA5, COL1A1, COMP, COL1A2, COL5A1, COL5A2, ITGB6, MET
KEGG	hsa04 05230:Central carbon metabolism in cancer	6	0.1E-1	PKM, SLC2A1, SLC16A3, MET, HK2, PFKP

Abbreviations- gene ontology: GO; biological process: BP; cell component: CC; Kyoto Encyclopedia of Genes and Genomes: KEGG (by the p value)

3.4. The PPI-network and KEGG pathway enrichment

Fig. 4 shows the PPI map between the set of input DEGs. The expressions of the nodes and their degree of connection were symbolized by yellow to red and fields, respectively in the visualized networks. The genes with the best three scores according to the *p value* is identified; these proteins also determines the functionality of the PPI network. Best scoring genes comprise with PPI network KEGG enrichment analysis represents involvement of ECM-receptor interaction (hsa04512), amoebiasis (hsa05146), protein digestion and absorption (hsa04974), focal adhesion (hsa04510), and PI3K-Akt signaling pathway (hsa04151). The other significantly enriched pathways associated with pancreatic tumor and normal cells DEGs are ECM-receptor interaction (hsa04512) and Central carbon metabolism in cancer (hsa0405230). Furthermore, the enriched KEGG pathways also involved the 'PI3K-Akt signaling pathway' due to playing a key function in cell and tissue growth which is also associated with the cancerous cells.

PPI network of DEGs was created and pictured using NetworkAnalyst. Progressive color change from yellow to the red represents relative gene expression value from down to up-regulated genes. CDK1 with the largest Betweenness Centrality (BC) was suggested to be central to the PPI network associated with pancreatic tumor and normal cells gene expression data. Moreover, CCNB1, CDC20, PPARG, MET, and ISG15 with the secondary highest degree and SFN with the secondary BC might be engaged in the progression pancreatic adenocarcinoma and other substance diseases.

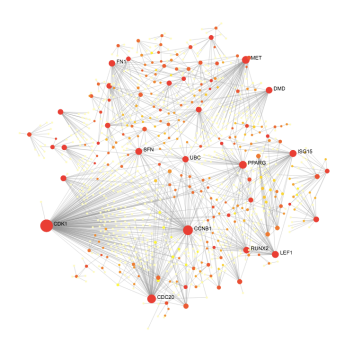


Fig.4 demonstrates PPI network of DEGs in pancreatic tumor and normal of all the DEGs . To conclude, PIP-network of pancreatic tumor and normal of all the DEGs (see Fig.4 and Table 3) in the expression data set. Hub genes in Fig.4 can be listed as CDK1, CCNB1, CDC20 , PPARG, MET, and ISG15. SFN gene is active in protein degradation, DNA repair, cell cycle regulation, kinase modification, endocytosis, and regulation of other cell signaling pathways. UBC is a key gene that directly engages with other genes such as LC3, propose that it may be a central component that leads to a bad prognosis of pancreatic cancer regulated by perineural invasion (Yang et al., 2019). ECT2 has a vital task in diseases comprising breast cancer and lung cancer. Related pathways of ECT2 are signaling pathways such as p75 NTR receptor-mediated signaling. GO annotations of molecular function related to this gene contain protein homodimerization activity and GTPase activator activity.

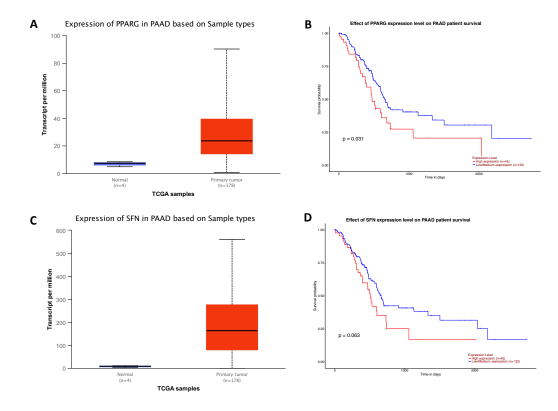
Table 3 Top 15 genes of PIP network of DEGs in pancreatic tumor and normal cells gene expression data.

