

## Exploring genes of rectal cancer for new treatments based on protein interaction network

Running Title: protein interaction network of rectal cancer

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Abstract

**Objective:** To develop a protein-protein interaction network of rectal cancer, which is based on genetic genes as well as to predict biological pathways underlying the molecular complexes in the network. In order to analyze and summarize genetic markers related to diagnosis and prognosis of rectal cancer.

**Methods:** the genes expression profile was downloaded from OMIM (Online Mendelian Inheritance in Man) database; the protein-protein interaction network of rectal cancer was established by Cytoscape; the molecular complexes in the network were detected by Clusterviz plugin and the pathways enrichment of molecular complexes were performed by DAVID online and Bingo (The Biological Networks Gene Ontology tool).

**Results and Discussion:** A total of 127 rectal cancer genes were identified to differentially express in OMIM Database. The protein-protein interaction network of rectal cancer was contained 966 nodes (proteins), 3377 edges (interactive relationships) and 7 molecular complexes (score>7.0). Regulatory effects of genes and proteins were focused on cell cycle, transcription regulation and cellular protein metabolic process. Genes of *DDK1*, *sparcl1*, *wisp2*, *cux1*, *pabpc1*, *ptk2* and *htral* were significant nodes in PPI network. The discovery of featured genes which were probably related to rectal cancer, has a great significance on studying mechanism, distinguishing normal and cancer tissues, and exploring new treatments for rectal cancer.

**Keywords:** Protein-protein interaction networks; rectal cancer; molecular complexes; pathway ,

genes

Rectal cancer is one of the common fatal malignant tumors in the world, which is secondly lethal cancer in United States as well as thirdly in Europe. The research has improved systemic treatment, such as micro diagnosis. However, despite advances in detection and care, morbidity and mortality from rectal cancer continues to be high[1]. Early detection and diagnosis can be great significance to reduce mortality and improve prognosis, as well as identify those who were at the highest risk due to improving triage for treatment has the greatest impact on rectal cancers treatment. Studies shown that rectal cancer has a prime paradigm for cancer genetics, which can be prevention by early detection of the pre-disease (neoplastic) state. Therefore, the roles of genetics in rectal cancer have become critical to the missions of disease prevention, early detection and effective treatment.

As report goes, approximately 10% of well defined hereditary rectal cancer has syndromes[2]. Hereditary rectal has some benefits, such as many of it are growing slowly, family members are armed with the knowledge of potential risk of associated carcinomas and empowerment to reduce the disease burden.

A large number of mutation, mismatch, inactivation of tumor suppressor genes and activations of oncogenes are involved in the process of rectal cancer[4]. Up to now, genetic marker of rectal cancer can be summarized by the following six aspects : genomic instability, CpG island methylator phenotype, specific microRNAs, histone modification, gene mutation and protein biomarkers. The protein-protein interaction network which is a model of biological molecular interactions can more clearly show the genes, proteins and pathogenesis in the development process of the disease[5]. These genetic traits may partially explain the geographical variance in rectal cancer incidence and mortality as well as the differences between hereditary and sporadic rectal cancer [6].

## 1 Materials and Methods

**Design:** Study on enrichment of genomic biological pathways.

**Method:**

**Data acquisition:** OMIM(Online Mendelian Inheritance in Man) is a comprehensive, authoritative, daily updated human phenotype database, containing more than 12000 genes of all human genetic diseases, and mainly focusing on hereditary diseases. In addition, text messages, related reference information, sequence records, maps, and related databases are available for each gene[7-8]. This study had started from August 22(2015), searched "rectal cancer" in the OMIM database and obtained human genes associated with rectal cancer information.

**The construction of gene/protein interaction networks:** rectal cancer associated genes were submitted to Cytoscape 3.2.1 plug-in Agilent Literature Search 2.7.7 (USA Agilent Technologies company) and Pubmed [9]. False positive interaction information was removed from retrieval results. Then, gene/protein interaction relations were read in Cytoscape 2.8.2 and visualized [10].

**Network analysis:** MODE algorithm in Cytoscape 3.2.1 web analytics plug-in Clusterviz of 1.2 was administrated to make the correlation analysis for the area of the construction of biological networks [11-12]. By analyzing the network structure, proteins were grouped to form molecular compounds in the entire network and were shown in Cytoscape according to the correlation integral value. The areas with integral value higher than 3 were regarded as molecular compounds. The gene/protein names contained in the molecular compounds were submitted to The Database for Annotation, Visualization and Integrated Discovery. [13-14] By retrieving Kyoto Encyclopedia of Genes and Genomes (KEGG) Database, biological pathways involved in chronic myelogenous leukaemia heredity were identified. Then the Biological pathways data were submitted to **Bingo** (in Networks Gene Ontology tool) for enrichment analysis.

**Main outcome measures:** Protein networks were constructed based on the rectal cancer-related genes, nodes (proteins) and edges (interaction between), molecular complexes in the network and its associated interaction points and nodes (protein) and the edge (interaction between), analyze the biological pathways has involved in the molecular complexes.

## 2 Results

## 2.1 Rectal cancer related genes in OMIM

According to the OMIM database retrieval, it can be found that 127 genes were reported to be associated with rectal cancer. After screening and deleting duplicate genes, 127 related genes were identified, which are shown in **Table 1**.

## 2.2 Protein interaction networks

Through text mining, 127 genetic-related genes were shown that there was a network diagram with 638 nodes (proteins) and 1830 edges. As shown in **Figure 1-3**, the diamond represents OMIM genetic disease related proteins, while the round represents the proteins obtained from text mining.

### **Figure 1(A+B). Network map of protein interaction (Overall)**

As shown in **Figure 1**, the protein-protein interaction network underlying rectal cancer is extremely complex. The edges intersect with each other and several clusters emerge in the figure. The more edges among the genes, the correlation of the genes is more tightening. So the genes formed the round network at the top of **Figure 1** connected with each other more tightly than the genes in the bottom.

### **Figure 2 the relationship of proteins**

After removing the protein molecule in the network, the relationship between proteins in the network became more clear. In the centre part of the protein-protein network, the relationship between protein complexes emerged is closer, however, at the farther edge of the network, the relationship of it became looser, as shown in **Figure 2**.

### **Figure 3(A): pathway jak2-stat1**

### **Figure 3(B): relationship of gene *Kras***

AS shown in **Figure3**, it can be found some common genes and pathways such as jak2-stat1, *Kras*, P53, etc as well as genes associated with them.

#### **2.3 Network topology attribute analysis**

Network topology attribute analysis shown that the connectivity of nodes in the network (the number of nodes in the network) obeys descending distribution, i.e. with the increase of edges connected to the node, correspondingly the number of nodes decrease, so it can be seen that the gene-protein interaction networks are scale-free networks [15]. We found that the connectivity of nodes in the network greater than or equal to 25 corresponds to a sharp reduction in the number of nodes **Figure 4**. Therefore, we regarded the nodes which the connectivity is greater than / equal to 20 as the key nodes (hub). Key nodes (connectivity score) included: *hif1a*(25), *cdkn1b*(25), *rb1*(26), *plau*(26), *brca1*(27), *asns*(28), *pcna*(28), *Vegfa*(29), *stat1*(29), *cdkn2a*(29), *egfr*(30), *tp53*(71).

#### **Figure 4. Connectivity degree of each node and betweenness Comparison**

(The horizontal axis represents betweenness, and the ordinate represents the connectivity degree of protein interaction network. And the graphic in the table represents each node in the network). It can be seen that the connectivity (the number of nodes in the network) of nodes in the network obeys descending distribution, while the connectivity is greater than / equal to 25, the number of nodes corresponding to a sharp decrease.

#### **2.4 The detection of molecular complexes**

According to MCOMD algorithm analysis, analyse the correlation between genes in network and calculate the score, the number of nodes, the number of edges. There is a total of 74 molecular complexes and 7 of them showed correlation integral values higher than 7.0.

## Figure 5.

**Rank1** score20 Node20 Edges190

**Rank2** score20 Node20 Edges190

**Rank3** score14 Node14 Edges91

**Rank4** score11.662 Node78 Edges449

**Rank5** score3.143 Node21 Edges66

**Rank6** score9 Node9 Edges36

**Rank7** score7.673 Node56 Edges221

**Figure 5 Molecular complexes obtained by MCOMD algorithm analysis**

The details of the relationship of proteins in molecular complexes 7 was shown in Figure 6

### **Figure6 the relationship of proteins in molecular complexes 7**

The whole network has a total of six parts, including mitogen-activated protein kinase, Specific microRNAs, SRY-related HMG-box, matrix metalloproteinases, cyclinD.

#### **2.5Molecular complex pathway enrichment**

Submit the 7 names of protein molecule complexes online to obtain the relevant pathways in

**Table 2.** The Protein molecule biological pathways of complexes 1 are not exist, and other complexes contain different pathways. Regulatory effects of genes and proteins mainly focused on cell cycle, transcription regulation, and cellular protein metabolic process.

**Table 2 Enrichment of pathways related to molecular complexes**

Bingo results show the gene oncology hierarchical network to the biological processes, **Figure 7**. The size of the node represents the number of the genes, the depth of the node color represents the P values. Diagrams have presented the main biological processes of cluster 2, 3, 5, containing metabolic regulation, transcriptional regulation, biosynthesis, cell differentiation and gene expression regulation and signal transduction, etc.

### **Figure 7**

**A Bingo results show that the basic structure of the biological cluster 1**

**B Bingo results show that the basic structure of the biological cluster 4**

**C Bingo results show that the basic structure of the biological cluster 6**

**C Bingo results show that the basic structure of the biological cluster 7**

## **3 Discussions**

Based on a large number of research references, the progression of rectal cancer is a multistep process, containing overcome apoptosis, inhibiting senescence (infinite proliferation), secretion proliferation signal itself, not sensitive to growth signals, angiogenesis and invasion, which requires a large number of genes and proteins in action [16]. Genetic markers of rectal cancer can be summarized from the following six aspects: the first part is genomic instability, close to 12-17% of sporadic rectal cancer cases exist with microsatellite instability (MSI). At present, microsatellite instability - high - H (MSI) has become a positive prognosis of rectal cancer patients' overall survival [17]. 50-85% of Rectal cancer patients has chromosome aberration frequency, chromosome instability (CIN+) usually be associated with colorectal cancer patients' overall survival, progression-free survival (PFS) and poor prognosis after 5 - fluorouracil therapy; the second part is CpG island methylator phenotype. Nearly 29.6% of the rectal cancer patients shown CpG island methylation phenotype - high (CIMP - H) [18], however, its value is still in research. Specific microRNAs and histone modification are another two aspects, which mainly associate with colorectal cancer patients' overall survival and progression-free serial, tumor metastasis, local

invasion, tumor volume, tumor staging, treatment results, relapse and drug resistance.

What is more, gene mutations and protein biomarkers have brought a special significance. The APC gene mutant p.D1822V containing homozygous V/V would reduce the risk of rectal cancer[19]. According to the APC (rs565453 y rs1816769) and CTNNB1 (rs229303) gene polymorphisms, the death risk stratification in patients with rectal cancer can be analysed. The APC gene mutant p.I1307K as risk factors of the rectal cancer among Ashkenazi Jewishes has been found[20]. The loss of PTEN gene in 22% of rectal cancer patients has caused no response to the EGFR inhibitors and higher risk of death [21]. And the clinical application of protein molecular markers colorectal cancer mainly included: early diagnosis(hnRNP A1,kininogen-1,adipophilin, Apo AI and C9,OLFM4), Prognosis(SM3,desmin,surviving,hTERT and NM23), Potential therapeutic targets(EB1)[22-23]. Therefore, exploring the protein interaction network of tumors, analysing the characteristics of the protein molecular interaction network, and excavating a variety of signalling pathway and the genes will provide a biological basis for the study of the molecular mechanism of tumor and the further treatment as well.

Bio-molecular network analysis is an important direction of systematic biology research. Large-scale human protein-protein network can provide new insights into protein functions, pathways, molecular machines and functional protein modules [24]. The function of bio-molecular often depends on modularization, the network module is made by a number of nodes in the conjunction of each other and has a stable structure which can often reflect a similar nature between the nodes [25,26]. Analysing the function module is the one of the most common method to analyse biological molecular network. According to 127 genes provided by OMIM, our research has built up protein interaction network of rectal cancer, which contain 996 nodes (proteins), 3377 edges (interaction). Due to the network is very large, the experimental introduced MCOMD algorithm to evaluate the network 's regional integration through the correlation integral. Correlation integral described the proteins associated with the degree in the region. Proteins of the same molecular complex generally have the same biological function, therefore we can discover the unknown gene functions or new molecular functional groups, such as cluster 1.

Gene *DDK1*,*sparcl1*,*wisp2*,*cux1*,*pabpc1*,*ptk2* and *htral1* as the composition members of the



cluster1(score 20) in the centre, have closely relationships with each other as well as other genes. It has reported that the secreted protein acidic and rich in cysteine-like 1 (*sparcl1*) is expressed in various normal tissues and many types of cancers. Another study has shown that marker *sparcl1* was significantly related to the prognosis and clinical pathological features of the CRC patients [27]. *Sparcl1* expression increased with RT and is related to a better prognosis in rectal cancer patients with RT but not in patients without RT. This result may help us to select the patients to the best suited preoperative RT. In the process of rectal cancer, gene WISP2 knockout significantly increased Caco-2 cell invasion and motility. Up-regulation of MMP2, -7 and -9 may indicate that WISP2 regulates invasion and motility through MMPs. Regulation of invasion by WISP2 may involve the WNT signalling pathway[28]. Thus, some studies have identified CUX1 as a pan-driver of tumorigenesis and uncover a potential strategy for treating CUX1-mutant tumors. So from cluster1 we predict the effect of gene Cut1 to rectal cancer may be that CUX1 deficiency activates phosphoinositide 3-kinase (PI3K) signalling through direct transcriptional downregulation of the PI3K inhibitor PIK3IP1 (phosphoinositide-3-kinase interacting protein 1), leading to increased tumor growth and susceptibility to PI3K-AKT inhibition. And we also can forecast a few clones existing in colorectal cancer, containing gene mutation of ptk2, htra1 and PABPC1.

