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A Motif-Based Network Analysis of Regulatory Patterns in Doxorubicin Effects on Treating Breast Cancer, a Systems Biology Study

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Abstract

Background: Breast cancer is the most common malignancy worldwide. Doxorubicin is an anthracycline used to treat breast cancer as the first treatment choice. Nevertheless, the molecular mechanisms underlying the response to Doxorubicin and its side effects are not comprehensively understood so far. We used systems biology and bio-informatics methods to identify essential genes and molecular mechanisms behind the body response to Doxorubicin and its side effects in breast cancer patients.

Methods: Omics data were extracted and analyzed to construct the protein-protein interaction and gene regulatory networks. Network analysis was performed to identify hubs, bottlenecks, clusters, and regulatory motifs to evaluate crucial genes and molecular mechanisms behind the body response to Doxorubicin and its side effects.

Results: Analyzing the constructed PPI and gene-TF-miRNA regulatory network showed that MCM3, MCM10, and TP53 are key hub-bottlenecks and seed proteins. Enrichment analysis also revealed cell cycle, TP53 signaling, Forkhead box O (FoxO) signaling, and viral carcinogenesis as essential pathways in response to this drug. Besides, SNARE interactions in vesicular transport and neurotrophin signaling were identified as pathways related to the side effects of Doxorubicin. The apoptosis in-duction, DNA repair, invasion inhibition, metastasis, and DNA replication are sug-gested as critical molecular mechanisms underlying Doxorubicin anti-cancer effect. SNARE interactions in vesicular transport and neurotrophin signaling and FoxO signaling pathways in glucose metabolism are probably the mechanisms responsible for side effects of Doxorubicin.

Conclusion: Following our model validation using the existing experimental data, we recommend our other newly predicted biomarkers and pathways as possible molecular mechanisms and side effects underlying the response to Doxorubicin in breast cancer requiring further investigations.

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Keyw ords: Breast cancer, Doxorubicin, Protein-protein interaction network, Regulatory motif, Systems biology

Introduction

Breast cancer is the most common cause of cancer and mortality caused by cancers in women worldwide ¹. Four subtypes of this cancer include luminal A and luminal B [expressing the Estrogen Receptor (ER)], basal-like, and Human Epidermal growth factor Receptor 2 (HER2)-enriched (without ER expression). This cancer is a heterogeneous disease at the molecular level. The characterization influence biologically-directed therapies and treatment de-escalation ². Breast cancer is often curable early, but the metastatic form is almost mortal due to therapeutic resistance ³. The estrogen hormone and its receptor play essential roles in breast cancer progression. The dysregulation of the Estrogen Receptor (ER) is attributed to two-thirds of all breast

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cancers. The ER receptor is one of the therapeutic targets for ER+ breast cancer ⁴. In clinical diagnosis, 75% of breast tumors are ER+ ⁵; however, the role of ER signaling in metastasis of breast cancer remains poorly understood. Several studies have shown an adverse effect of ER signaling on motility and invasion of cells ^{6,7}, while a few studies suggested a positive effect of ER signaling on motility and invasion ^{8,9}.

Doxorubicin (DXR) is an anthracycline and chemotherapeutic drug isolated from Streptomyces peucetius ¹⁰. This drug is used to treat several cancers, including breast, gastric, lung, ovarian, thyroid, sarcoma, non-Hodgkins and Hodgkins lymphoma, multiple myeloma, and pediatric cancers ^{11,12}. DXR induces Reactive Oxygen Species release (ROS) that ROS lead to DNA damage, lipid peroxidation and membrane damage, and apoptotic cell death pathways ¹³. DXR is among the chemotherapy drugs approved to treat ER+breast cancer. The response rates to DXR in patients exposed to DXR for the first time is reported to be 48%, and for more than once is 28%¹⁴. Nevertheless, little is known about the molecular basis of its effect on cell proliferation, estrogen/estrogen receptor signaling, and cell cycle progression ¹⁵⁻¹⁷. Some investigations have even reported cardiotoxic side effects for DXR that their molecular mechanisms remain to be deciphered in detail 18.

Systems biology and network-based methods are recently used to decipher the molecular mechanisms behind drugs and their possible side effects. Several such studies rely on network topology analysis to identify the effect of chemotherapy on various cancers. These networks can help understand how drugs influence the disease at the molecular level and identify the crucial gene sets underlying various drug effects ¹⁹⁻²². Several network analysis studies of drug-disease associations have been used to predict drug side effects with high accuracy. Global expression data-based computational approaches can utilize gene interaction information for modeling Protein-Protein Interaction Networks (PPINs) and Gene Regulatory Networks (GRNs).

Identifying network modules and their biological functions helps decipher the molecular mechanisms of drug effects, identify new drug targets, predict body response to drugs, and organism behavior ²³⁻²⁵. Gene regulatory networks contain information about regulatory elements of gene expression. These networks can identify regulatory programs and help understand the molecular basis of drug pharmacodynamics and even pharmacogenetics ²⁶. In 2020, a study analyzed the gene regulatory network of breast cancer and identified gene-specific personalized drug treatments ²⁷. Rao Zheng et al also constructed a gene regulatory network of diabetic nephropathy; they recognized essential genes using this method. These findings provide targets for drug development ²⁸. Adel Aloraini et al, in 2018, performed the identification of breast anti-cancer Docetaxel drug targets (DAXX and FGR1) using analysis of gene regulatory network and molecular docking ²⁹.

Molecular mechanisms mediating in breast cancer treatment by DXR and the mechanisms underlying its side effects are not still comprehensively understood. Therefore, in this study, we used protein-protein interaction and gene regulatory networks to identify essential molecular mechanisms and biological functions in response to DXR and the molecular mechanisms responsible for its side effects. We utilize a systems biology approach and bioinformatics analysis of proteinprotein interaction network and Gene Regulatory Networks (GRNs) on omics data of breast cancer treatment using the DXR chemotherapeutic agents. Here, we utilize the protein-protein interaction modules and gene regulatory network motifs to predict and identify drug targets, Gene Ontology (GO) and biochemical pathways mediating in response to ER+ breast cancer and mechanisms underlying its side effects.

Materials and Methods

Data collection

Datasets on breast cancer (MCF-7 cell)/DXR were searched and collected from the *Gene Expression Om-nibus* (*GEO*) database (http://www.ncbi.nlm.nih.gov/geo/) and proteomic publications ^{30,31}. Three datasets (GSE124597 (GPL 15207), GSE39870 (GPL 571), and GSE13477 (GPL 570) were selected to compare breast cancer (MCF-7 cell line)/DXR and non-treatment breast cancer (MCF-7 cell line) for analysis in this study. Figure 1 shows the workflow of this study.