		Node	Betweenness	
Gene ID	Genes	Degree	centrality	Expression
983	CDK1	191	170861.12	6.95
891	CCNB1	91	42477.27	6.05
991	CDC20	66	30778.32	5.08
5468	PPARG	48	43020.9	6.84
4233	MET	47	76573.47	9.87
9636	ISG15	37	31351.4	8.86
51176	LEF1	36	52887.37	6.81
2810	SFN	34	28856.95	8.26
1756	DMD	32	37877.67	7.59
2335	FN1	30	36750.92	12.61
860	RUNX2	30	26338.94	7.37

7316	UBC	28	156179.54	8.81
7153	TOP2A	24	18346.88	6.74
1894	ECT2	23	16342.09	7.29

3.5. Tumor gene expression and survival analyses through UALCAN portal

Notably, PPARG and SFN genes were the two genes related to survival. Once the detection of hub genes of PPI network was revealed, survival analysis for PAAD was carried out through UALCAN. PPARG; peroxisome proliferator-activated receptor gamma and SFN; Stratifin were shown to be significantly related with survival (P<0.05). Afterwards, the differentially expressed genes in pancreatic adenocarcinoma were compared; PPARG and SFN were identified to be significantly differentially expressed. The expression levels of PPARG were analyzed using TCGA database, and the outcomes were consistent with those of the mentioned differential gene analysis; PPARG was significantly upregulated in PAAD tumor tissues compared with normal tissues $p=1.62\times10^{-12}$. In addition, SFN is also upregulated with in PAAD tumor and normal tissues with a p-value $< 10^{-12}$.



3.6. Validation of PPARG and SNF

45 tumor and 45 normal specimens of DEGs were studied. The expression level of PPARG and SFN was studied specifically. The outcomes of the analysis to validate the significance of PPARG and SFN are demonstrated in Table 4; it was noted that PPARG and SFN was significantly upregulated in pancreatic adenocarcinoma cells in the two of the GEO datasets.

Table 4 Base-2 logarithmic scale of differential expression of most significant hub genes in pancreatic cancer cells in two different datasets.

Datasets	Genes	FC	P-value	
GSE16515	PPARG	6.85	1.62E-12	
	SFN	5.04		
GSE28735	PPARG	6.17	<1E-12	
	SFN	4.8		

3.7. Analysis of the PI3K-Akt signaling pathway

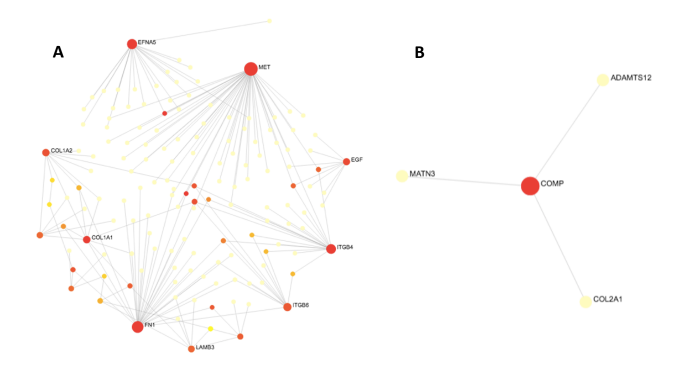
The core of this project due to it is close association with cancer within all of the significantly (*p value* < 0.01) enriched pathways of DEGs is the PI3K-Akt signaling pathway. There were 17 DEGs particularly engaged in this pathway, containing LAMB3, ITGB4, EGF, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, EFNA5, COL1A1, COMP, COL1A2, COL5A1, COL5A2, ITGB6, MET (Fig. 5 and Table 2 and 4). We have performed the primary DEGs associated with PI3K-Akt signaling pathway in Fig.6. We implemented PI3K-Akt signaling pathway genes to construct Fig.5A showing Subnetwork 1 and Fig.5B showing Subnetwork 2 protein-protein interaction networks.