Rectal cancer is demonstrated not simply controlled by a particular gene or signaling pathways, but also by the complex process of network system co-ordinately regulated which consisted of a variety of signalling pathways and multiple genes. In the signalling network, it is likely there is some "key regulatory point". At last, a best understanding of chemotherapy molecular targets allowed the identification of genetic markers that can predict the response and/or the toxicity of anti-cancer drugs used in rectal cancers, which could be helpful in the future to propose for each patient a personalized treatment [29-31]. Mutations that can predict the response of new target therapies such as the inhibitors of the JAK kinase inhibitor AG4 90 in colorectal cancer have also been found and will allow the selection of patients who can have benefit from these new therapeutic drugs. The experiment dig out a variety of signalling pathways and genes which can provide reliable directions for molecular mechanism research of treatment, and it need to be further verified.

## Compliance with Ethical Standards

**Conflict of interest** : We declare that we have no financial and personal relationships with other people or organizations that can inappropriately influence our work, there is no professional or other personal interest of any nature or kind in any product, service and/or company that could be construed as influencing the position presented in, or the review of, the manuscript entitled exploring genes of rectal cancer for new treatments based on protein interaction network

**Ethical approval:** This article does not contain any studies with human participants or animals performed by any of the authors.

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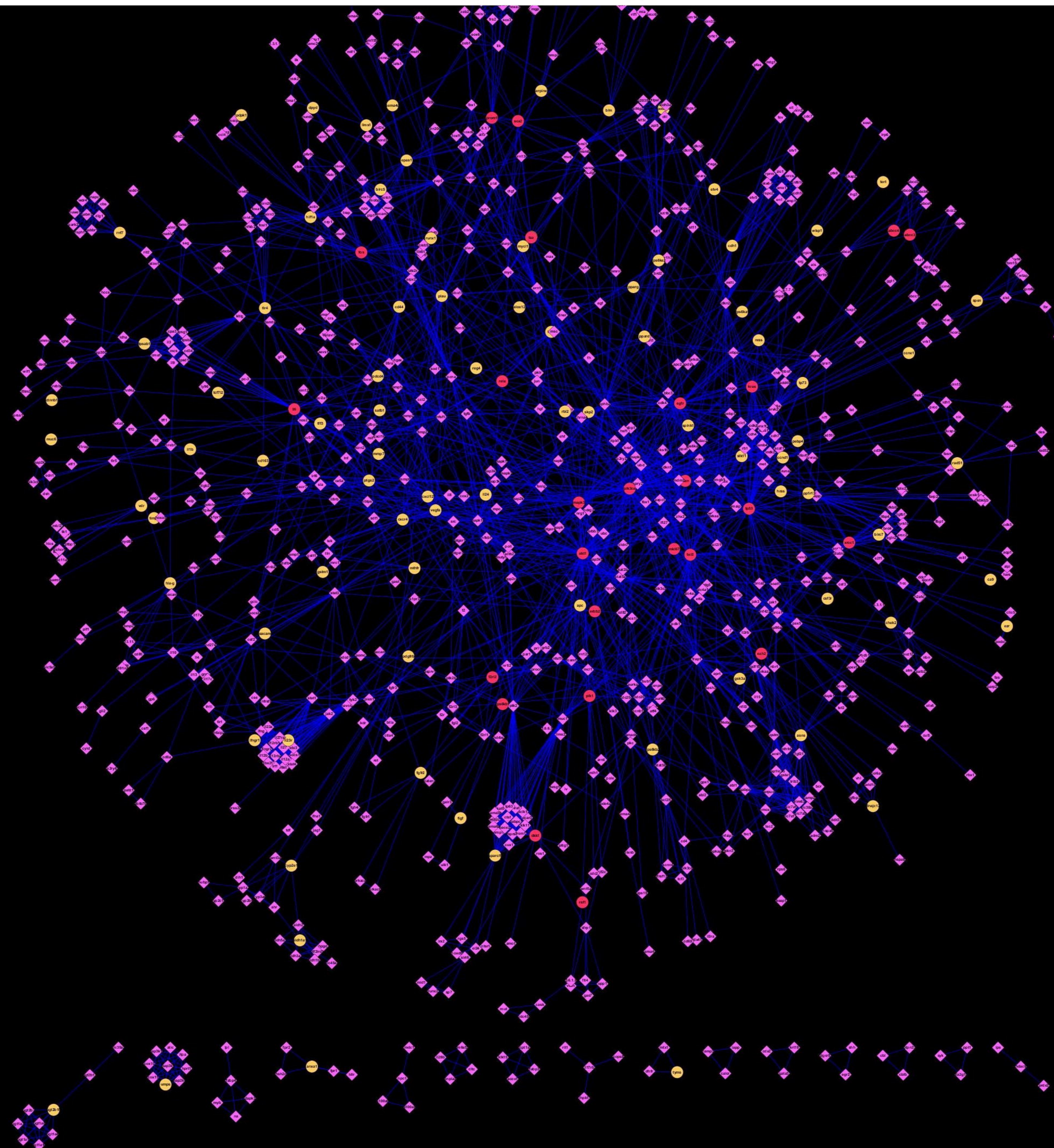
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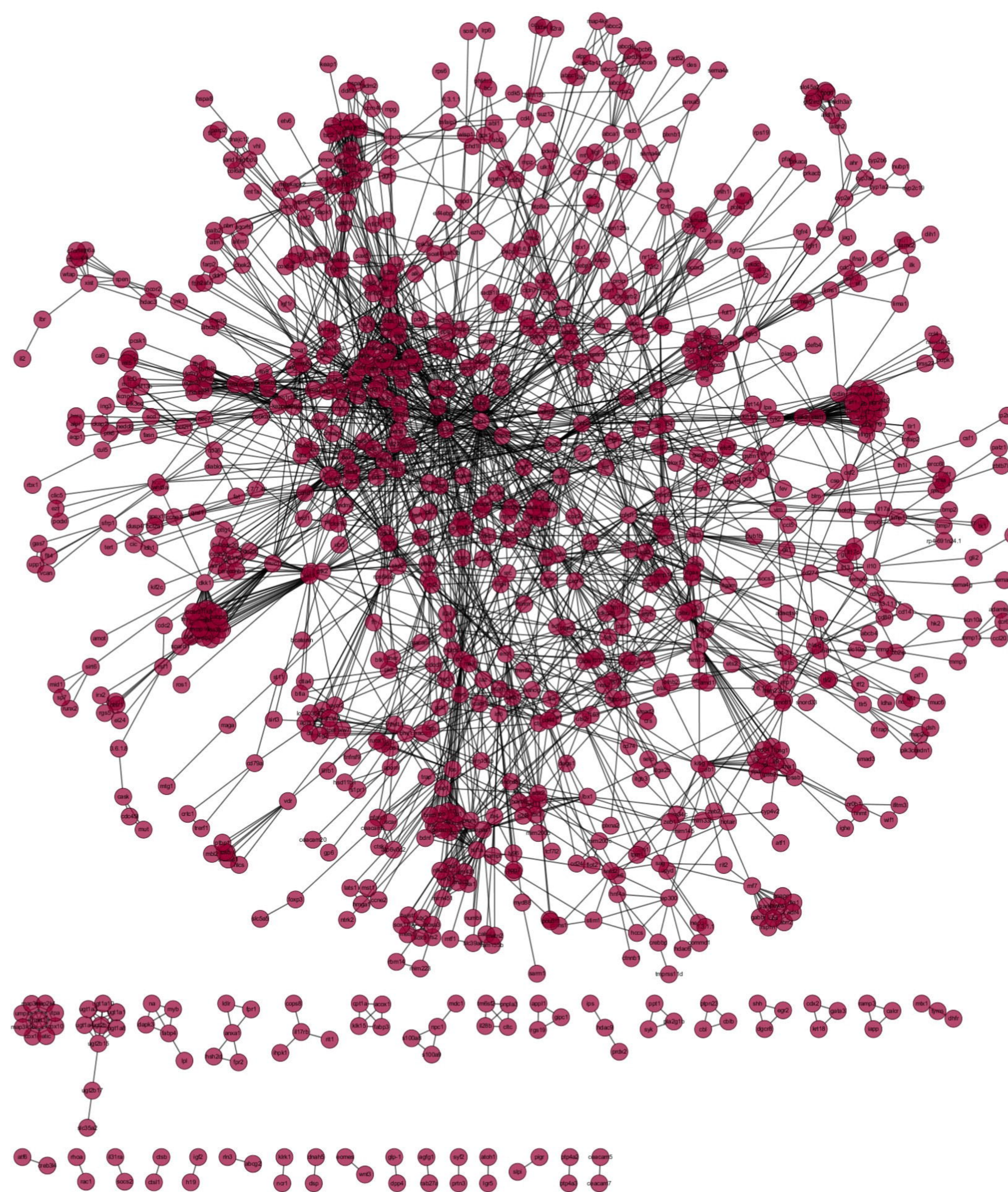
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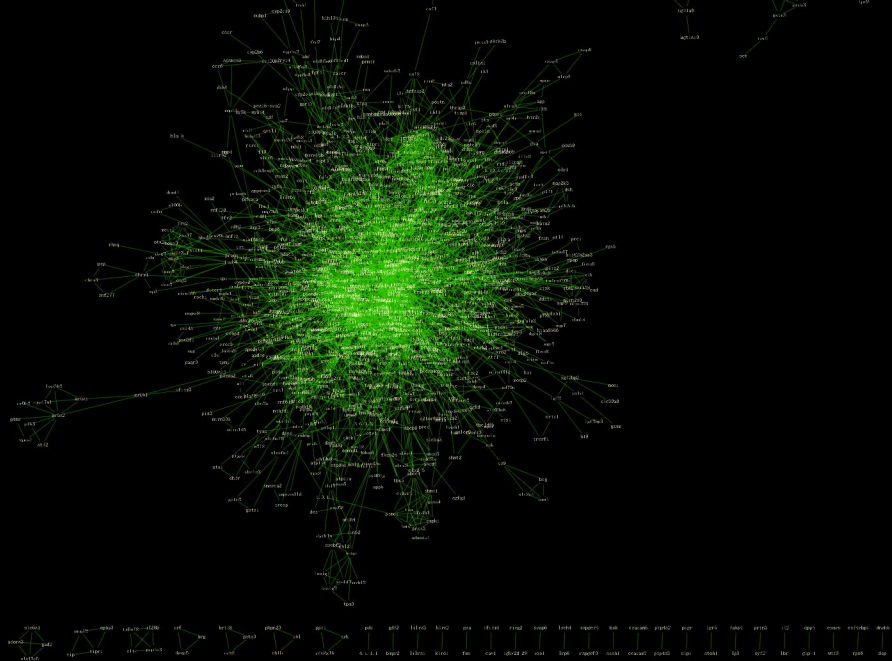