Raw data processing and Data analysis

The datasets' Differentially Expressed Genes (DEGs) were analyzed and identified using GEO2R (https://www.ncbi.nlm.nih.gov/geo/geo2r/), which normalized the data using the GEO query and limma R package. The differentially expressed genes were iden-

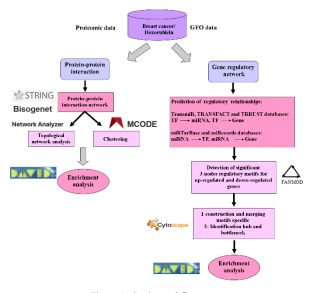


Figure 1. Study workflow.

tified according to p-value<0.05 and Fold Change cutoff of >0.5 and <-0.5 as a threshold. The genes obtained from the three datasets and proteomic publications (mass spectroscopy on MCF-7 treated compared to untreated) were used for further analysis.

Protein-protein interaction network construction

The shared DEGs between the three GSE datasets were obtained using the Venn diagram Tool ³² and unioned with DEGs data extracted from proteomic publications. We applied the STRING (Search Tool for the Retrieval of Interacting Genes/Proteins, https://stringdb.org) with a confidence score of more than 0.7 and Bisogenet app [Human Protein Reference Database (HPRD)] to map the interactions of DEGs obtained from the shared bodes among the three GSE DEGs and unioned with proteomic publications. STRING is a biological database of known and predicted proteinprotein interaction (physical and functional) for many organisms ³³. Bisogenet could build a relation between genes and their products in a fast and user-friendly manner and has multiple applications, including genomics information, protein-protein interactions, protein-DNA interactions, and gene ontology ³⁴. Bisogenet is available in Cytoscape software.

Topological network analysis

The PPI networks obtained from STRING and Bisogenet app were merged using Cytoscape software to analyze the interactions and connections between proteins (http://www.cytoscape.org/). Cytoscape software in bioinformatics for visualizing biomolecular interaction networks (protein-protein, protein-DNA, and genetics interactions) was available for humans and model organisms ³⁵.

This software contains several plugins for functional analysis in PPI networks. Cytoscape network analyzer is a tool that determines the degree and betweenness-centrality of every node as the hub and bottleneck genes. The hubs are node proteins with many interactions, and bottlenecks are nodes with high betweenness centrality ^{35,36}. Finding PPIN hubs and bottlenecks is used to candidate drug targets when drug designing. Besides, it is used to candidate possible disease markers ^{37,38}. We selected the top 10% of nodes with a higher degree and betweenness as hub-bottlenecks for further analysis.

Molecular complex detection (MCODE) cluster subnetworks

The STRING and Bisogenet PPI (HPRD database) networks were merged, and the resulted network was used to identify clusters using the MCODE Plug-in. The MCODE algorithm, one of the Cytoscape plugins, was used to identify highly interconnected sub-networks with parameter settings, including Degree Cut-off=2, Node Score Cutoff=0.2, K-Core=2, and Max-Depth=100³⁹. We considered the MCODE score>3 and the number of nodes>10 as the final clusters' cut-off criteria.

Functional enrichment analysis for hub-bottlenecks and MCODE clusters

The enrichment analysis for Biological process, molecular function, and cellular component and KEGG biochemical pathways (Kyoto Encyclopedia of Genes and Genomes) were performed for the top 10% of the hub and bottleneck genes using the DAVID Tool (Database for Annotation, Visualization, and Integrated Discovery; https://david.ncifcrf.gov/). DAVID is a bioinformatics resource for functional interpretation of a list of genes and can identify GO terms and visualize genes on the KEGG pathway ⁴⁰. The functional enrichment analysis was then performed for pathways of the sub-networks using the STRING database. For the enrichment analysis, STRING uses known systems such as Gene Ontology and KEGG ⁴¹.

TF-miRNA-gene regulatory networks Construction (for UP and down-regulated DEGs)

The up and down-regulated DEGs among the three GSE datasets were identified separately using Venn diagram Tool ³². The identified shared DEGs were unioned with up- and down-regulated proteins retrieved from proteomic publications results, separately. These up- and down-regulated genes were finally used to construct two separate regulatory networks for up- and down-regulated DEGs. The four relationships, including TF-gene, TF-miR, miR-gene, miR-TF, were extracted using the following tools and database to construct two gene regulatory networks for the gene sets.

MiRNAs regulating DEGs

The miRTarBase (http://miRTarBase.mbc.nctu.edu. tw/) and miRecords (http://c1.accurascience.com/mi-Records/) databases were used for identifying miRNAs regulating genes and transcription factors. MiRecords is a database of experimentally validated miRNAtarget interaction ⁴². Besides, miRTarBase is a curated database of experimentally validated miRNA targets with high quality, and its miRNA-target interactions data are collected by receptor assay, microarray, nextgeneration sequencing, and western blot ⁴³.

Transcription factors regulating DEGs

The TFs regulating our target genes were extracted from the TRANSFAC (TRANScription FACtor; https: //genexplain.com/transfac/) and TRRUST databases (transcriptional regulatory relationships unravelled by sentence-based text-mining; https://www.grnpedia.org/ trrust/). TRANSFAC is a database of eukaryotic transcription factors and their experimentally-proven binding sites ⁴⁴. TRRUST is a curated database of human and mouse transcriptional regulatory networks, including 8444 TF-target interactions for 800 TFs in humans and 6552 regulatory interactions for 828 mouse TFs ⁴⁵.

miRNAs inhibiting TFs

The TFs regulating our target genes were fed into the miRTarBase and miRecords databases to obtain miRNAs targeting TFs.

TFs regulating miRNAs

For identifying TFs regulating miRNAs, we used the TransmiR database (http://www.cuilab.cn/transmir). This database contains 3730 TF-miRNA regulations among 19 species from 1349 reports manually curated by surveying >8000 publications and more than 1.7 million tissue-specific. TF-miRNA regulations incorporated based on ChIP-seq data ⁴⁶.

Network construction, motif detection and motif specific sub-networks generation

In a gene regulatory network, network motifs are composed of nodes and regulations that connect the nodes. Some of these regulatory interaction patterns may be significantly high in some networks ⁴⁷. The molecular interactions of motifs are necessary to understand each motif's biological function ⁴⁸. To find the regulatory motifs in up-regulated and down-regulated gene networks, we used FANMOD software. The regulatory relationships (TF-miRNA, TF-Gene, miRNA-Gene, and miRNA-TF) were fed into the FANMOD to identify the motifs with three nodes. FANMOD is a tool for network motifs detection with detection motifs in a big network and analyzes colored networks ⁴⁹. This tool was used to build random networks 1000 times and compared it with the original input network. When randomizing the network in a constant global model, they indicate the frequency of motifs observed in the real network minus the mean of their occurrence in the random network divided by the standard deviation. The motifs with Z-score>2.0 and p-value <0.05 were selected as the significant motifs. TFs, genes and miR-NAs participating in each motif were detected. The motif-related sub-networks of the up- and down-regulated DEGs were then merged (union) in Cytoscape software 3.5.1, separately. Finally, the top 10% of nodes with the highest degree (hub) and betweenness centrality (bottlenecks) were identified in the new networks, separately.