Associated genes with the DEGs of the data set enriched with the PI3K-Akt signaling pathway deciphered as new hub genes. In subnetwork 1 of PI3K-Akt pathway (Fig.5A), MET, FN1, and ITGB4 is the most significant genes in terms of BC. Moreover, collagen (COL) gene family also performs a key position which is demonstrated in the subnetwork of the PI3K-Akt signaling pathway. As shown in subnetwork 2, cartilage oligomeric matrix protein, COMP might be a gene that is associated with PI3K-Akt signaling pathway. We also found that the proteins associated with COMP gene in PI3K-Akt signaling pathway can be listed as ADAMTS12, MATN3, and COL2A1 in pancreatic tumor and normal cells gene expression data set. Even though the functional roles of other family members, such as ADAMTS12, remain unknown, downregulation or upregulation of particular ADAMTS proteins activated in serious human diseases, involving cancer. (Llamazares et al., 2007). These outcomes confirm the key task of the PI3K-Akt signaling pathway engaged in pancreatic cancer and related diseases treatment, offering new molecular therapeutic targets to improve fundamental drug agents.

Table 5 Top 10 most excessive KEGG pathway enrichment analysis of global DEGs in pancreatic tumor and normal cells micro-array gene expression data set.

Term	Count	P-value	Genes
hsa04512:ECM-receptor interaction	15	3.15E-09	LAMB3, ITGB4, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, COL1A1, COMP, COL1A2, COL5A1, COL5A2, SDC1, ITGB6
hsa05146:Amoebiasis	13	2.27E-06	SERPINB3, CXCL8, SERPINB2, LAMB3, IL1R2, COL11A1, LAMA3, FN1, LAMC2, COL1A1, COL1A2, COL5A1, COL5A2
hsa04974:Protein digestion and absorption	12	2.28E-06	COL17A1, COL1A1, CPA2, COL1A2, CTRL, COL5A1, COL11A1, COL12A1, COL5A2, COL10A1, SLC16A10, KCNN4
hsa04510:Focal adhesion	17	7.13E-06	LAMB3, ITGB4, EGF, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, COL1A1, COMP, COL1A2, COL5A1, COL5A2, ITGB6, PAK3, MET
hsa04151: PI3K-Akt signaling pathway	17	0.00271604	LAMB3, ITGB4, EGF, COL11A1, ITGA2, LAMA3, FN1, LAMC2, THBS2, EFNA5, COL1A1, COMP, COL1A2, COL5A1, COL5A2, ITGB6, MET
hsa05230:Central carbon metabolism in cancer	6	0.01154629	PKM, SLC2A1, SLC16A3, MET, HK2, PFKP
hsa00830:Retinol metabolism	6	0.01154629	SDR16C5, DHRS9, ADH1B, AOX1, UGT1A8, CYP2C18
hsa04610:Complement and coagulation cascades	6	0.01564074	C6, PLAU, F11, PLAUR, CD55, F5
hsa05144:Malaria	5	0.02015542	COMP, CXCL8, SDC1, THBS2, MET
hsa05200:Pathways in cancer	16	0.02027915	EGLN3, CXCL8, LAMB3, MMP1, EGF, ZBTB16, ITGA2, LAMA3, LEF1, SLC2A1, FN1, LAMC2, CKS2, PPARG, MET, WNT2

 $\begin{tabular}{ll} Figure 5 & Hub genes associated with $PI3K$-$Akt signaling pathway. (A) Subnetwork 1 and (B) Subnetwork 2 protein-protein interaction networks. \\ \end{tabular}$



4 Discussions

The prevalence of pancreatic adenocarcinoma and the related survival rates have demonstrated a decreasing in tendency in the past years (Siegel et al., 2018). One study showed that PAAD patients survive for only 4 months typically without therapies. Moreover, patients who undergo surgery and take required therapies the survival is not significantly increased (Wang et al., 2015). As a result, precise quick identification of PAAD and the advancement of powerful specific remedy is of fundamental significance.