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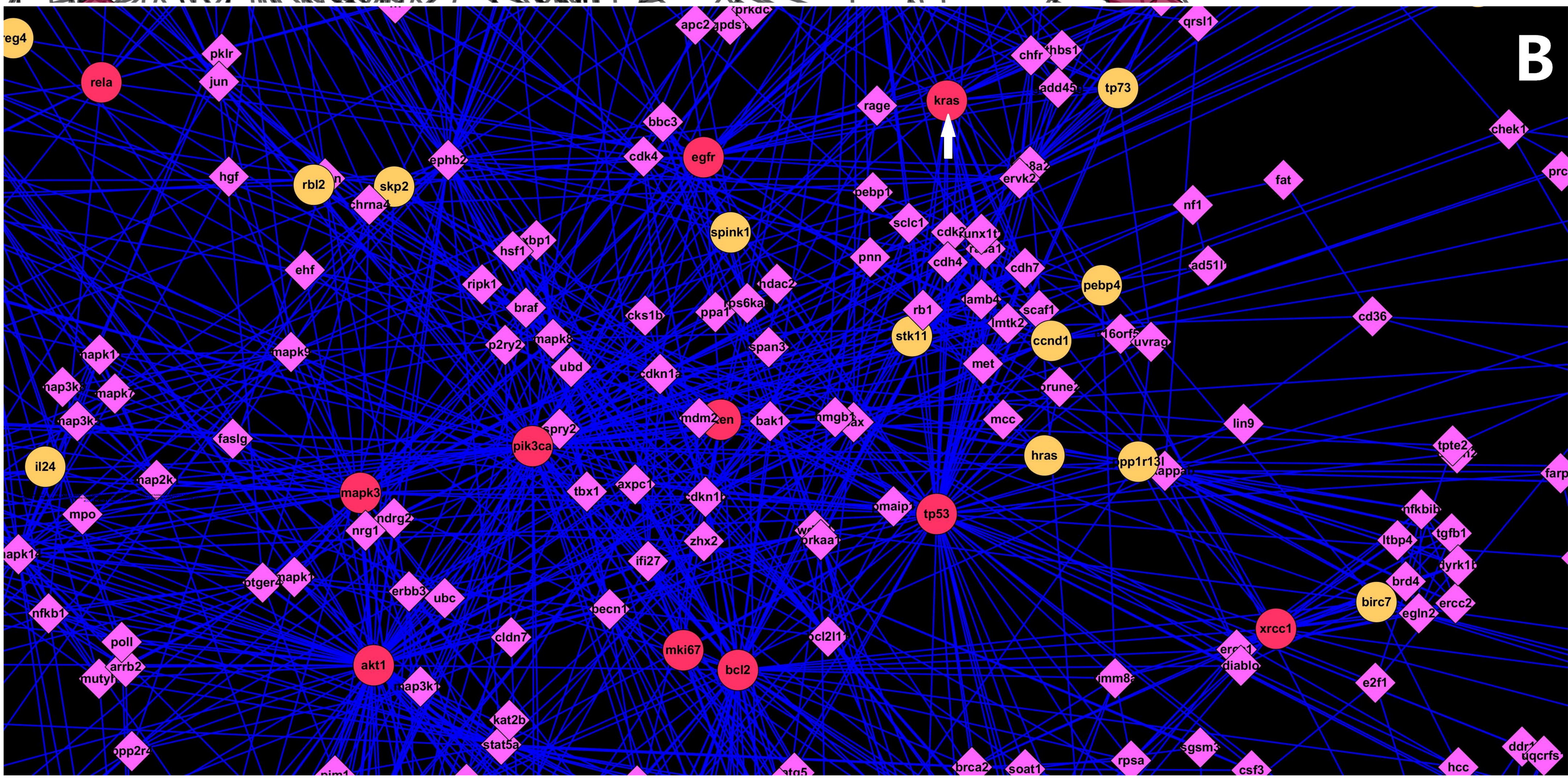
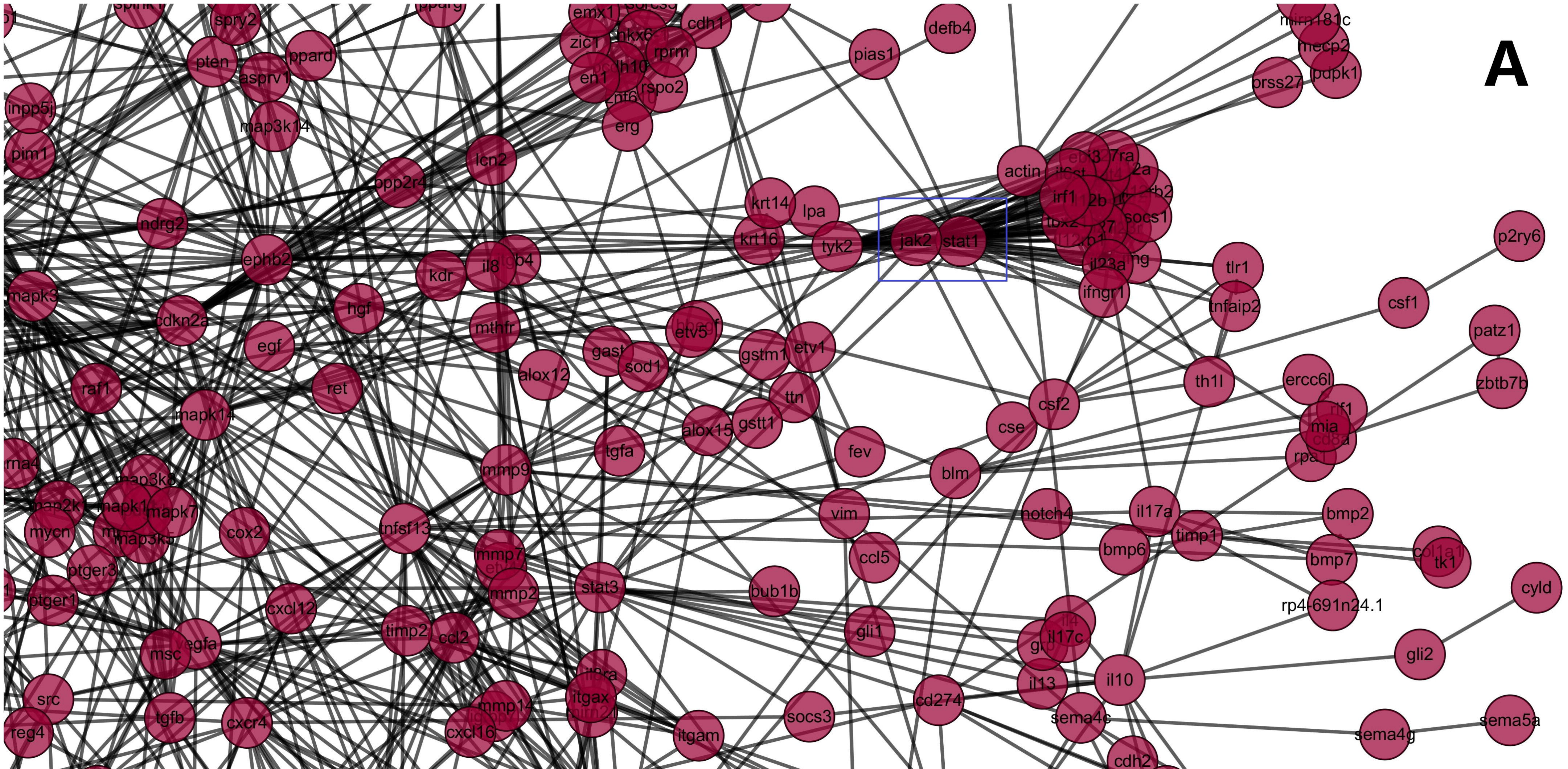
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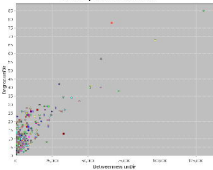
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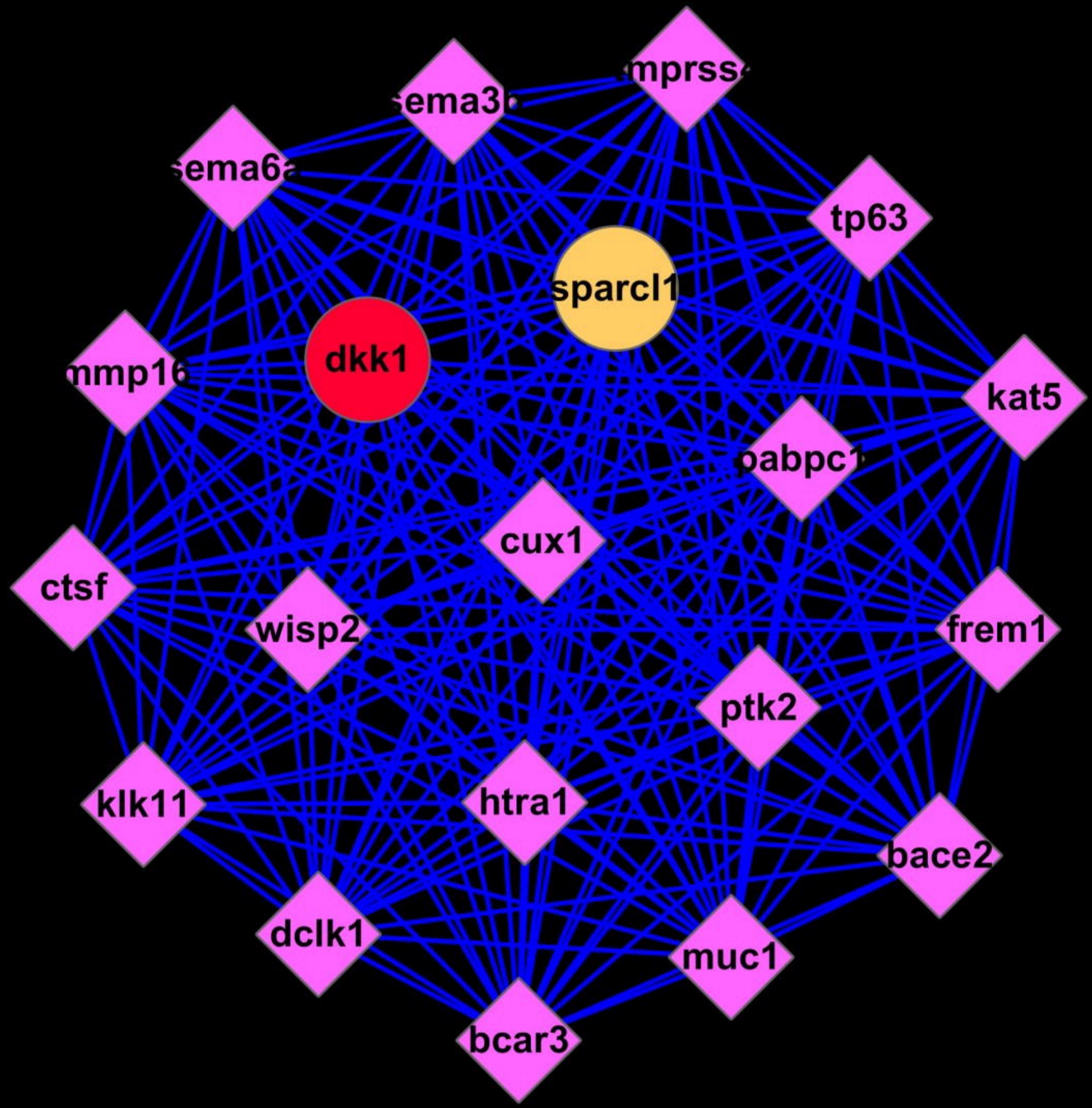




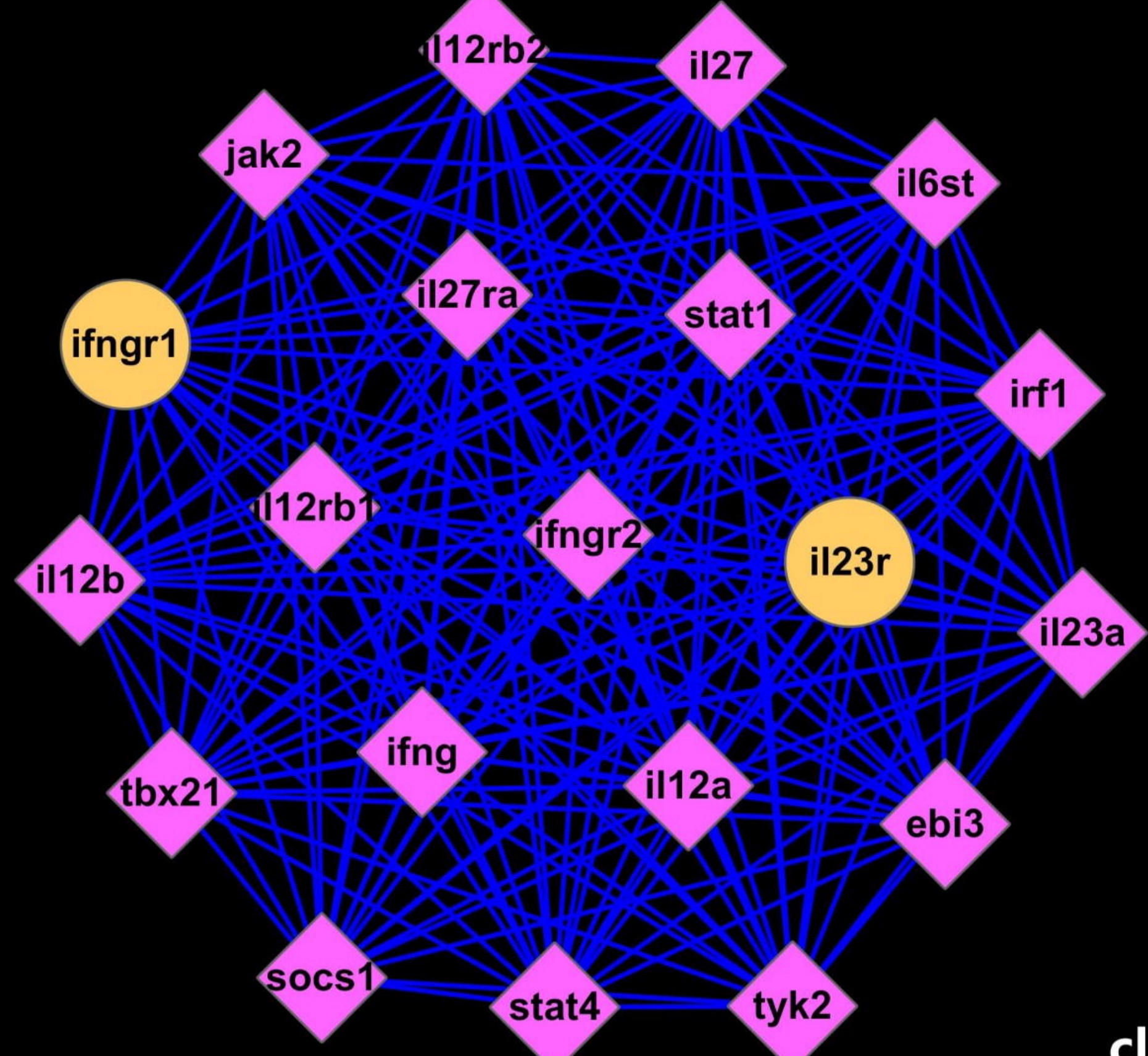
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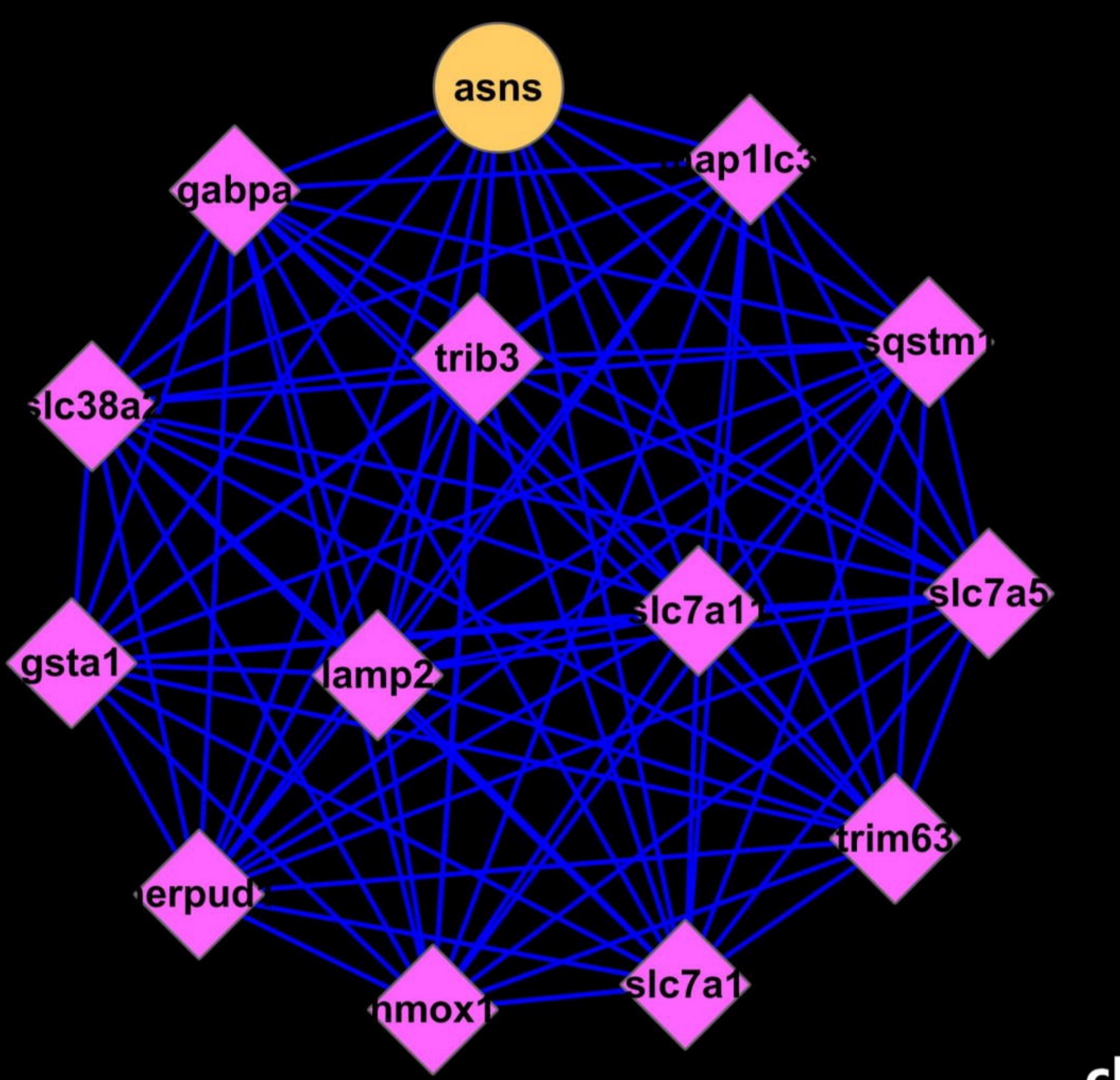