Functional enrichment analysis of GRN

The up- and down-regulated DEGs of motif-related sub-networks were selected for functional enrichment analysis. The sets of DEGs participating in the union of the up- and down-regulated motif-related sub-networks were enriched by the DAVID Tool. The GO terms with p-value <0.05 were selected as significant.

Results

Raw data gathering and analysis

A total of 320 DEGs, including 126 up- and 194 down-regulated genes, were retrieved after analysing the datasets (GSE124597, GSE39870, and GSE1347) and collecting proteomics publications data. Supplementary figure 1 shows the resulted Venn diagram. Supplementary table S1 represents all the up- and down-regulated DEGs

Construction of PPI network

The PPI network was constructed for DEGs using STRING and Bisogenet app (HPRD database) for map-

ping interactions and then merging. The resulted network consisted of 320 nodes and 2519 edges.

Topological analysis

The network analyzer tool was used to study the topological network properties and identify the crucial hub and bottleneck nodes. The topological network properties included the clustering coefficient of 0.385, the shortest path of 60362, network density of 0.045, and diameter of 7. Figure 2 represents a sub-network including the 10% of the genes with the highest degree and betweenness centrality as hubs and bottlenecks, respectively. The top ten hubs and bottlenecks are listed in table 1. The list of 10% of the genes with the highest degree and betweenness centrality are reported in Supplementary table S2.

Module detection

Further analysis of complexes by MCODE app in Cytoscape software revealed 13 sub-networks. The PPI sub-networks are highly connected regions of the network. Three sub-networks were selected according to score>3 and nodes>10 (Table 2, Figure 3). The seed-

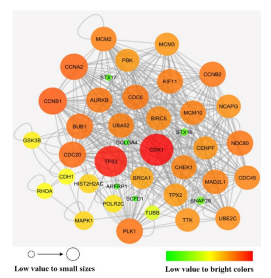


Figure 2. Protein-protein interaction network. The sub-network constructed by Cytoscape software encompasses 10% of hubs and bottlenecks. The nodes' size and color are based on their degree value, and Nodes with dark color (red) have the highest degree.

Table 1. Hub genes related to the breast cancer-doxorubicin network obtained from Cytoscape software

Name genes	Degree	Betweenness centrality
CDK1	111	0.06099772
TP53	105	0.29304896
CCNB1	82	0.03575425
CCNA2	78	0.01470353
CDC20	76	0.01682273
BUB1	73	0.0130227
PLK1	72	0.02161398
CCNB2	71	0.00729474
NDC80	71	0.00428032
CDC6	70	0.014245

Table 2. The PPI sub-networks with Score>3 and nodes>10

Sub- networks	Score	Density nodes	Number of Interactions	Seed node
1	40.227	45	916	MCM10
2	7	15	52	MCM3
3	4.941	18	43	NAPRT

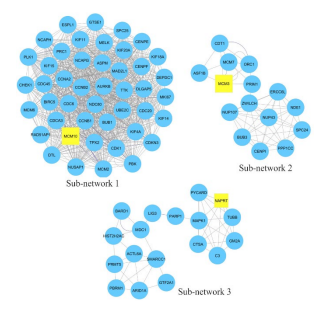


Figure 3. The PPI sub-networks based on highly connected-regions. Sub-networks 1, 2, and 3 were selected based on Score>3 and nodes> 10. Yellow rectangles represent seed nodes.

nodes of these sub-networks included MCM10 (Minichromosome Maintenance 10 Replication Initiation Factor) for sub-network No.1, MCM3 (Minichromosome Maintenance 3 Replication Initiation Factor) for sub-network No.2, and NAPRT (Nicotinate Phosphoribosyltransferase) for sub-network No.3. The results are depicted in table 3. The nodes related to sub-networks are shown in Supplementary table S3.

Gene ontology and KEGG pathway enrichment

We performed Gene Ontology analyses for 10% of hubs and bottleneck genes using the DAVID database.

Table 4 shows the resulting gene ontology terms. Biological process terms reveal that most of the hub genes participate in regulating the cell cycle. The top 10 related molecular functions identified using the DAVID database mediated in protein binding, nucleic acidbinding, *etc.* The cellular component terms showed that most hub genes were present in the cytoskeleton, chromosomes, *etc.*

The DAVID database's KEGG pathway analysis demonstrated that hub and bottleneck genes were involved in the cell cycle, Tumor protein 53 (Tp53) signaling pathway, viral carcinogenesis, viral infections, Forkhead box O (FoxO) signaling pathway, and adherent junctions. Besides, the KEGG pathway analysis showed that SNARE interactions in vesicular transport and neurotrophin signaling pathway were of a significant p-value in enrichment. They could be hypothesized and studied as a signaling possibly related to some side effects of DXR. Table 4 contains the top results of the KEGG pathway analysis by the DAVID database. Supplementary table S4 contains all gene ontology and pathways data related to 10% hub and bottleneck genes.

The top pathway terms significantly enriched in sub-network No.1 included the cell cycle, p53 signaling pathway, viral carcinogenesis, FoxO signaling pathway, and DNA replication. Biochemical pathways involved in sub-network No.2 included cell cycle and DNA replication. The nodes in sub-network No.3 were related to viral carcinogenesis (Table 3).

TF-miRNA-gene regulatory network construction

Identification of miRNA-gene/TF and TF-miRNA/gene interactions: In this study, miRNAs regulating posttranscriptional mRNAs were retrieved from the two experimentally validated databases, including miRTar-Base and miRecords. The up-regulated genes obtained from three GSE and proteomics data were targeted with 1082 miRNAs through 2103 interaction, and the transcription factors regulating genes were identified using TRANSFAC and TRRUST databases. The results revealed that 227 TFs regulated the target genes through 1088 interactions. The number of 1444 miR-NAs targeted 152 TFs with 5979 interactions. TFs regulating miRNAs identified by the validated data of the

	rable 5. KEGG pathway analysis of sub-networks			
KEGG ID	Terms	p-value	Genes	
Sub-network 1				
hsa04110	Cell cycle	1.17E-22	PLK1, TTK, CDC6, CCNA2, CDC20, CCNB2,	
hsa04115	p53 signaling pathway	1.78E-05	CCNB2, CCNB1, CHEK1, CDK1, GTSE1	
hsa05203	Viral carcinogenesis	0.013025793	CCNA2, CDC20, CHEK1, CDK1	
hsa04068	FoxO signaling pathway	0.042317087	CCNB2, CCNB1, PLK1	
Sub-network 2				
hsa04110	Cell cycle	1.90E-04	MCM7, ORC1, MCM3, BUB3	
hsa03030	DNA replication	5.50E-04	MCM7, PRIM1, MCM3	
Sub-network 3				
hsa05203	Viral carcinogenesis	0.040723	C3, GTF2A1, MAPK1	