A recent research detected hub genes in PAAD that were stated to be of diagnostic significance (Lv et al., 2019). In our study, the microarray GSE16515 from the GEO was conducted, holding data of 36 tumor and 16 normal cells (Pei et al., 2009). Differentially expressed genes were only analyzed between tumor and normal cells. 278 upregulated and 77 downregulated differentially expressed genes were identified using R, and GO (Gene Ontology Consortium, 2004) and KEGG pathway analyses which revealed the information of locations and functions of these DEGs. Upregulated DEGs were primarily located in the collagen trimers and extracellular exosome, and were involved in cell adhesin, ECM organization and proteolysis, positive regulation of cell proliferation, collagen catabolic process and signal transduction. Contrary, downregulated DEGs were primarily enriched in proteolysis, digestion, and apoptotic process.

Finally, protein-protein interaction network construction reveal information about the hub genes is used to identify function and disease associated with proteins. The highest interactive proteins CDK1, CCNB1,

CDC20, PPARG, MET, LEF1, SFN, DMD, FN1, UBC, TOP2A, ECT2, WNT2, EFNA5, PAK3, PKM, ITGB4, NEK2, and ALB are predicted that are involved in several types of cancers like retinoblastoma, breast cancer, rhabdomyosarcoma, diabetes, childhood obesity, ovarian cancer, acute lymphoblastic leukemia, oral squamous cell carcinoma, colorectal and pancreatic cancer (Santiago et al., 2017; Sun et al., 2020; Tovar & Graveel, 2017; Tyagi et al., 2011). ISG15 pathway plays an important role in the tumorigenesis and treatment of cancers of the digestive system. (Zuo et al., 2016, p. 15). RUNX2 gene can be employed as biomarkers for diagnosis of early stage of pancreatic cancer can also operate as promising a pro-oncogenic potential (Ozaki et al., 2018).

Finally, 21 hub genes were identified using a PPI network and the prognostic value of these hub genes for patients with PAAD was studied using UACLAN. Utilizing data from TCGA that both *PPARG* and *SFN* were significantly upregulated in PAAD cells compared with control cells, as per the results of the differential gene analysis. Further, it was revealed that *SFN* and *PPARG* were considerably related with poorer survival. Thus, these two genes may be associated with the carcinogenesis and angiogenesis of PAAD.

After GO and KEGG pathway enrichment analyses, the functional enrichment of peroxisome proliferator activated receptor gamma (PPARG) was studied. *PPARG* is primarily involved in the modulation of fatty acid storage and glucose metabolism. SFN plays a function in kinase modification, endocytosis, protein degradation, DNA repair, cell cycle regulation, and modulation of other cell signaling pathways. Increased expression of PPARG might be the cause of the pathology of various diseases such as diabetes, obesity, atherosclerosis and pancreatic cancer. Therapies of inhibiting SFN downregulates the expression of Bcl-2 and XIAP to stimulate apoptosis that directs to the downsizing in EMT, car, carcinogenic, angiogenic biomarkers with considerable suppression in tumor development in mice (Li et al., 2013).

This investigation further highlighted the PI3K-Akt signaling pathway involving differentially expressed genes in a broad various kinds of human cancer (Jiang et al., 2020). The significant role of the PI3K-Akt signaling network in regulating cell proliferation, the cell cycle and apoptosis over the last 20 years was known (Hennessy et al., 2005; Rodon et al., 2013). Furthermore, mutations employ in PI3K pathway genes and other-associated regulated pathway genes contribute to pancreatic ductal adenocarcinoma tumorigenesis. In addition their standard roles, PI3K further rules metabolism characteristics of cancer cell and tumor microenvironment-mediated mechanisms of neoplasm development and survival (Murthy et al., 2018).

The presented project has certain limitations. In studying the expression level both PPARG and SFN only two datasets were analyzed, and more gene expression data should be investigated with a higher number of control samples is a need to verify the presented results. Secondly, additional studies is required for clinical lab confirmation of predicted proteins that are expressed in pancreatic tumor and normal cells data set and express at the developmental stage of pancreatic adenocarcinoma. More research is needed in the field of cancer biology to detect pancreatic cancer and subset diseases at its early stage. This paper also emphasizes the importance of microarray experiment in comprehending pancreatic cancer and related diseases and approach to study several results of gene expression data, like differentially expressed genes analysis, pathway and process identification, and protein-protein interaction network study.

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