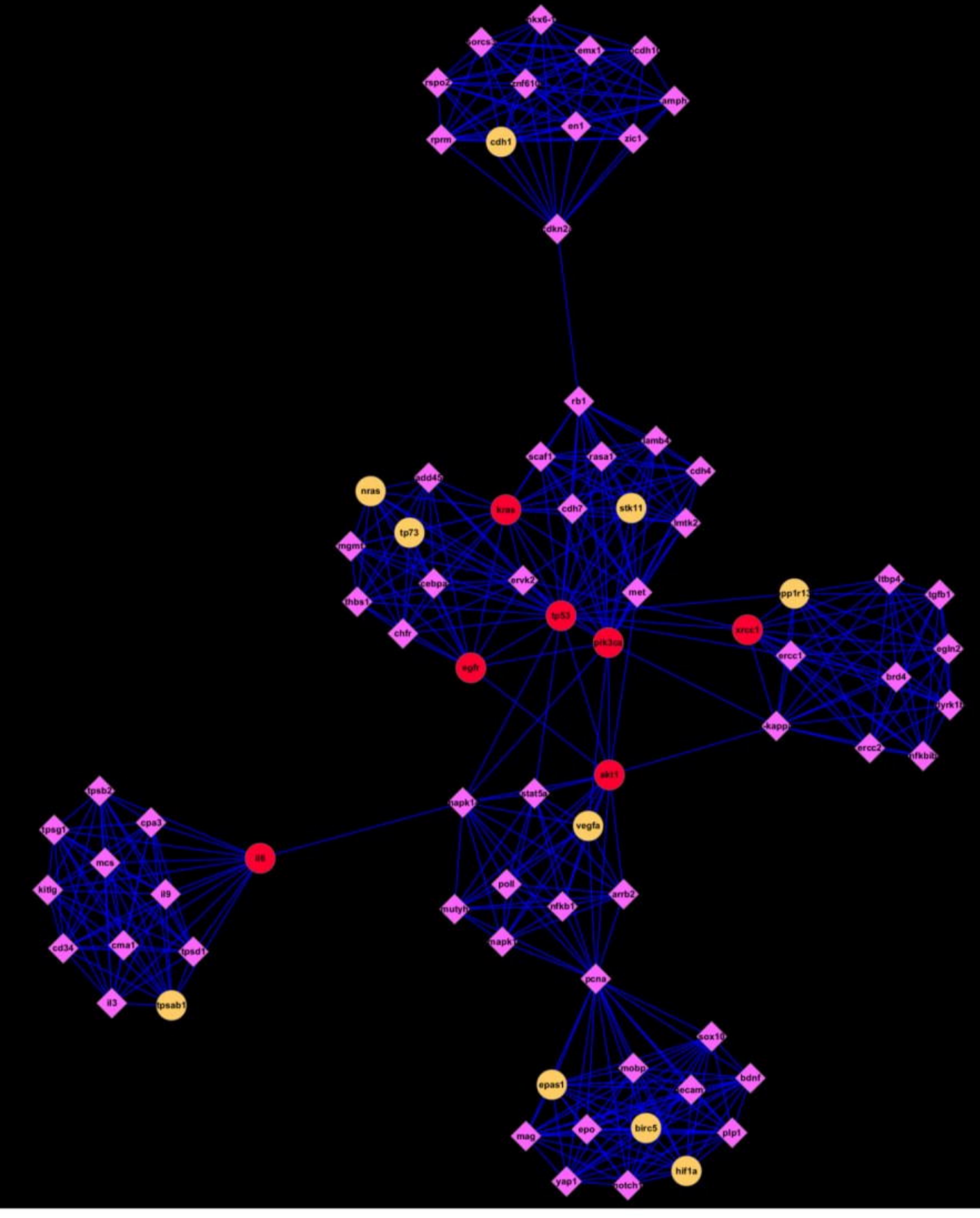
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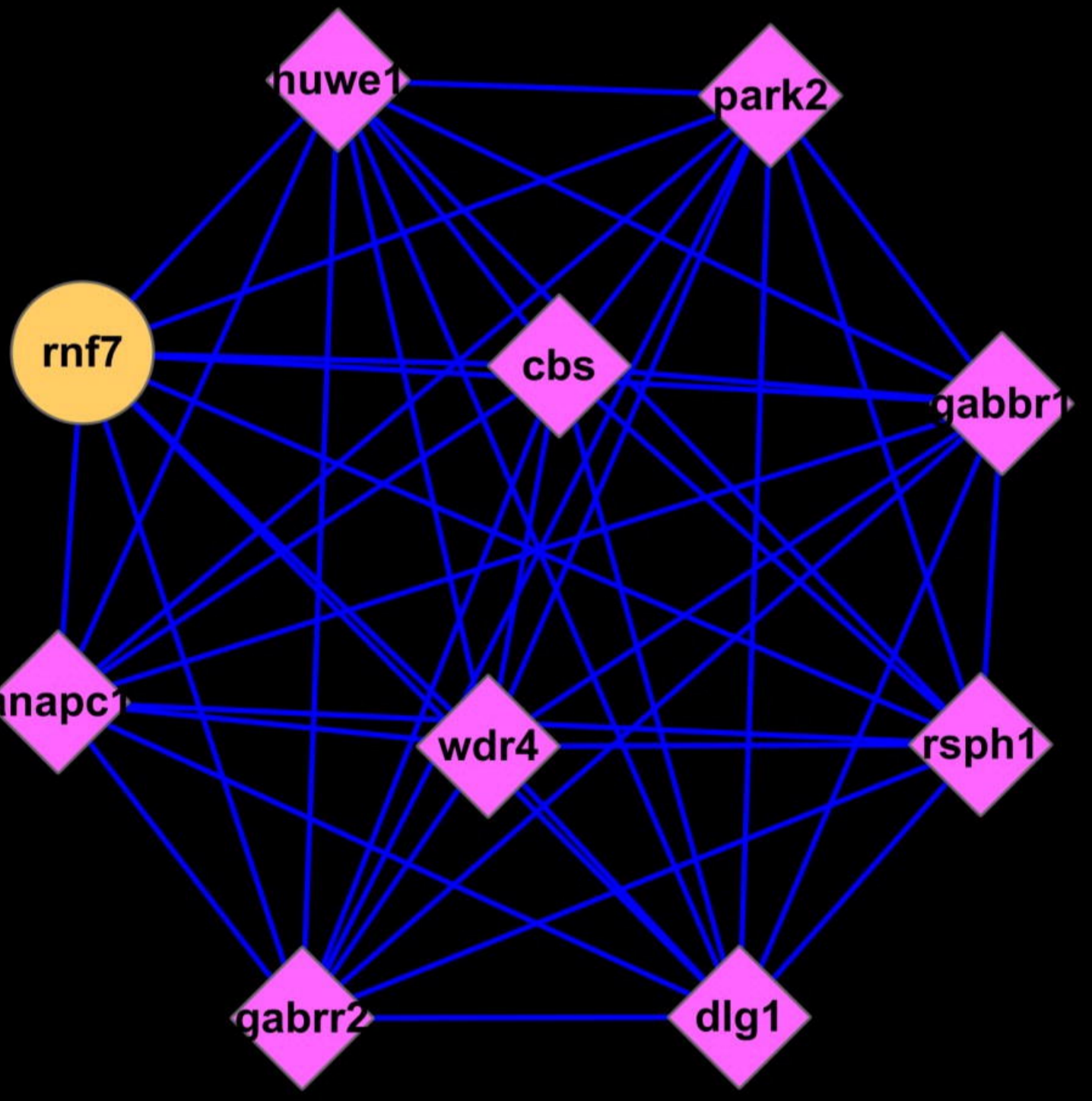
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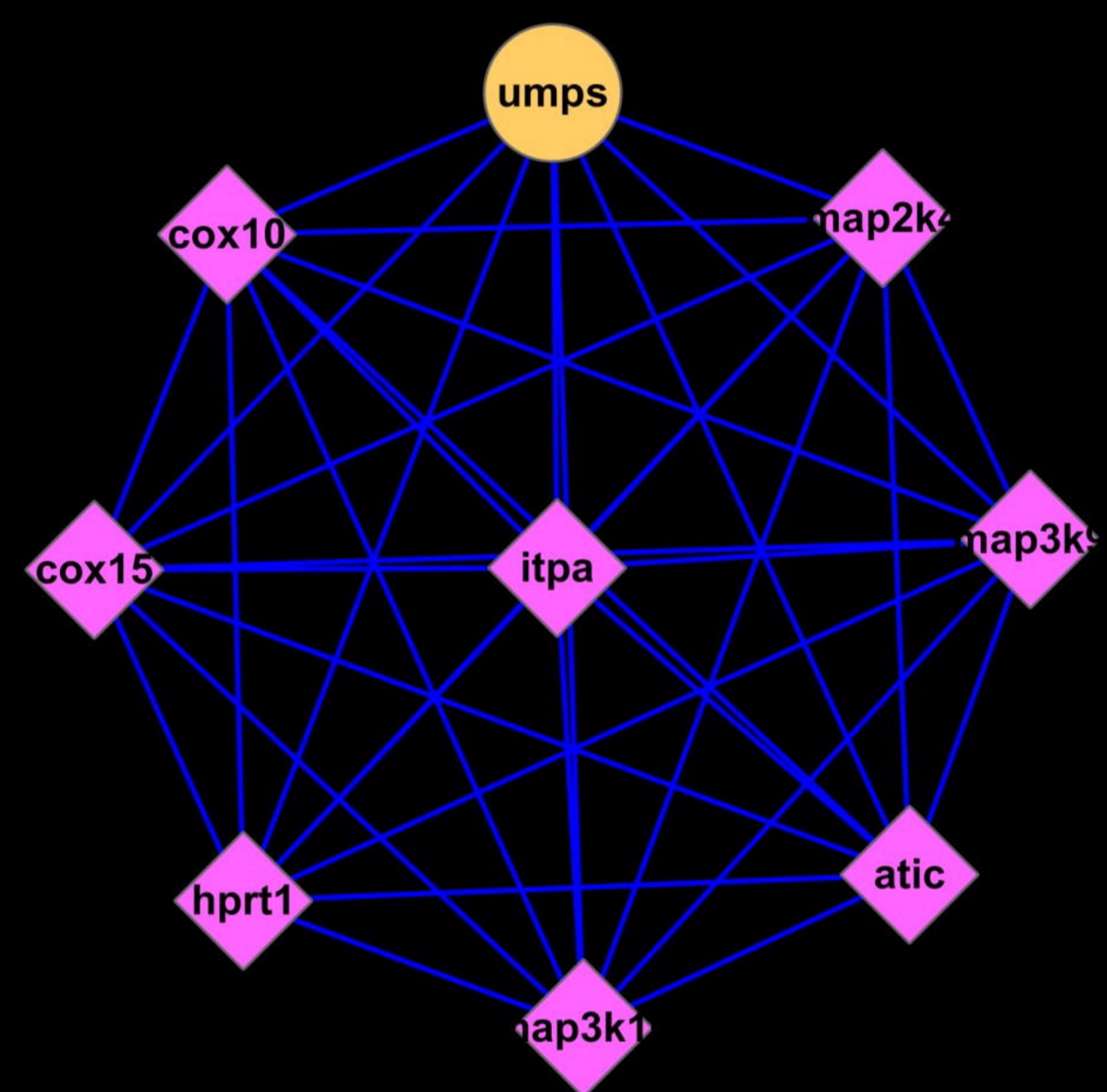
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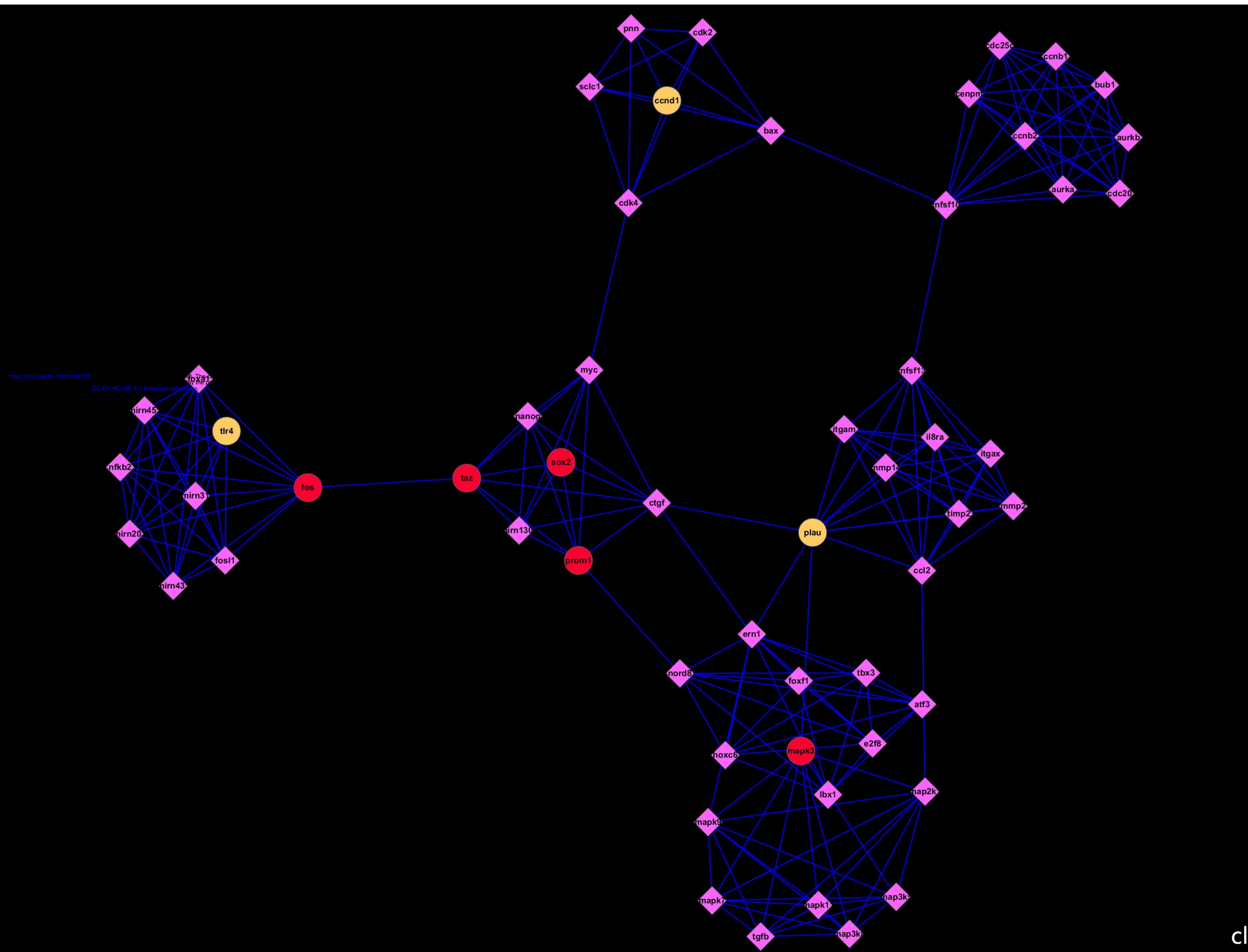
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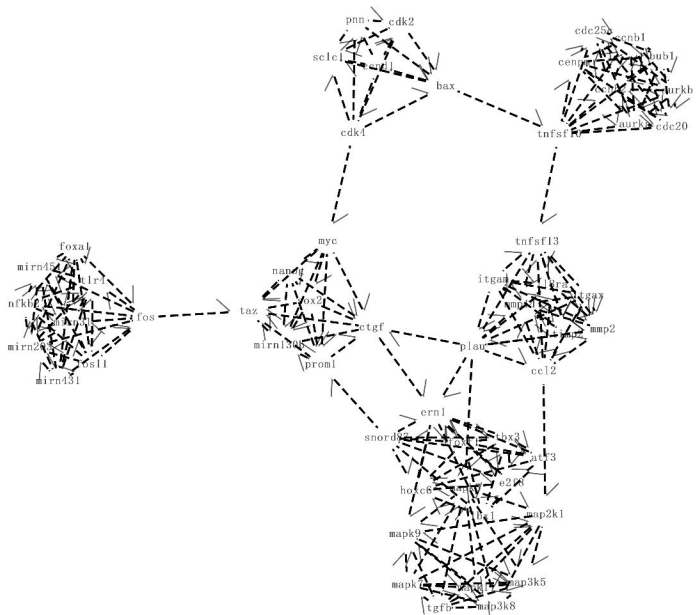