Table 3. KEGG pathway analysis of sub-networks

Regulatory Motifs in Doxorubicin Effects

Table 4. Top 10 biological processes, molecular functions, cellular components, and KEGG pathways of 10% hub and bottleneck genes identified using the DAVID database (sorted based on p-value <0.05)

ID	Terms	p-value	Genes
Biological proces	\$\$	•	
GO:1903047	mitotic cell cycle process	5.11E-26	NCAPG, MCM10, TTK, KIF11, AURKB, CDC20, CCNB2,
GO:0000278	mitotic cell cycle	4.81E-25	NCAPG, MCM10, TTK, KIF11, AURKB, CDC20, CCNB2,
GO:0022402	Cell cycle process	4.38E-24	GSK3B, NCAPG, MCM10, TTK, BRCA1, KIF11,
GO:0044772	mitotic cell cycle phase transition	9.66E-24	UBE2C, TUBB, PLK1, TTK, MCM10, CDC6, NDC80,
GO:0044770	Cell cycle phase transition	3.77E-23	UBE2C, TUBB, PLK1, TTK, MCM10, CDC6, NDC80,
GO:0007049	Cell cycle	5.82E-23	GSK3B, NCAPG, MCM10, TTK, BRCA1, KIF11,
GO:0007067	mitotic nuclear division	4.28E-22	UBE2C, PLK1, NCAPG, TTK, KIF11, CDC6, RHOA,
GO:0010564	Regulation of cell cycle process	8.14E-21	UBE2C, PLK1, TTK, BRCA1, KIF11, CDC6, RHOA,
GO:0000075	Cell cycle checkpoint	2.37E-20	PLK1, TTK, BRCA1, CDC6, NDC80, AURKB, CCNA2,
GO:0000280	Nuclear division	1.70E-19	UBE2C, PLK1, NCAPG, TTK, KIF11, CDC6, RHOA,
Molecular functi		1.701 17	
GO:0005515	Protein binding	2.71E-08	GSK3B, STX17, STX16, NCAPG, MCM10, TTK, BRCA1,
GO:0035639	Purine ribonucleoside triphosphate binding	2.87E-08	<i>GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,</i>
GO:0032550	Purine ribonucleoside binding	3.10E-08	GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,
GO:0032549	Ribonucleoside binding	3.18E-08	GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,
GO:0001883	Purine nucleoside binding	3.18E-08	<i>GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,</i>
GO:0001882	Nucleoside binding	3.37E-08	<i>GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,</i>
GO:0032555	Purine ribonucleotide binding	4.12E-08	GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,
GO:0017076	Purine nucleotide binding	4.59E-08	GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,
GO:0032553	Ribonucleotide binding	4.70E-08	GSK3B, UBE2C, TUBB, PLK1, TTK, KIF11, CDC6,
GO:0004674	Protein serine/threonine kinase activity	5.55E-08	GSK3B, CCNB2, CCNB1, PLK1, CHEK1, PBK, CDK1,
Cellular compon			,,,
GO:0015630	microtubule cytoskeleton	1.04E-18	GSK3B, TUBB, PLK1, NCAPG, TTK, BRCA1, KIF11,
GO:0005856	Cytoskeleton	5.37E-15	GSK3B, NCAPG, TTK, BRCA1, KIF11, AURKB, CDC20,
GO:0044427	Chromosomal part	1.42E-14	PLK1, NCAPG, TTK, MCM10, BRCA1, NDC80, AURKB,
GO:0044430	Cytoskeletal part	2.93E-14	GSK3B, TUBB, PLK1, NCAPG, TTK, BRCA1, KIF11,
GO:0043228	Non-membrane-bounded organelle	3.89E-14	GSK3B, NCAPG, MCM10, TTK, BRCA1, KIF11,
GO:0043232	Intracellular non-membrane-bounded organelle	3.89E-14	GSK3B, NCAPG, MCM10, TTK, BRCA1, KIF11,
GO:0005694	Chromosome	1.20E-13	PLK1, NCAPG, TTK, MCM10, BRCA1, NDC80, AURKB,
GO:0005819	Spindle	9.41E-13	PLK1, TTK, KIF11, CDC6, AURKB, CDC20, TPX2,
GO:0044446	Intracellular organelle part	1.61E-12	GSK3B, STX17, STX16, NCAPG, MCM10, TTK, BRCA1,
GO:0005815	microtubule-organizing center	1.74E-12	GSK3B, PLK1, NCAPG, BRCA1, AURKB, CDC20,
KEGG	6 6		
hsa04110	Cell cycle	8.81E-20	GSK3B, PLK1, TTK, CDC6, CCNA2, CDC20, CCNB2,
hsa04115	p53 signaling pathway	1.41E-04	CCNB2, CCNB1, CHEK1, CDK1, TP53
hsa05203	Viral carcinogenesis	1.42E-04	CCNA2, CDC20, CHEK1, CDK1, MAPK1, TP53, RHOA
hsa04130	SNARE interactions in vesicular transport	0.008270639	STX17, STX16, SNAP29
hsa04722	Neurotrophin signaling pathway	0.012342719	GSK3B, MAPK1, TP53, RHOA
hsa04068	FoxO signaling pathway	0.016591619	CCNB2, CCNB1, PLK1, MAPK1
hsa05130	Pathogenic Escherichia coli infection	0.018014118	CDH1, TUBB, RHOA
hsa05166	HTLV-I infection	0.018496934	CDC20, GSK3B, CHEK1, TP53, MAD2L1
hsa04520	Adherens junction	0.033403027	CDH1, MAPK1, RHOA

TransmiR database revealed 356 TFs regulated 323 miRNAs with 2011 interactions.

The 194 down-regulated genes obtained from the three GSE datasets and proteomics data were regulated by 1027 miRNAs and 209 TFs through 2457 and 1458 interactions. The number of 1451 miRNAs targeted 133 TFs with 5862 interactions. The analysis of TFs regulating miRNAs by TransmiR revealed that 354

TFs regulated 320 miRNAs with 2008 interactions obtained from experimentally validated data. Finally, the miRNA-gene, TF-Gene, miR-TF, and TF-miR interactions were incorporated to construct two regulatory networks in Cytoscape. The results are shown in table 5. Supplementary table S5 and table S6 contain all relationships in up-regulated and down-regulated, respectively.