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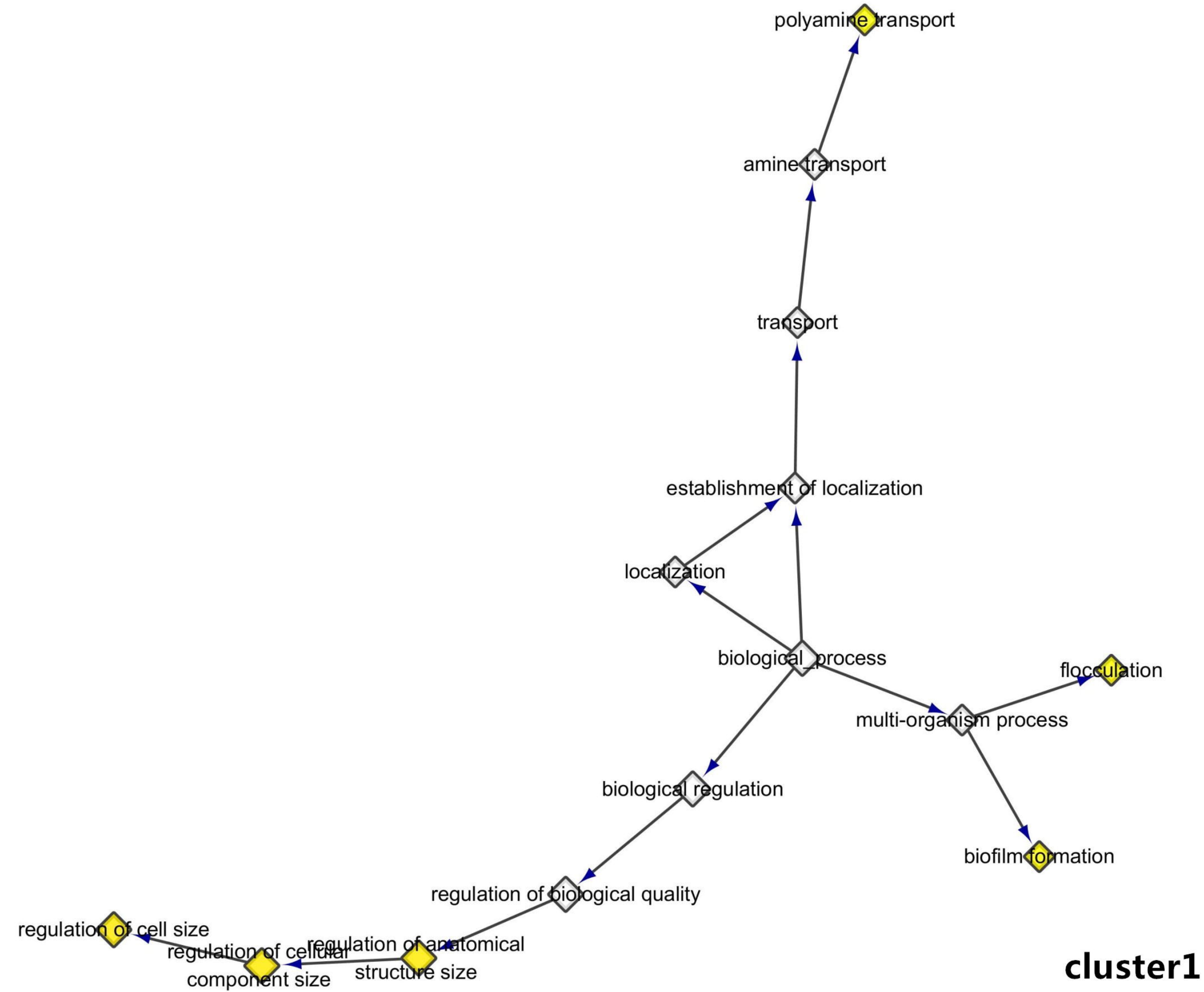


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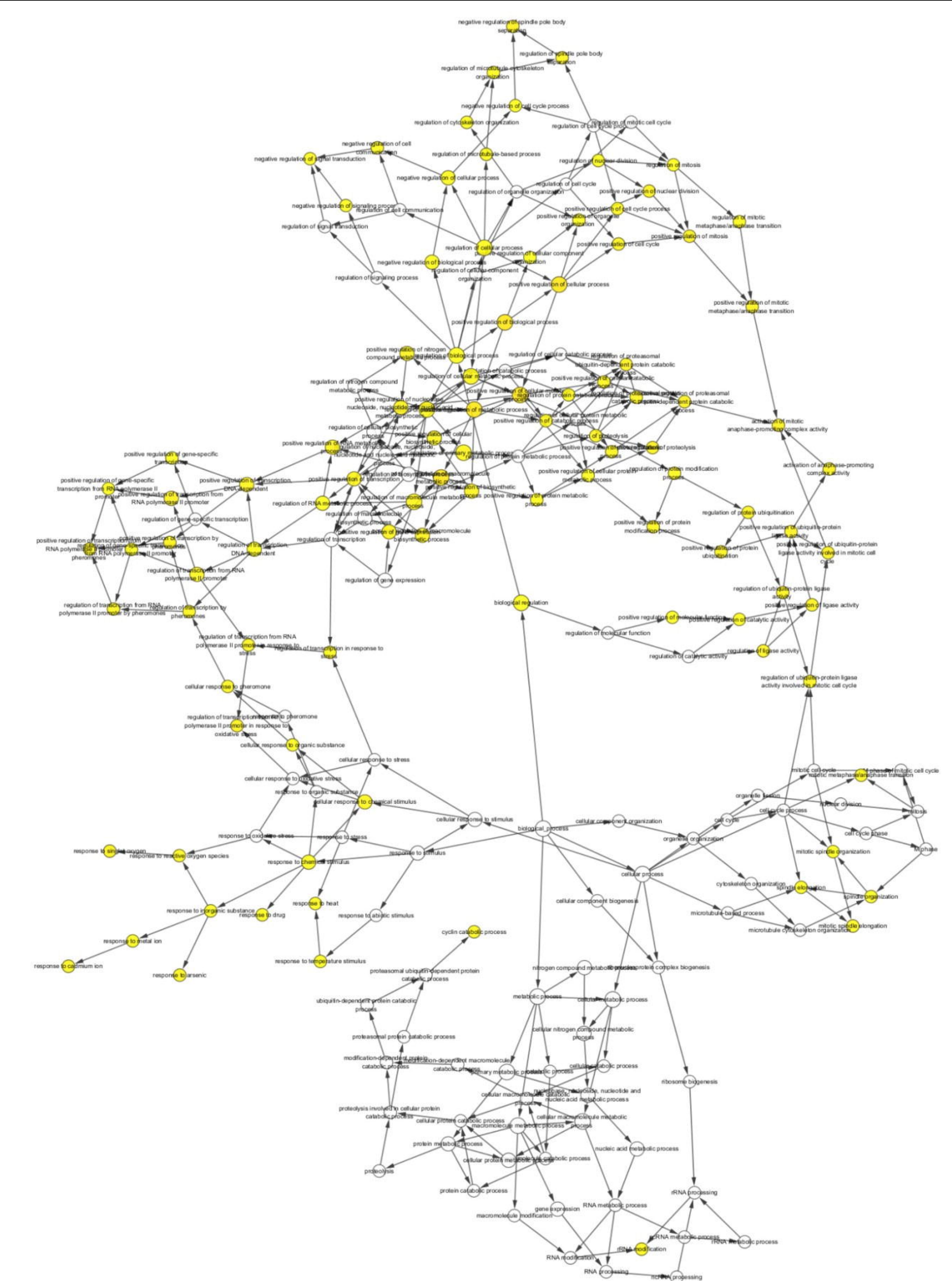






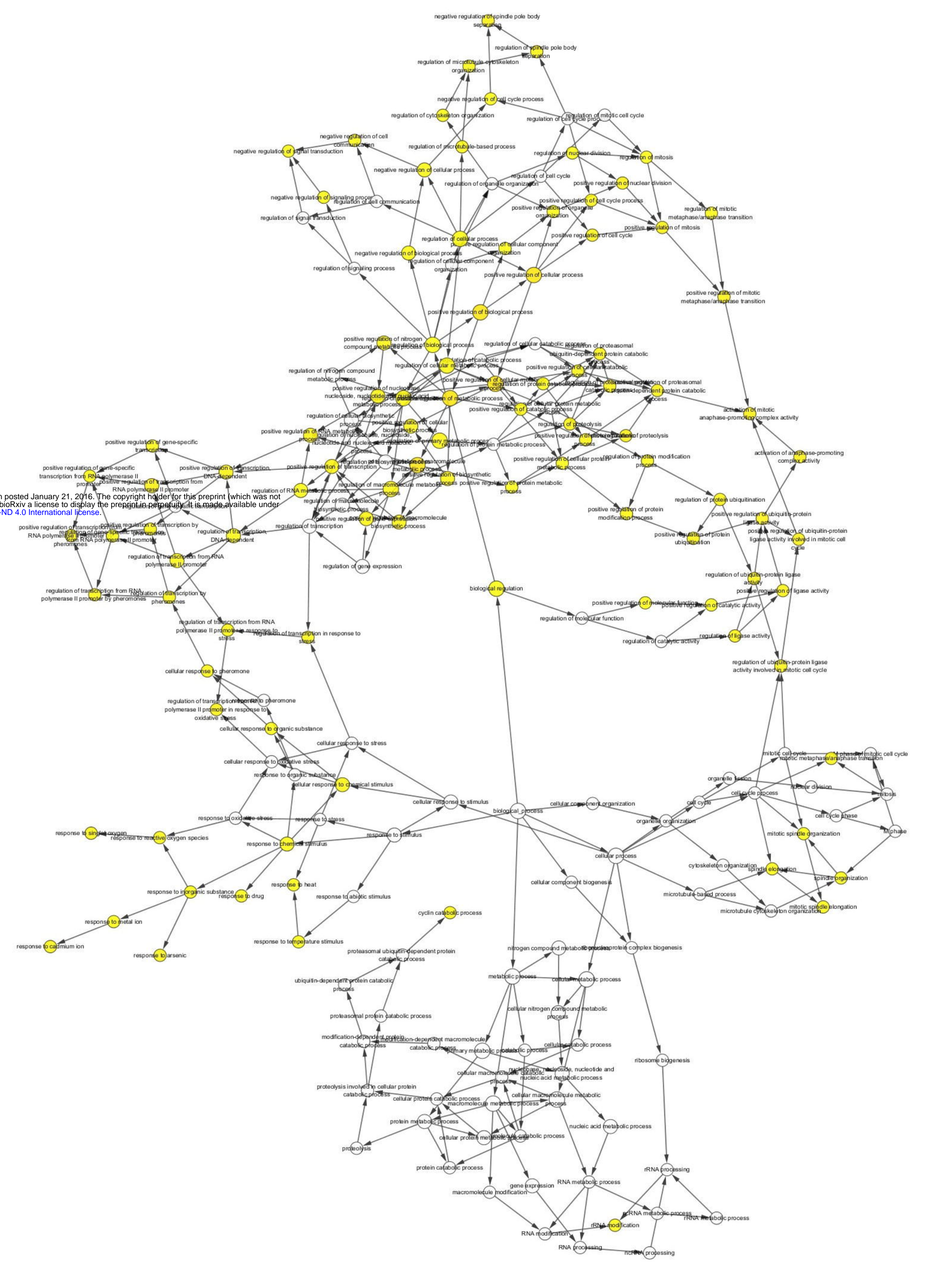


cluster1

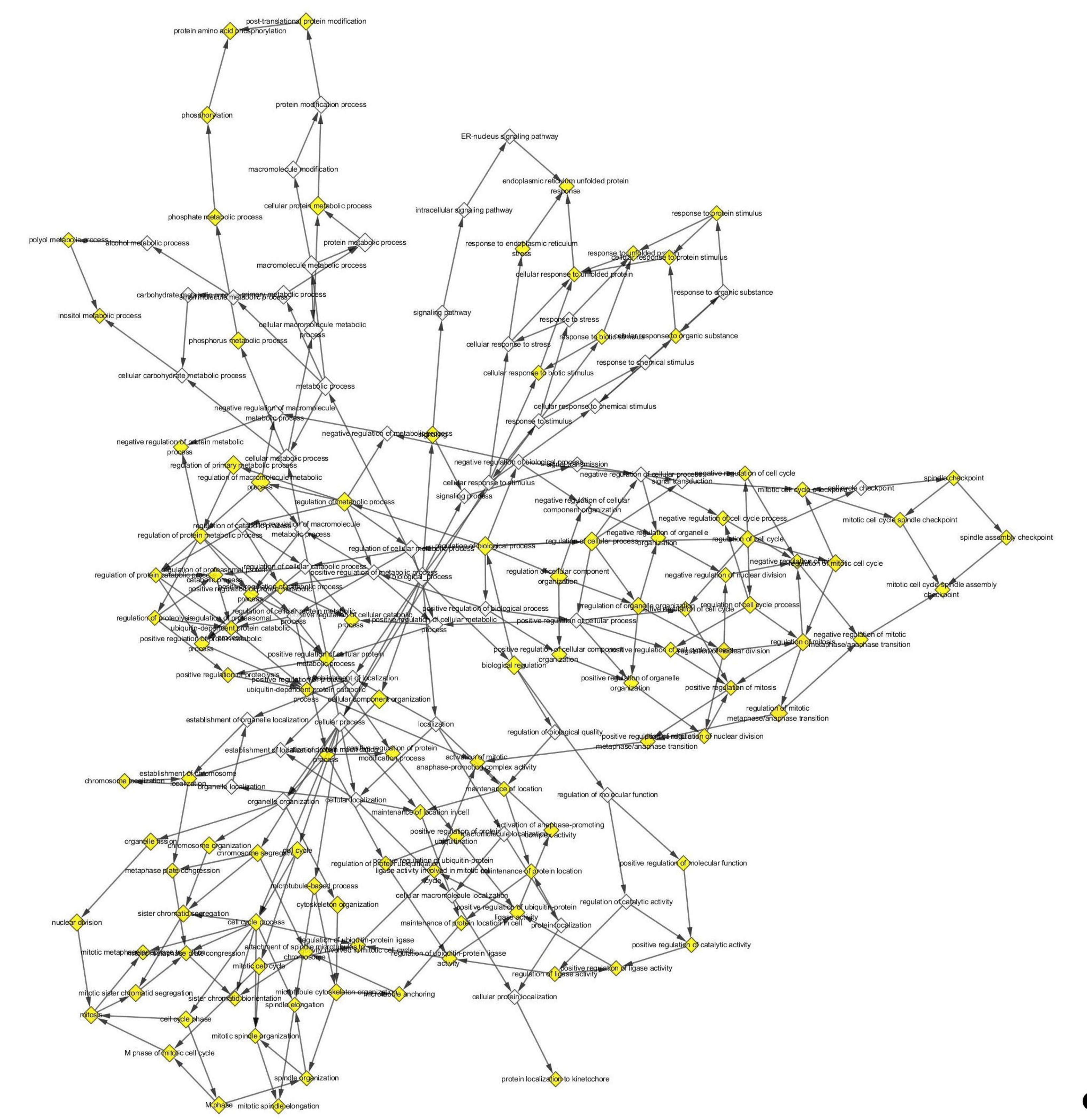


cluster4

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cluster6



cluster7



Symbol	Aliases
KRAS	C-K-RAS, CFC2, K-RAS2A, K-RAS2B, K-RAS4A, K-RAS4B, KI-RAS1, KRAS2, NS, NS3,
HIF1A	HIF-1A, HIF-1alpha, HIF1, HIF1-ALPHA, MOP1, PASD8, bHLHe78
TP53	BCC7, LFS1, P53, TRP53
TYMS	HST422, TMS, TS
EGFR	ERBB, ERBB1, HER1, NISBD2, PIG61, mENA
VEGFA	MVCD1, VEGF, VPF
CXCL12	IRH, PBSF, SCYB12, SDF1, TLSF, TPAR1
SLC2A1	CSE, DYT17, DYT18, DYT9, EIG12, GLUT, GLUT-1, GLUT1, GLUT1DS, HTLVR
PROM1	AC133, CD133, CORD12, MCDR2, MSTP061, PROML1, RP41, STGD4
ALDH1A1	ALDC, ALDH-E1, ALDH1, ALDH11, HEL-9, HEL-S-53e, HEL12, PUMB1, RALI
MTHFR	
ERBB2	CD340, HER-2, HER-2/neu, HER2, MLN 19, NEU, NGL, TKR1
CD44	CDW44, CSPG8, ECMR-III, HCELL, HUTCH-I, IN, LHR, MC56, MDU2, MDU3, MIC
EZR	CVIL, CVL, HEL-S-105, VIL2
MMP7	MMP-7, MPPL1, PUMP-1
CEACAM7	CGM2
PEBP4	CORK-1, CORK1, GWTM1933, HEL-S-300, PEBP-4, PRO4408, hPEBP4
PPARG	CIMT1, GLM1, NR1C31, PPARG2, PPARGgamma, PPARG
CTNNB1	CTNNB, MRD19, armadillo
SERPINE1	PAI, PAI-1, PAI1, PLANH1

BIRC5	API4, EPR-1
XRCC1	RCC
PIK3CA	CLOVE, CWS5, MCAP, MCM, MCMTc, PI3K, p110-alpha
HRAS	C-BAS/HAS, C-H-RAS, C-HA-RAS1, CTLO, H-RASIDX, HAMSv1, RASH1, p21ras
CYP2E1	CPE1, CYP2E, P450-J, P450C2E
EZH2	ENX-1, ENX1, EZH1b, KMT6, KMT6A, WVS, WVS2, EZH2
MKI67	KIA, MIB-, MIB-1, PPP1R105
CA9	CAIX, MN
CEACAM1	BGP, BGP1, BGPI
DPYD	DHP, DHPDHASE, DPD
TGFA	TFGA
CEACAM5	CD66e, CEA
PDPK1	PDK1, PDPK2, PDPK2P, PRO0461
LGR5	FEX, GPR49, GPR67, GRP49, HG38
ETV4	E1A-F, E1AF, PEA3, PEAS3
APOE	AD2, APO-E, LDLCQ5, LPG
IL6	BSF2, HGF, HSF, IFNB2, IL-6
TGFB1	CED, DPD1, LAP, TGFB, TGFBeta
PTGS2	COX-2, COX2, GRIPGHS, PGG/HS, PGHS-2, PHS-2, hCox-2
CRP	PTX1
IL1B	IL-1, IL1-BETA, IL1F2

AKT1	AKT, CWS6, PKB, PKB-ALPHA, PRKBA, RAC, RAC-ALPHA
VDR	NR1I1, PPP1R163
TLR4	ARMD10, CD284, TLR-4, TOLL
GSTM1	GST1-1, GSTM1a-1a, GSTM1b-1b, GTH4, GTM1, H-B, MU, MU-1, GSTM1
PTEN	10q23del, BZS, CWS1, DEC, GLM2, MHAM, MMAC11, TEP1, PTEN
CDH1	Arc-1, CD324, CDHE, ECAD, LCAM, UVO
TERT	CMM9, DKCA2, DKCB4, EST2, PFBMFT1, TCS1, TP2, TRT, hEST2, hTRT
BCL2	Bcl-2, PPP1R50
CXCR4	CD184, D2S201E, FB22, HM89, HSY3RR, LAP-3, LAP3, LCR1, LESTR, NPY3R, NPYR, NPYF WHIMS
CCND1	BCL1, D11S287E, PRAD1, U21B31
HLA-DQB1	CELIAC1, HLA-DQB, IDDM1
RELA	NFKB3, p65
CASP3	CPP32, CPP32B, SCA-1
HLA-G	MHC-G
MAPK3	ERK-1, ERK1, ERT2, HS44KDAP, HUMKER1A, P44ERK1, P44MAPK, PRKM3, p44-ERK
MMP1	CLG, CLGN
APC	BTPS2, DP2, DP2.5, DP3, GS, PPP1R46
CYP19A1	ARO, ARO1, CPV1, CYAR, CYP19, CYPXIX, P-450AROM
ERCC2	COFS2, EM9, TFIIH, TTD, TTD1, XPD
TCF7L2	TCF-4, TCF4

F3	CD142, TF, TFA
TP73	P73
LCN2	24p3, MSFI, NGAL
RUNX1	AML1, AML1-EVI-1, AMLCR1, CBF2alpha, CBFA2, EVI-1, PEBP2aB, PEBP2a
FGFR2	BBDS, BEK, BFR-1, CD332, CEK3, CFD1, ECT1, JWS, K-SAM, KGFR, TK14, 1
CHEK2	CDS1, CHK2, HuCds1, LFS2, PP1425, RAD53, hCds1
PLAU	ATF, BDPLT5, QPD, UPA, URK, u-PA
TIMP1	CLGI, EPA, EPO, HCI, TIMP
RAD51	BRCC5, FANCR, HRAD51, HsRad51, HsT16930, MRMV2A, RECA, RAD51
PLK1	PLK, STPK13
POU5F1	OCT3, OCT4, OTF-3, OTF3, OTF4, Oct-3, Oct-4
XRCC3	CMM6
SOX2	ANOP3, MCOPS3
PDGFRB	CD140B, IBGC4, IMF1, JTK12, PDGFR, PDGFR-1, PDGFR1
STK11	LKB1, PJS, hLKB1
DKK1	DKK-1, SK
EPAS1	ECYT4, HIF2A, HLF, MOP2, PASD2, bHLHe73
CFLAR	CASH, CASP8AP1, CLARP, Casper, FLAME, FLAME-1, FLAME1, FLIP, I-FLICE, MRIT, c-FLIF c-FLIPS
FOS	AP-1, C-FOS, p55
SKP2	FBL1, FBXL1, FLB1, p45