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Relationship	Number of pairs	Number of genes	Number of TFs	Number of miRNAs
Up-regulated				
miRNA-gene	2103	52	-	1082
miR-TF	5979	-	152	1445
TF-Gene	1088	56	227	-
TF-miR	2011	-	356	323
Down-regulated				
miRNA-gene	2457	88	-	1027
miR-TF	5862	-	133	1451
TF-Gene	1558	87	209	-
TF-miR	2008	-	354	320

Table 5. Summary of four types of regulatory relationships among miRNA-gene, TF-Gene, miR-TF, and TFmiR interactions

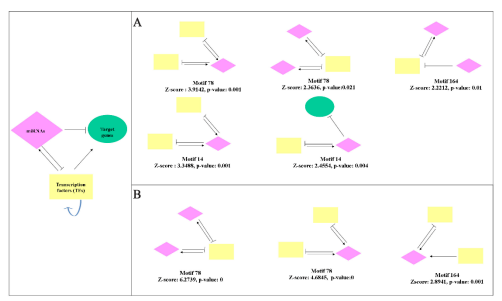


Figure 4. Regulatory motifs consist of miRNAs, TFs, and target genes detected in up and down-regulated gene networks with their Z-score and their p-value. Three types of relationships involved in these motifs included miRNA-gene (miRNA represses gene expression); miRNA-TF (miRNA represses TF gene expression); and TF-miRNA (TF regulates miRNA expression).

Motif detection and generating motif-specific sub-networks

The miRNA-gene, TF-Gene, miRNA-TF, TF-miRNA relationships were combined, and the regulatory networks were constructed. The up- and down-regulated gene networks contained 2250 and 2251 nodes, respectively. The FANMOD software was used to detect the motifs. The types of identified motifs are represented in figure 4 for up- and down-regulated gene networks.

We selected motifs with Z-score>2, p-value<0.05, and at least two-color edges (two types of interactions). Motifs with identification numbers 14, 78, and 164 were finally selected in the up-regulated network. The sub-networks related to these motifs were merged to create a network, including 64 miRNAs, 53 genes, and 321 TFs. The regulatory sub-networks were visualized by Cytoscape 3.5.1. (Figure 5A). Motifs No.78 and 164 were selected and merged in the down-regulated network to create a sub-network including 77 miRNAs, eight genes, and 274 TFs (Figure 5B).

The topological analysis of up and down-regulated GRNs identified the BTG Anti-Proliferation Factor 2

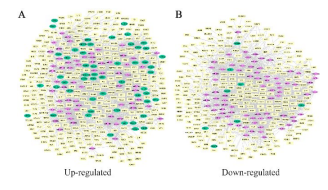


Figure 5. Regulatory sub-networks. A) The sub-network was generated by merging motifs No.14, 78, and 164 in the up-regulated gene network. B) Merging motifs No.78 and 164 in the down-regulated network. Pink diamond nodes are miRNAs, green circular show genes, and yellow rectangles represent the transcription factors.

(BTG2), Specificity Protein 1 (SP1), and TP53 as hubbottleneck, which were also present in up-regulated sub-network. However, none of the genes in the downregulated sub-network was among the GRN 10% of hub-bottlenecks.

Gene ontology and biochemical pathway enrichment analysis

For Gene Ontology analysis, genes obtained from motif detection were submitted in the DAVID database. The top 10 Biological process terms included regulation of cell death and the metabolic process, nucleic acid-binding, transcription binding are top 10 in molecular function terms. The cellular components showed the nucleus and organelle as the top related to the gene-set. The significant pathways were identified from analysis of genes in motif-related sub-networks using the DAVID database. The significant pathways included the p53 signaling pathway, transcriptional misregulation in cancer, cell cycle, PI3K-Akt signaling pathway, viral carcinogenesis, viral infection, Measles, and FoxO signaling pathway (Table 6). Supplementary table S7 contains all gene ontology and KEGG pathways data related to the gene set.

Discussion

Breast cancer is the most common malignancy in women. Its molecular heterogeneity influences the selection of methods in the effective treatment of this cancer². Treatment of routine surgery, radiation therapy, chemotherapy, stabilizing agents, enzyme inhibitors, and immunotherapy are used to treat breast cancer. DXR is an effective chemotherapeutic drug of the anthracycline family used to treat breast cancer ⁵⁰. A comprehensive understanding of the molecular mechanisms of DXR in the treatment of breast cancer is still lacking ⁵¹. The systems biology approach and bioinformatical network analysis for breast cancer in response to DXR help candidate essential genes and pathways mediating in response to this drug for further experimental examinations. The identified and validated targets and pathways may even be used to repurpose new drugs.

Network-based approaches have recently appeared to be a powerful tool to investigate pathobiological processes and the molecular complexity of disease aetiology by identifying disease-specific network clusters such as MCODE clusters in PPI networks. The nodes participating in these particular regions usually have critical roles and biological functions 52. Nodes participating in regulatory motifs also are of biological importance in GRNs ⁵³. We applied the MCODE clusters and GRN motifs to predict the molecular mechanisms underlying the treating effect of DXR on breast cancer and its side effects. This study selected genomics and proteomics data to integrate and explore critical genes and molecular pathways. The present study is the first in silico analysis that uses bioinformatics analysis to

predict the essential genes and pathways of breast cancer treated with DXR and its side effects.

Our systematic analysis of the PPI MCODE clusters and GRN motif-related sub-networks of the MCF7 cell line in response to DXR demonstrated that TP53, MCM10, and MCM3 are the top hub-bottlenecks and MCODE cluster seeds in response to DXR (Supplementary table S8). The functional enrichment analysis indicated that hub-bottleneck and cluster nodes were involved in the cell cycle, P53 signaling pathway, FoxO signaling pathway, and viral carcinogenesis.

TP53 is a hub-bottleneck protein in our PPIN and GRN. TP53 is a gene with a high degree and betweenness centrality over-expressed in the MCF-7 cell line in response to DXR. This protein can recognize DNA damage, stop the cell cycle at the G1/S regulation point, and activate DNA repair proteins. Therefore, TP53 can initiate apoptosis if DNA damage is irreparable ⁵⁴. TP53 was up-regulated in MCF-7 cells treated with DXR. Therefore, it can be concluded that DXR activates the repair system and instigates apoptosis in cancer cells possible through P53 mediation.

The Minichromosome Maintenance proteins (MCM) are critical regulators in DNA replication 55. These proteins are implicated in cancer initiation and progression, and their expression is up-regulated in a wide range of epithelial malignancies ⁵⁶. MCM10, an MCM family member, is an essential factor for DNA replication by binding with Cell Division Cycle 45 (CDC45) and is essential in breast cancer progression ⁵⁷. Alcivar AL et al reported that cells depleted of MCM10 showed instability of replication fork 58. Wei-Dong Yang et al in 2019 showed MCM10 was significantly overexpressed in breast carcinoma and involved in proliferation, migration, and invasion. Therefore, it can induce metastasis via the Wnt/β-catenin pathway in breast cancer ⁵⁹. Our results identified MCM10 as a critical node in the network. Given the importance of this protein and the lack of experimental literature about its mediation in response to DXR, we suggest that its experimental investigation seems necessary.

MCM3, another MCM member, is over-expressed in various human cancers ⁶⁰. MCM3 is one of the cell cycle markers that regulates the growth, migration, and invasion of cells ⁶¹. Our study showed that MCM3 was also a down-regulated protein of importance in the networks. Therefore, we hypothesize that DXR can probably inhibit DNA replication, invasion, and metastasis by down-regulating the MCM10 and MCM3 genes.

The functional enrichment analysis showed regulation of the cell cycle, p53 signaling, viral carcinogenesis, Human T-Lymphotropic Virus type 1 (HTLV-1) infection, and FoxO signaling pathway were the top terms related to hubs and bottlenecks in PPIN, MCODE clusters, and GRNs. Besides, Soluble Nethylmaleimide-sensitive factor Attachment protein

Table 6. The table represents the top 10 biological	processes, molecular function	, cellular components,	and KEGG pathways identified using the
]	DAVID tool (sorted based on p	o-value<0.05)	

GO ID	Terms	p-value	Genes
Biological proce		•	
GO:0010941	Regulation of cell death	4.37E-10	CDKN1A, TIGAR, BTG2, CEBPB, GADD45A,
GO:0010604	Positive regulation of macromolecule metabolic	4.69E-10	FOXA1, KDM5B, CDKN1A, BTG2, CEBPB,
GO:0042981	process Regulation of the apoptotic process	6.86E-10	CDKN1A, TIGAR, BTG2, CEBPB, GADD45A,
GO:0043067	Regulation of programmed cell death	8.18E-10	CDKN1A, TIGAR, BTG2, CEBPB, GADD45A,
GO:0008219	Cell death	2.29E-09	CDKN1A, BTG2, CEBPB, GLS2, GATA3,
GO:0009893	Positive regulation of the metabolic process	2.54E-09	FOXA1, KDM5B, CDKN1A, BTG2, CEBPB,
GO:0009095	Positive regulation of cellular metabolic process	1.09E-08	FOXA1, CDKN1A, BTG2, CEBPB, SRSF1,
GO:0031323 GO:0012501	Programmed cell death	2.31E-08	CDKN1A, TIGAR, BTG2, CEBPB, GADD45A,
GO:0012501 GO:0010628	Positive regulation of gene expression	2.33E-08	FOXA1, KDM5B, CEBPB, NFYC, SRSF1,
GO:0006915	Apoptotic process	2.33E-08 4.17E-08	CDKN1A, TIGAR, BTG2, CEBPB, GADD45A,
Molecular funct		4.17L-08	CDKNIA, HOAK, BIO2, CEBI B, GADD43A,
violeculai funci			
GO:0000982	Transcription factor activity, RNA polymerase II core promoter proximal region sequence-specific binding	1.27E-09	FOXA1, BTG2, CEBPB, NFYC, GATA3,
GO:0044212	Transcription regulatory region DNA binding	1.61E-09	FOXA1, PRMT5, CEBPB, GADD45A, NFYC,
GO:0000975	Regulatory region DNA binding	1.70E-09	FOXA1, PRMT5, CEBPB, GADD45A, NFYC,
GO:0001067	Regulatory region nucleic acid binding	1.73E-09	FOXA1, PRMT5, CEBPB, GADD45A, NFYC,
GO:0001228	Transcriptional activator activity, RNA polymerase II transcription regulatory region sequence-specific binding	9.84E-09	FOXA1, FOSL1, CEBPB, CREB1, MAF, MAFB,
GO:0000981	RNA polymerase II transcription factor activity, sequence-specific DNA binding	2.86E-08	FOXA1, BTG2, CEBPB, NFYC, GATA3,
GO:0000987	Core promoter proximal region sequence-specific DNA binding	4.26E-08	FOSL1, MUC1, CEBPB, CREB1, MAFB, SP1,
GO:0003690	Double-stranded DNA binding	4.45E-08	PRMT5, CEBPB, NFYC, XPC, GATA3, RUNX2,
GO:0001159	Core promoter proximal region DNA binding	4.50E-08	FOSL1, MUC1, CEBPB, CREB1, MAFB, SP1,
GO:0000976	Transcription regulatory region sequence-specific DNA binding	4.98E-08	PRMT5, CEBPB, NFYC, GATA3, RUNX2,
Cellular compo	nent		
GO:0070013	Intracellular organelle lumen	1.61E-06	FOXA1, KDM5B, CDKN1A, CEBPB, GLS2,
GO:0043233	Organelle lumen	2.34E-06	FOXA1, KDM5B, CDKN1A, CEBPB, GLS2,
GO:0031974	Membrane-enclosed lumen	3.13E-06	FOXA1, KDM5B, CDKN1A, CEBPB, GLS2,
GO:0005667	Transcription factor complex	7.01E-06	CEBPB, CREB1, MAFB, CDK4, NFYC, GATA3,
GO:0031981	Nuclear lumen	1.77E-05	FOXA1, KDM5B, CDKN1A, CEBPB, SRSF1,
GO:0000785	Chromatin	1.93E-05	MUC1, CEBPB, CREB1, MAF, SP1, CDK4,
GO:0005654	Nucleoplasm	2.81E-05	CTSA, KDM5B, PRMT5, CDKN1A, CEBPB,
GO:0044428	Nuclear part	3.71E-05	FOXA1, KDM5B, CDKN1A, CEBPB, SRSF1,
GO:0005634	Nucleus	5.48E-05	FOXA1, KDM5B, CDKN1A, CEBPB, SRSF1,
GO:0044422	Organelle part	2.45E-04	FOXA1, KDM5B, CDKN1A, CEBPB, SYNM,
KEGG	-		
hsa04115	p53 signaling pathway	2.01E-06	CDKN1A, ZMAT3, CDK4, GADD45A, MDM2, FAS, TP53
hsa05202	Transcriptional misregulation in cancer	4.16E-05	CDKN1A, CEBPB, MAF, SP1, MDM2, TP53, RUNX2, PBX1
hsa04110	Cell cycle	6.97E-04	CDKN1A, CDK4, GADD45A, PLK1, MDM2, TP
hsa04151	PI3K-Akt signaling pathway	0.013789	CDKNIA, CREBI, CDK4, MDM2, BRCAI, TP5. EPHA2
hsa05166	HTLV-I infection	0.015101	FOSL1, CDKN1A, CREB1, CDK4, TP53, ATF3
hsa05203	Viral carcinogenesis	0.030309	CDKN1A, CREB1, CDK4, MDM2, TP53
hsa05162	Measles	0.041521	CDK4, FAS, TNFRSF10B, TP53
hsa04068	FoxO signaling pathway	0.042304	CDKN1A, GADD45A, PLK1, MDM2

Receptor (SNARE) interactions in vesicular transport and neurotrophin signaling pathway were identified in our PPIN KEGG pathway results that could be related to side effects of DXR. Deregulation of the cell cycle is one of the mechanisms involved in the malignant phenotype of cancer. Regulation of the cell cycle can be used as a therapeutic targeting strategy against cancer ⁶². The chemother-

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apeutic agent DXR can cause cell arrest in the G1phase of the cell cycle ⁶³. In addition, Kim HS *et al* in 2009 reported that this drug could induce intracellular apoptotic signaling through up-regulation of Fas expression ⁶⁴.