NRAS	ALPS4, CMNS, N-ras, NCMS1, NS6, NRAS
CCNE1	CCNE, pCCNE1
IL23R	
PPARD	FAAR, NR1C2, NUC1, NUCI, NUCII, PPARB
RPS6KB1	PS6K, S6K, S6K-beta-1, S6K1, STK14A, p70 S6KA, p70(S6K)-alpha, p70-S6K, p7
ANXA1	ANX1, LPC1
PTGS1	COX1, COX3, PCOX1, PES-1, PGG/HS, PGHS-1, PGHS1, PHS1, PTGHS
BLM	BS, RECQ2, RECQL2, RECQL3
CD163	M130, MM130
SPINK1	PCTT, PSTI, Spink3, TATI, TCP
CLDN1	CLD1, ILVASC, SEMP1
PDCD4	H731
IFNGR1	CD119, IFNGR, IMD27A, IMD27B
IL24	C49A, FISP, IL10B, MDA7, MOB5, ST16
ABCC4	MOAT-B, MOATB, MRP4
ALOX12	12-LOX, 12S-LOX, LOG12
BIRC7	KIAP, LIVIN, ML-IAP, MLIAP, RNF50
SATB1	
TFF3	ITF, P1B, TFI
CSF3R	CD114, GCSFR
RBL2	P130, Rb2

RPS6KA1	HU-1, MAPKAPK1A, RSK, RSK1, p90Rsk
FIGF	VEGF-D, VEGFD
TPSAB1	TPS1, TPS2, TPSB1
GSK3A	
ABCC3	ABC31, EST90757, MLP2, MOAT-D, MRP3, cMOAT2
MIR137	MIRN137, miR-137
TAZ	BTHS, CMD3A, EFE, EFE2, G4.5, LVNCX, Taz1
UGT2B15	HLUG4, UDPGT 2B8, UDPGT2B15, UDPGTH3, UGT2B8
PPP1R13L	IASPP, NKIP1, RAI, RAI4
REG4	GISP, REG-IV, RELP
WISP1	CCN4c, WISP1i, WISP1tc, WISP1
ASNS	ASNSD, TS11
UMPS	OPRT
F2RL2	PAR-3, PAR3
LIMS1	PINCH, PINCH-1, PINCH1
MUC6	MUC-6
RSF1	HBXAP, RSF-1, XAP8, p325
MYCL	L-Myc, LMYC1, bHLHe38, MYCL
RPS6KB2	KLS, P70-beta, P70-beta-1, P70-beta-2, S6K-beta2, S6K2, SRK, STK14B, p70(S6K)-be
RNF7	CKBBP1, ROC2, SAG
RPS6KA2	HU-2, MAPKAPK1C, RSK, RSK3, S6K-alpha, S6K-alpha2, p90-RSK3, pp90Rsk

SPARCL1

MAST 9, MAST9, FIG33, SC1

SPEN

HIAA0929, MINT, RBM15C, SHARP

SEMA4C

M-SEMA-F, SEMACL1, SEMAF, SEMAI

DNAJC12

JDP1

Category	Term	PValue	Genes	FDR
cluster1				
KEGG_PATHWAY	hsa04360:Axon guidance	0.006073822	SEMA6A, PTK2, SEMA3B	3.843791544
EC_NUMBER	3.4.21.-	0.009036531	HTRA1, KLK11, TMPRSS4	4.848129606
PANTHER_PATHWAY	P00004:Alzheimer disease-presenilin pathway	0.019326937	BACE2, MMP16, KAT5	11.44756514
cluster2				
KEGG_PATHWAY	hsa04630:Jak-STAT signaling pathway	3.32E-22	IL23R, IL6ST, SOCS1, STAT1, TYK2, IL12RB2, STAT4, IL23A, IL12RB1, IFNG, IL12A, JAK2, IL12B, IFNGR2, IFNGR1	2.53E-19
BIOCARTA	h_no2il12Pathway:N O2-dependent IL 12 Pathway in NK cells	1.28E-10	IL12RB2, TYK2, STAT4, IL12RB1, IFNG, IL12A, JAK2	1.10E-07
BIOCARTA	h_th1th2Pathway:Th1/Th2 Differentiation	2.81E-09	IL12RB2, IL12RB1, IFNG, IL12A, IL12B, IFNGR2, IFNGR1	2.42E-06
BIOCARTA	h_IL12Pathway:IL12 and Stat4 Dependent Signaling Pathway in Th1 Development	2.81E-09	IL12RB2, TYK2, STAT4, IL12RB1, IFNG, IL12A, JAK2	2.42E-06
KEGG_PATHWAY	hsa04060:Cytokine-cytokine receptor interaction	3.57E-09	IL12RB2, IL23A, IL12RB1, IL23R, IL6ST, IFNG, IL12A, IL12B, IFNGR2, IFNGR1	2.72E-06
BIOCARTA	h_ifngPathway:IFN gamma signaling pathway	4.16E-08	IFNG, JAK2, STAT1, IFNGR2, IFNGR1	3.57E-05
BIOCARTA	h_nktPathway:Selective expression of chemokine receptors during T-cell polarization	7.64E-07	IL12RB2, IL12RB1, IFNG, IL12A, IFNGR2, IFNGR1	6.57E-04
PANTHER_PATHWAY	P00035:Interferon-gamma signaling pathway	6.85E-06	IFNG, JAK2, STAT1, IFNGR2, IFNGR1	0.004406679
PANTHER_PATHWAY	P00038:JAK/STAT signaling pathway	8.34E-05	STAT4, SOCS1, JAK2, STAT1	0.053656257
BBID	12.IL-6_type_cytokine-signaling-transduct	8.71E-05	TYK2, IL6ST, SOCS1, JAK2, STAT1	0.07288153
PANTHER_PATHWAY	P00036:Interleukin signaling pathway	9.85E-05	IL12RB2, STAT4, IL23A, IL12RB1, IL6ST, IL12A, STAT1	0.063347144
BIOCARTA	h_tidPathway:Chaper	2.84E-04	IFNG, JAK2, IFNGR2, IFNGR1	0.243475935

	ones modulate			
	interferon Signaling			
	Pathway			
BBID	48.mice_minus_JAKs _and_STATs	7.11E-04	TYK2, STAT4, JAK2, STAT1	0.593475439
	P00031:Inflammation			
PANTHER_ PATHWAY	mediated by chemokine and cytokine signaling pathway	7.32E-04	TYK2, STAT4, IFNG, JAK2, STAT1, IFNGR2, IFNGR1	0.470260756
BBID	75.Stats_activators_o f_Apoptosis	0.00193497	IFNG, JAK2, STAT1	1.607237902
KEGG_PAT HWAY	hsa05330:Allograft rejection	0.004204138	IFNG, IL12A, IL12B	3.155617774
KEGG_PAT HWAY	hsa04940:Type I diabetes mellitus	0.005691714	IFNG, IL12A, IL12B	4.251279397
EC_NUMBE R	2.7.10.2	0.009766127	TYK2, JAK2	0.976612696
cluster3				
REACTOME _PATHWAY	REACT_15518:Trans membrane transport of small molecules	1.40E-04	SLC38A2, SLC7A1, SLC7A5, SLC7A11	0.07222524
REACTOME _PATHWAY	REACT_13:Metabolis m of amino acids	5.32E-04	SLC38A2, SLC7A1, ASNS, SLC7A5, SLC7A11	0.274131661
cluster4				
KEGG_PAT HWAY	hsa05200:Pathways in cancer	8.49E-14	EGFR, CEBPA, IL6, EPAS1, STAT5A, MET, TP53, KITLG, CDH1, BIRC5, NFKB1, RB1, TGFB1, AKT1, MAPK1, NRAS, LAMB4, CDKN2A, HIF1A, KRAS, VEGFA, PIK3CA	9.07E-11
KEGG_PAT HWAY	hsa05219:Bladder cancer	6.35E-11	EGFR, NRAS, MAPK1, CDKN2A, KRAS, VEGFA, TP53, CDH1, RB1, THBS1	6.79E-08
KEGG_PAT HWAY	hsa05218:Melanoma	4.01E-10	EGFR, AKT1, NRAS, MAPK1, CDKN2A, KRAS, MET, TP53, PIK3CA, CDH1, RB1	4.29E-07
KEGG_PAT HWAY	hsa05212:Pancreatic cancer	4.63E-10	EGFR, AKT1, MAPK1, CDKN2A, KRAS, VEGFA, TP53, PIK3CA, NFKB1, RB1, TGFB1	4.94E-07
KEGG_PAT HWAY	hsa05220:Chronic myeloid leukemia	7.02E-10	AKT1, NRAS, MAPK1, CDKN2A, KRAS, STAT5A, TP53, PIK3CA, NFKB1, RB1, TGFB1	7.50E-07
KEGG_PAT HWAY	hsa05211:Renal cell carcinoma	7.70E-09	AKT1, NRAS, MAPK1, HIF1A, KRAS, EPAS1, MET, VEGFA, PIK3CA, TGFB1	8.23E-06
KEGG_PAT HWAY	hsa05223:Non-small cell lung cancer	1.81E-08	EGFR, AKT1, NRAS, MAPK1, CDKN2A, KRAS, TP53, PIK3CA, RB1	1.93E-05

KEGG_PATHWAY	hsa05214:Glioma	6.34E-08	EGFR, AKT1, NRAS, MAPK1, CDKN2A, KRAS, TP53, PIK3CA, RB1	6.78E-05
PANTHER_PATHWAY	P04398:p53 pathway feedback loops 2	2.17E-07	AKT1, NRAS, CDKN2A, KRAS, MAPK14, TP53, PIK3CA, RB1, TP73	1.98E-04
KEGG_PATHWAY	hsa05213:Endometrial cancer	3.04E-07	EGFR, AKT1, NRAS, MAPK1, KRAS, TP53, PIK3CA, CDH1	3.25E-04
KEGG_PATHWAY	hsa05210:Colorectal cancer	6.20E-07	EGFR, AKT1, MAPK1, KRAS, MET, TP53, PIK3CA, BIRC5, TGFB1	6.62E-04
KEGG_PATHWAY	hsa05221:Acute myeloid leukemia	6.57E-07	CEBPA, AKT1, NRAS, MAPK1, KRAS, STAT5A, PIK3CA, NFKB1	7.01E-04
KEGG_PATHWAY	hsa05215:Prostate cancer	9.71E-07	EGFR, AKT1, NRAS, MAPK1, KRAS, TP53, PIK3CA, NFKB1, RB1	0.001037757
KEGG_PATHWAY	hsa04722:Neurotrophin signaling pathway	1.17E-06	AKT1, NRAS, MAPK1, BDNF, KRAS, MAPK14, TP53, PIK3CA, NFKB1, TP73	0.00125362
KEGG_PATHWAY	hsa04010:MAPK signaling pathway	2.69E-06	EGFR, AKT1, MAPK1, NRAS, BDNF, KRAS, ARRB2, MAPK14, GADD45G, TP53, NFKB1, TGFB1, RASA1	0.002871532
KEGG_PATHWAY	h_erythPathway:Erythrocyte Differentiation Pathway	3.22E-06	IL3, IL6, IL9, KITLG, TGFB1, EPO	0.003887971
REACTOME_PATHWAY	REACT_16888:Signaling by PDGF	9.59E-06	NRAS, MAPK1, KRAS, STAT5A, PIK3CA, THBS1, RASA1	0.007693883
PANTHER_PATHWAY	P00056:VEGF signaling pathway	1.68E-05	AKT1, NRAS, MAPK1, HIF1A, KRAS, MAPK14, VEGFA, PIK3CA	0.015328038
EC_NUMBER	3.4.21.59	4.75E-05	TPSAB1, TPSB2, TPSD1, TPSG1	0.030526793
KEGG_PATHWAY	hsa04370:VEGF signaling pathway	4.94E-05	AKT1, NRAS, MAPK1, KRAS, MAPK14, VEGFA, PIK3CA	0.052708911
REACTOME_PATHWAY	REACT_11061:Signaling by NGF	5.63E-05	AKT1, MAG, NRAS, MAPK1, KRAS, MAPK14, PIK3CA, NFKB1, SORCS3	0.045131702
KEGG_PATHWAY	hsa04664:Fc epsilon RI signaling pathway	6.17E-05	AKT1, NRAS, MAPK1, IL3, KRAS, MAPK14, PIK3CA	0.065876411
KEGG_PATHWAY	hsa04150:mTOR signaling pathway	9.18E-05	AKT1, MAPK1, HIF1A, STK11, VEGFA, PIK3CA	0.097993218
PANTHER_PATHWAY	P00005:Angiogenesis	9.21E-05	AKT1, NRAS, MAPK1, NOTCH1, HIF1A, KRAS, MAPK14, VEGFA, PIK3CA, BIRC5, RASA1	0.084242254
PANTHER_PATHWAY	P00018:EGF receptor signaling pathway	1.11E-04	EGFR, AKT1, NRAS, MAPK1, KRAS, MAPK14, STAT5A, PIK3CA, RASA1	0.101309728
KEGG_PATHWAY	hsa04012:ErbB	1.14E-04	EGFR, AKT1, NRAS, MAPK1, KRAS,	0.121813791