Viral carcinogenesis and HTLV-1 infection were other pathways related to hubs and bottlenecks enrichments. The virus has known oncogenic potential in specific cancers, including the cervix, liver, head and neck, some lymphomas, and breast cancer ⁶⁵. HTLV-1 is one of the viruses that encode oncogenic protein *Tax1* Binding Protein 1 (TAX1) and help breast cancer progression ⁶⁶. TAX1 protein can inactivate the function of cellular TP53 and postpone the G1 cell cycle arrest required for repairing DNA in response to DNA damage ⁶⁷. DXR can induce apoptotic cell death in HTLV-1 infected cells ⁶⁸.

Our model identified that the P53 signaling pathway was a significantly enriched KEGG pathway related to hub-bottlenecks and MCODE clusters. McSweeney *et al* in 2019 reported TP53 as a critical regulator of transcriptomic changes induced by DXR ⁶⁹. Ru-Wei Lin *et al* in 2018 showed DXR-induced apoptosis in response to DNA damage by overexpression of TP53 ⁷⁰. In addition, p53 interferes in cell metabolism, ferroptosis, autophagy, and generation of ROS ⁷¹. These validate the predictions performed by our model and justify performing experimental examinations on its other findings.

The FoxO signaling pathway was another identified signaling predicted by the model. FoxO transcription factors are tumor suppressors that mediate redox homeostasis, proliferation, survival, and Phosphatidylinositol-4,5-Bisphosphate 3-Kinase (PI3K) ⁷². Rosaline CY et al. reported that the cancer treatment with DXR increased FOXO3a activity 73. During apoptosis, FOXOs are involved in expressing death receptor ligands such as Fas ligand, TNF, Bim, bNIP3, and Bcl-XL ⁷⁴. The enhanced FOXO3a activity increased the expression of ABCB1, a plasma membrane P-glycoprotein, which functions as an efflux for various anti-cancer agents ⁷³. FOXO proteins play an essential role in glucose homeostasis by promoting gluconeogenic enzyme expression 75. The dysfunction of FoxO1 pathways involves several metabolic diseases, including atherosclerosis, diabetes, non-alcoholic fatty liver disease, and obesity 76. Notably, FOXO proteins are involved in physiological processes. Activation and inhibition of these proteins could have intolerable side effects.

Other signaling pathways significantly enriched in our study were SNARE interactions in vesicular transport and neurotrophin signaling pathway. SNAREs are a group of transmembrane proteins which create a bridge for interaction vesicle to its fusion partner. This vesicle trafficking is regulated by a separate process and stimulates the SNARE complex formation ⁷⁷. The dysfunction of membrane trafficking is associated with cardiovascular events ⁷⁸. DXR disrupts the trafficking membrane by reducing Syntaxin 17 (STX17), Syntaxin 16 (STX16), and Synaptosome Associated Protein 29 (SNAP29) protein expression, thereby probably having side effects on the heart in this way. Besides, neurotrophins and their receptors are regulatory factors in heart and vascular development. These molecules regulate angiogenesis and vasculogenesis, controlling the survival of endothelial cells, vascular smooth muscle cells and cardiomyocytes 79. Therefore, DXR may lead to cardiotoxicity through dysfunction of the neurotrophin signaling pathway with a change in expression of Glycogen Synthase Kinase 3 Beta (GSK3B), Mitogen-Activated Protein Kinase 1 (MAPK1), TP53, and Ras Homolog Family Member A (RHOA) proteins. In addition, the SNARE complex is vital in the formation of vesicle fusion, vesicle recycling and neurotransmitter release. The defects in the formation of the SNARE complex, SNARE-dependent exocytosis, and SNAREmediated vesicle fusion are associated with neurological diseases 80.

Altogether we suggest that DXR regulates repair, apoptosis, invasion and metastasis of breast cancer cells. Its side effects are probably mediated by SNARE interactions in vesicular transport and neurotrophin signaling pathway and FoxO signaling pathway through up- and down-regulated genes primarily identified in our model. Further studies *in vitro* and *in vivo* are required to validate some of our novel findings.

Conclusion

This study applied a network-based approach (PPIN and GRN) to reveal the network hubs and bottlenecks and 3-nodes motifs consisting of TFs, miRNAs, and target genes underlying the DXR effect on breast cancer. We identified the molecular mechanisms and pathways mediating in response to DXR treatment. The hubs and bottlenecks of PPIN and GRN and PPIN MCODE clusters of differentially expressed genes in the MCF-7 cell line treated with DXR revealed that the essential biological processes and pathways are related to cell cycle, p53, viral carcinogenesis, and FoxO signaling pathway. Besides, SNARE interactions in vesicular transport and neurotrophin signaling pathway and FoxO signaling pathway were identified as pathways possibly mediating in its side effects. MCM10 and MCM3 were identified as essential DEGs mediating in response to DXR and are recommended for further investigations since their role is not studied sufficiently so far. We hope that our analysis results can understand the mechanisms involved in response to DXR and its side effects and help design further experimental investigations.

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Conflict of Interest

The authors declare no conflicts of interest.

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Supplementary

Complementer Table Cl	Un /D aslated anotating and anotating the later.
Supplementary Table S1.	Up/Down-regulated proteins are represented below