HWAY	signaling pathway		STAT5A, PIK3CA	
KEGG_PAT	hsa05216:Thyroid	1.18E-04	NRAS, MAPK1, KRAS, TP53, CDH1	0.126019896
HWAY	cancer			
	h_hcmvPathway:Human Cytomegalovirus and Map Kinase Pathways			
BIOCARTA		1.34E-04	AKT1, MAPK1, MAPK14, NFKB1, RB1	0.161807042
KEGG_PAT	hsa04115:p53	3.30E-04	CDKN2A, GADD45G, RPRM, TP53, THBS1, TP73	0.352478227
HWAY	signaling pathway			
KEGG_PAT	hsa04660:T cell receptor signaling	3.75E-04	AKT1, NRAS, MAPK1, KRAS, MAPK14, PIK3CA, NFKB1	0.399792892
HWAY	pathway			
	h_telPathway:Telomeres, Telomerase, Cellular Aging, and Immortality			
BIOCARTA		3.86E-04	EGFR, AKT1, KRAS, TP53, RB1	0.465899729
PANTHER_PATHWAY	P04393:Ras Pathway	3.87E-04	AKT1, NRAS, MAPK1, CDKN2A, KRAS, MAPK14, PIK3CA	0.353518988
REACTOME_PATHWAY	REACT_9417:Signaling by EGFR	5.16E-04	EGFR, NRAS, MAPK1, KRAS, PIK3CA	0.413199584
KEGG_PAT	hsa04662:B cell receptor signaling	5.22E-04	AKT1, NRAS, MAPK1, KRAS, PIK3CA, NFKB1	0.556225736
HWAY	pathway			
KEGG_PAT	hsa05222:Small cell lung cancer	8.79E-04	AKT1, LAMB4, TP53, PIK3CA, NFKB1, RB1	0.934847379
HWAY	pathway			
	h_RacCycDPathway:Influence of Ras and Rho proteins on G1 to S Transition			
BIOCARTA		0.001228749	AKT1, MAPK1, PIK3CA, NFKB1, RB1	1.475202636
KEGG_PAT	hsa04320:Dorso-ventral axis formation	0.001386389	EGFR, MAPK1, NOTCH1, KRAS	1.471198696
HWAY	pathway			
REACTOME_PATHWAY	REACT_6900:Signaling in Immune system	0.001592713	MAG, NRAS, MAPK1, KRAS, CD34, PECAM1, PIK3CA, NFKB1, CDH1	1.270531117
PANTHER_PATHWAY	P00059:p53 pathway	0.001789034	AKT1, CDKN2A, GADD45G, TP53, PIK3CA, THBS1, TP73	1.624912242
KEGG_PAT	hsa04510:Focal adhesion	0.001957264	EGFR, AKT1, MAPK1, LAMB4, MET, VEGFA, PIK3CA, THBS1	2.071273278
HWAY	pathway			
KEGG_PAT	hsa04620:Toll-like receptor signaling	0.002011203	AKT1, MAPK1, IL6, MAPK14, PIK3CA, NFKB1	2.127799484
HWAY	pathway			
KEGG_PAT	hsa04060:Cytokine-cytokine receptor interaction	0.002108909	EGFR, IL3, IL6, MET, VEGFA, IL9, KITLG, TGFB1, EPO	2.230116541
HWAY	pathway			

PANTHER_PATHWAY	P00010:B cell activation	0.002214285	NRAS, MAPK1, KRAS, MAPK14, PIK3CA, NFKB1	2.007674836
REACTOME_PATHWAY	REACT_604:Hemostasis	0.002330368	MAG, NRAS, KRAS, VEGFA, PECAM1, PIK3CA, THBS1, TGFB1	1.854165541
KEGG_PATHWAY	hsa04630:Jak-STAT signaling pathway	0.002487246	AKT1, IL3, IL6, STAT5A, IL9, PIK3CA, EPO	2.62539532
PANTHER_PATHWAY	P00021:FGF signaling pathway	0.003144809	AKT1, NRAS, MAPK1, KRAS, MAPK14, PIK3CA, RASA1	2.840604013
BIOCARTA	h_il17Pathway:IL 17 Signaling Pathway	0.003157025	IL3, IL6, CD34, KITLG	3.750040719
BIOCARTA	h_stemPathway:Regulation of hematopoiesis by cytokines	0.003157025	IL3, IL6, IL9, EPO	3.750040719
PANTHER_PATHWAY	P04397:p53 pathway by glucose deprivation	0.003458487	AKT1, STK11, TP53, TP73	3.119958155
REACTOME_PATHWAY	REACT_498:Signaling by Insulin receptor	0.003756119	NRAS, MAPK1, KRAS, PIK3CA	2.973670175
KEGG_PATHWAY	hsa04110:Cell cycle	0.005068716	CDKN2A, GADD45G, PCNA, TP53, RB1, TGFB1	5.284031958
BIOCARTA	h_badPathway:Regulation of BAD phosphorylation	0.005426648	AKT1, MAPK1, IL3, KITLG	6.365799458
BIOCARTA	h_crebPathway:Transcription factor CREB and its extracellular signals	0.005426648	AKT1, MAPK1, MAPK14, PIK3CA	6.365799458
KEGG_PATHWAY	hsa04062:Chemokine signaling pathway	0.006278908	AKT1, NRAS, MAPK1, KRAS, ARR2, PIK3CA, NFKB1	6.507599178
BIOCARTA	h_keratinocytePathway:Keratinocyte Differentiation	0.006958557	EGFR, CEBPA, MAPK1, MAPK14, NFKB1	8.094303481
KEGG_PATHWAY	hsa04914:Progesterone-mediated oocyte maturation	0.007248802	AKT1, MAPK1, KRAS, MAPK14, PIK3CA	7.477850945
KEGG_PATHWAY	hsa04640:Hematopoietic cell lineage	0.007248802	IL3, IL6, CD34, KITLG, EPO	7.477850945
KEGG_PATHWAY	hsa04210:Apoptosis	0.007548465	AKT1, IL3, TP53, PIK3CA, NFKB1	7.7757735
PANTHER_PATHWAY	P00036:Interleukin signaling pathway	0.007614161	AKT1, NRAS, MAPK1, IL6, KRAS, STAT5A, PIK3CA, RASA1	6.753977015
REACTOME	REACT_216:DNA	0.009001	MGMT, PCNA, XRCC1, ERCC1, ERCC2	6.996374809



_PATHWAY Repair				
PANTHER_	P00053:T cell	0.009738791	AKT1, NRAS, MAPK1, KRAS, PIK3CA,	8.564577054
PATHWAY	activation		NFKB1	
cluster5				
KEGG_PAT	hsa04120:Ubiquitin	6.19E-04	ANAPC1, RNF7, HUWE1, PARK2	0.370875716
HWAY	mediated proteolysis			
EC_NUMBE	6.3.2.-	0.069187942	HUWE1, PARK2	14.94398228
R				
cluster6				
REACTOME	REACT_1698:Metabli	5.07E-04	UMPS, ATIC, HPRT1	NaN
_PATHWAY	sm of nucleotides			
KEGG_PAT	hsa00983:Drug	0.001025542	ITPA, UMPS, HPRT1	0.657964401
HWAY	metabolism			
	h_mapkPathway:MA			
BIOCARTA	PKinase Signaling	0.00306271	MAP3K9, MAP2K4, MAP3K10	2.110065352
	Pathway			
PANTHER_	P05918:p38 MAPK	0.005195037	MAP3K9, MAP2K4, MAP3K10	3.701051163
PATHWAY	pathway			
KEGG_PAT	hsa00230:Purine	0.012459659	ITPA, ATIC, HPRT1	7.749728301
HWAY	metabolism			
EC_NUMBE	2.7.11.25	0.030208436	MAP3K9, MAP3K10	16.18260742
R				
KEGG_PAT	hsa00860:Porphyrin	0.038330298	COX10, COX15	22.23317069
HWAY	and chlorophyll			
	metabolism			
PANTHER_	P00029:Huntington	0.04121774	MAP3K9, MAP2K4, MAP3K10	26.27015356
PATHWAY	disease			
	h_p38mapkPathway:			
BIOCARTA	p38 MAPK Signaling	0.045417275	MAP3K9, MAP2K4	27.61443797
	Pathway			
BBID	100.MAPK_signaling	0.058659218	MAP2K4, MAP3K10	17.32821753
	_cascades			
cluster7				
KEGG_PAT	hsa04914:Progesterone-mediated oocyte	2.08E-08	CCNB1, CCNB2, MAP2K1, MAPK3, BUB1,	2.13E-05
HWAY	maturation		MAPK9, MAPK11, CDC25C, CDK2	
KEGG_PAT	hsa04114:Oocyte	1.46E-07	CCNB1, CCNB2, MAP2K1, MAPK3, BUB1,	1.50E-04
HWAY	meiosis		CDC20, AURKA, CDC25C, CDK2	
KEGG_PAT	hsa04110:Cell cycle	3.96E-07	CCNB1, CCND1, CCNB2, BUB1, CDC20,	4.05E-04
HWAY			CDK4, CDC25C, MYC, CDK2	
KEGG_PAT	hsa05219:Bladder	3.68E-06	CCND1, MAP2K1, MAPK3, CDK4, MYC,	0.00376453
HWAY	cancer		MMP2	

KEGG_PATHWAY	hsa05210:Colorectal cancer	7.34E-06	FOS, CCND1, MAP2K1, BAX, MAPK3, MAPK9, MYC	0.007502731
KEGG_PATHWAY	hsa05200:Pathways in cancer	1.02E-05	FOS, CCND1, MAP2K1, BAX, MAPK3, MAPK9, NFKB2, CDK4, MYC, MMP2, CDK2	0.010452597
KEGG_PATHWAY	hsa04010:MAPK signaling pathway	1.43E-05	FOS, MAP3K5, MAP2K1, MAPK3, MAP3K8, MAPK9, MAPK11, NFKB2, MAPK7, MYC	0.014636534
REACTOME_PATHWAY	REACT_152:Cell Cycle, Mitotic	1.79E-05	CCNB1, CENPM, CCND1, CCNB2, BUB1, CDC20, AURKA, AURKB, CDK4, CDC25C, CDK2	0.012929573
KEGG_PATHWAY	hsa04912:GnRH signaling pathway	1.79E-05	MAP2K1, MAPK3, MAPK9, MAPK11, MAPK7, MMP14, MMP2	0.018347497
EC_NUMBER	2.7.11.24	2.02E-05	MAPK3, MAPK9, MAPK11, MAPK7	0.012135676
KEGG_PATHWAY	hsa04620:Toll-like receptor signaling pathway	2.13E-05	FOS, MAP2K1, MAPK3, MAP3K8, MAPK9, MAPK11, TLR4	0.02182043
PANTHER_PATHWAY	P00006:Apoptosis signaling pathway	2.68E-05	FOS, TNFSF10, MAP3K5, ATF3, BAX, MAPK3, MAPK9, NFKB2, MAPK7	0.02355655
KEGG_PATHWAY	hsa04660:T cell receptor signaling pathway	3.13E-05	FOS, MAP2K1, MAPK3, MAP3K8, MAPK9, MAPK11, CDK4	0.032014891
PANTHER_PATHWAY	P00054:Toll receptor signaling pathway	3.16E-05	MAP2K1, MAPK3, MAP3K8, MAPK9, MAPK11, TLR4, NFKB2	0.02781923
KEGG_PATHWAY	hsa04115:p53 signaling pathway	4.05E-05	CCNB1, CCND1, CCNB2, BAX, CDK4, CDK2	0.041438576
KEGG_PATHWAY	hsa04722:Neurotrophin signaling pathway	6.84E-05	MAP3K5, MAP2K1, BAX, MAPK3, MAPK9, MAPK11, MAPK7	0.069959614
BIOCARTA_PATHWAY	h_mapkPathway:MAPK Kinase Signaling Pathway	3.55E-04	FOS, MAP3K5, MAP2K1, MAPK3, MAP3K8, MAPK9, MAPK11, MAPK7	0.400316292
KEGG_PATHWAY	hsa05216:Thyroid cancer	6.11E-04	CCND1, MAP2K1, MAPK3, MYC	0.622621821
PANTHER_PATHWAY	P00052:TGF-beta signaling pathway	6.33E-04	FOS, FOXF1, FOXA1, MAPK3, MAPK9, MAPK11, MAPK7, FOSL1	0.555596842
KEGG_PATHWAY	hsa05212:Pancreatic cancer	7.67E-04	CCND1, MAP2K1, MAPK3, MAPK9, CDK4	0.781138732
KEGG_PATHWAY	hsa05220:Chronic myeloid leukemia	8.95E-04	CCND1, MAP2K1, MAPK3, CDK4, MYC	0.91105286
PANTHER_PATHWAY	P00010:B cell activation	0.001400729	FOS, MAP2K1, MAPK3, MAPK9, MAPK11, NFKB2	1.225108429
BIOCARTA_PATHWAY	h_p53Pathway:p53 Signaling Pathway	0.003024948	CCND1, BAX, CDK4, CDK2	3.364924295

KEGG_PAT	hsa05213:Endometri	0.003370738	CCND1, MAP2K1, MAPK3, MYC	3.393288113
HWAY	al cancer			
KEGG_PAT	hsa05223:Non-small	0.003753245	CCND1, MAP2K1, MAPK3, CDK4	3.77171584
HWAY	cell lung cancer			
PANTHER_	P00035:Interferon-ga			
PATHWAY	mma signaling	0.004182552	MAPK3, MAPK9, MAPK11, MAPK7	3.618778393
PATHWAY	pathway			
KEGG_PAT	hsa05221:Acute	0.004595499	CCND1, MAP2K1, MAPK3, MYC	4.600275355
HWAY	myeloid leukemia			
PANTHER_	P00034:Integrin		MAP3K5, MAP2K1, ITGAX, MAPK3, MAPK9,	
PATHWAY	signalling pathway	0.005122253	MAPK11, MAPK7, ITGAM	4.415663351
KEGG_PAT	hsa04621:NOD-like			
HWAY	receptor signaling	0.005544037	CCL2, MAPK3, MAPK9, MAPK11	5.525680097
HWAY	pathway			
KEGG_PAT	hsa05214:Glioma	0.005798209	CCND1, MAP2K1, MAPK3, CDK4	5.772273682
HWAY				
REACTOME	REACT_1538:Cell			
_PATHWAY	Cycle Checkpoints	0.007137442	CCNB1, CCNB2, CDC20, CDC25C, CDK2	5.054206969
KEGG_PAT	hsa05218:Melanoma	0.008084875	CCND1, MAP2K1, MAPK3, CDK4	7.964786431
HWAY				
	h_RacCycDPathway:			
BIOCARTA	Influence of Ras and	0.008275776	CCND1, MAPK3, CDK4, CDK2	8.961786045
	Rho proteins on G1			
	to S Transition			
	h_cellcyclePathway:			
BIOCARTA	Cyclins and Cell	0.009291865	CCNB1, CCND1, CDK4, CDK2	10.010086
	Cycle Regulation			