Up regulated	Down regulated	Up regulated	Down regulated	Up regulated	Down regulate
BTG2	ESPL1	CAT	DLGAP5		ZFR
GDF15	E2F8	TP53I3	RAD51AP1		PBRM1
WISP2	ORC3	CCS	MKI67		NKRF
HIC2	TPX2	PML	DTL		PRDM15
TIGAR	GPSM2	PRKD1	PAICS		ZFP2
EPHA2	CCNB1	PXN	MPHOSPH9		TTC5
ATF3	BRCA1	HBA2	MCM6		C17orf49
FAS	BIRC5	LIG1	CENPI		ZBTB9
FAP1	PLK1	CDK1	CCNF		HIST2H2AB
XPC	ST8SIA4	PRKAA1	SLC25A12		YY2
NADSYN1	TUBB	ASNS	KIF20A		HIST2H3D
KRT15	CHEK1	CDH1	TTK		RFX8
MOSPD1	MUC1	CDK4	CDKN3		CREB1
STOM	KIF11	CDKN1A	NCAPG		PNKP
YPEL5	SKP2	MPST	CENPF		PURB
AVPI1	ORC1	SCFD1	AKAP8		NABP2
OPYSL4		HMGCL	PES1		TAF2
	CDC6				
GM2A	ZWILCH	SMARCA5	H2AF		TAF7
DUSP1	DBF4	FOXD4L1	MKI67IP		MTA2
WSB1	KIF14	MSH2	CSNK1A1		HOXC11
РНҮН	MAD2L1	SRSF1	PPP1CC		HOXD10
ZMAT3	POLA1	GNA13	BUB3		ALX1
JMK2	AURKB	PITPNB	NDE1		MYBBP1A
P53I3	MCM3	TANC2	ERCC6L		VSX1
AK1	TIPIN	EPN1	SPC24		POLE4
PSG9	C17orf75	MBD3	NUP43		STX16
EPPK1	KIF4A	RIC8A	LIG3		SNAP29
FMEM158	BARD1	GRSF1	NSMCE2		STX17
GADD45A	MCM10	AMBRA1	PARP1		TLK1
MDM2	FBXO5	PPP1R13L	BCL3		HIRIP3
FRAF4	MELK	MLL	CCNT1		ASF1B
CSAD	CDC20	POU4F3	DNA2		ACTL6A
SLC6A8	GART	PRKRA	ENO1		SUPT4H1
INFRSF10B		RUNX2	GATA3		LEO1
	NDC80				
ARFGAP3	CCNA2	DLG1	GTF2A1		MRGBP
MAFB	GTSE1	ALB	H1F0		UTP3
CABYR	BUB1	GSTK1	HIST1H1E		SRPK1
CDKN1A	PBK	PRDX4	HIST1H1B		BRD8
MORC4	DEPDC1	SNAPIN	HMGN1		PRMT5
MAF	NCAPH	DTNBP1	AGFG1		PYGO2
FDXR	MDC1	SLC6A17	MCM3		GPI
PIDD1	PFAS	TRIM37	NFYB		C3
GPR87			PBX1		MTPN
	HNRNPD	CHP1			
ACTA2	CENPE	PYCARD	POLR2C		OGDH
ANXA4	ASPM	NR3C2	MAPK1		HPRT1
SYNM	UBE2S	NFYC	RFC2		MYCBP2
CYFIP2	LMNB1	ZNF24	RPL6		MAP2
FOSL1	PRIM1	MTA1	SMARCA1		ANXA1
PDE4A	CDCA3	ZGPAT	SMARCC1		RYK
GLS2	ATAD2	ARX	SP100		UACA
SAT1	BRIP1	FOSL2	SURF6		CD276
GABPA		NUP93			
	STIL		BRPF1		CTNNA2
FP53	UBE2C	NUP153	DEK		MYO1D
CEBPB	CCNB2	NUP107	KDM5D		DDN
FOXA1	MCM7	GSK3A	ARID1A		LIMS1
SP1	PRC1	GSK3B	HIST2H2AC		RHOA
ANK3	CDT1	PREP	NCOR2		LTBP2
GOLGA4	CDC45	CTSA	BCLAF1		PPM1A
ARFRP1	GEMIN2	SCPEP1	TOX4		PPP1CB
MACF1					
	SPC25	ARHGEF2	SRA1		UBA52
NQO1	WDHD1		HUWE1		SNX6
NUDT1	MCM2		PQBP1		ANKRD17
NDUFS8	DUT		SRRM1		
RRM2B	KIF18A		KDM5B		
NAPRT1	KIF15		AKAP8L		
SRXN1	MSH2		NUSAP1		



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10% degree and betweenness centrality CDK1 TP53 CCNB1 CCNA2 CDC20 BUB1 PLK1 NDC80 CCNB2 CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1 CDC45
TP53 CCNB1 CCNA2 CDC20 BUB1 PLK1 NDC80 CCNB2 CDC6 AURKB MAD2L1 UBA52 MCM2 MCM10 CHEK1
CCNB1 CCNA2 CDC20 BUB1 PLK1 PLK1 NDC80 CCNB2 CCNB2 CCNB2 CDC6 AURKB CDC6 AURKB SCDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 AURKB CDC6 CDC6 CDC6 CDC6 CDC6 CDC6 CDC6 CDC
CCNA2 CDC20 BUB1 PLK1 PLK1 NDC80 CCNB2 CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
CDC20 BUB1 PLK1 PLK1 NDC80 CCNB2 CCNB2 CDC6 CDC6 AURKB AURKB MAD2L1 UBA52 UBA52 MCM2 MCM2 MCM10 CHEK1
BUB1 PLK1 NDC80 CCNB2 CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
PLK1 NDC80 CCNB2 CDC6 AURKB AURKB MAD2L1 UBA52 UBA52 MCM2 MCM10 CHEK1
NDC80 CCNB2 CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
CCNB2 CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
CDC6 AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
AURKB KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
KIF11 MAD2L1 UBA52 MCM2 MCM10 CHEK1
MAD2L1 UBA52 MCM2 MCM10 CHEK1
UBA52 MCM2 MCM10 CHEK1
MCM2 MCM10 CHEK1
MCM10 CHEK1
CHEK1
CDC45
00043
BIRC5
UBE2C
NCAPG
TTK
CENPF
MCM3
РВК
TPX2
STX16
SNAP29
GOLGA4
ARFRP1
SCFD1
MAPK1
STX17
HIST2H2AC
RHOA
BRCA1
GSK3B
POLR2C
TUBB
CDH1

Supplementary Table S2. Hubs and bottlenecks (top 10%) related to PPI
network obtained by Cytoscape software



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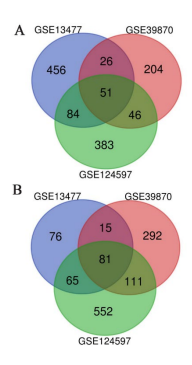
Regulatory Motifs in Doxorubicin Effects

Subnetwork 1	Subnetwork 2	Subnetwork 3
CDC45	ASF1B	GTF2A1
TTK	PRIM1	NAPRT
MELK	PPP1CC	GM2A
BUB1	MCM3	C3
CCNB2	ORC1	PBRM1
DLGAP5	MCM7	MAPK1
KIF15	ZWILCH	TUBB
KIF20A	NDE1	PARP1
CDKN3	ERCC6L	HIST2H2AC
CDCA3	NUP43	ARID1A
KIF14	CDT1	SMARCC1
BIRC5	SPC24	BARD1
MKI67	CENPI	ACTL6A
KIF11	NUP107	LIG3
DTL	BUB3	PRMT5
CDK1		MDC1
KIF4A		CTSA
NDC80		PYCARD
CENPE		
CDC6		
CENPF		
CDC20		
GTSE1		
PRC1		
CCNA2		
UBE2C		
ASPM		
PLK1		
CCNB1		
MCM2		
MCM6		
NCAPG		
CHEK1		
MAD2L1		
RAD51AP1		
PBK		
AURKB		
NCAPH		
KIF18A		
TPX2		
ESPL1		
SPC25		
NUSAP1		
MCM10		
DEPDC1		
DLIDCI		

Supplementary Table S3. Each column represents proteins available in one MCODE cluster







Supplementary Figure S1. Venn diagram for DEGs of GEO datasets (GSE124597, GSE39870, and GSE13477) related to MCF-7 cell line treated with doxorubicin. A) Venn diagram related to up-regulated genes B) Venn diagram related to down-regulated genes.

