PRLR and MNAT1 are promising druggable targets for treating Parkinson Disease that control activity of PGR, NR3C1 and RAD21 transcription factors on promoters of differentially expressed genes

Demo User geneXplain GmbH info@genexplain.com Data received on 02/07/2020 ; Run on 06/07/2020 ; Report generated on 06/07/2020

Genome Enhancer release 2.0.1 (TRANSFAC®, TRANSPATH® and HumanPSD™ release 2020.2)



Abstract

In the present study we applied the software package "Genome Enhancer" to a data set that contains *transcriptomics* data. The study is done in the context of *Parkinson Disease*. The goal of this pipeline is to identify potential drug targets in the molecular network that governs the studied pathological process. In the first step of analysis pipeline discovers transcription factors (TFs) that regulate genes activities in the pathological state. The activities of these TFs are controlled by so-called master regulators, which are identified in the second step of analysis. After a subsequent druggability checkup, the most promising master regulators are chosen as potential drug targets for the analyzed pathology. At the end the pipeline comes up with (a) a list of known drugs and (b) investigational active chemical compounds with the potential to interact with selected drug targets.

From the data set analyzed in this study, we found the following TFs to be potentially involved in the regulation of the differentially expressed genes: PGR, NR3C1, ZEB1, RAD21, POU2F1 and YY1. The subsequent network analysis suggested

- TFIIH-CAK
- PKCtheta
- prlr:tec:Vav
- Cdc25A

as the most promising molecular targets for further research, drug development and drug repurposing initiatives on the basis of identified molecular mechanism of the studied pathology. Having checked the actual druggability potential of the full list of identified targets, both, via information available in medical literature and via cheminformatics analysis of drug compounds, we have identified the following drugs as the most promising treatment candidates for the studied pathology: Acetylcysteine, Ruxolitinib, D-Myo-Inositol-Hexasulphate and 2,4-Dihydroxybenzoic Acid.

1. Introduction

Recording "-omics" data to measure gene activities, protein expression or metabolic events is becoming a standard approach to characterize the pathological state of an affected organism or tissue. Increasingly, several of these methods are applied in a combined approach leading to large "multiomics" datasets. Still the challenge remains how to reveal the underlying molecular mechanisms that render a given pathological state different from the norm. The disease-causing mechanism can be described by a re-wiring of the cellular regulatory network, for instance as a result of a genetic or epigenetic alterations influencing the activity of relevant genes. Reconstruction of the disease-specific regulatory networks can help identify potential master regulators of the respective pathological process. Knowledge about these master regulators can point to ways how to block a pathological regulatory cascade. Suppression of certain molecular targets as components of these cascades may stop the pathological process and cure the disease.

Conventional approaches of statistical "-omics" data analysis provide only very limited information about the causes of the observed phenomena and therefore contribute little to the understanding of the pathological molecular mechanism. In contrast, the "upstream analysis" method [1-4] applied here has been deviced to provide a casual interpretation of the data obtained for a pathology state. This approach comprises two major steps: (1) analysing promoters and enhancers of differentially expressed genes for the transcription factors (TFs) involved in their regulation and, thus, important for the process under study; (2) re-constructing the signaling pathways that activate these TFs and identifying master regulators at the top of such pathways. For the first step, the database TRANSFAC® [6] is employed together with the TF binding site identification algorithms Match [7] and CMA [8]. The second step involves the signal transduction database TRANSPATH® [9] and special graph search algorithms [10] implemented in the software "Genome Enhancer".

The "upstream analysis" approach has now been extended by a third step that reveals known drugs suitable to inhibit (or activate) the identified molecular targets in the context of the disease under study. This step is performed by using information from

HumanPSD^M database [5]. In addition, some known drugs and investigational active chemical compounds are subsequently predicted as potential ligands for the revealed molecular targets. They are predicted using a pre-computed database of spectra of biological activities of chemical compounds of a library of 2507 known drugs and investigational chemical compounds from HumanPSD^M database. The spectra of biological activities for these compounds are computed using the program PASS on the basis of a (Q)SAR approach [11-13]. These predictions can be used for the research purposes - for further drug development and drug repurposing initiatives.

2. Data

For this study the following experimental data was used:

Table	1.	Experimental	datasets	s used	in	the	stud

File name	Data type
GSE145804_DESeq2_final	Transcriptomics



Figure 1. Annotation diagram of experimental data used in this study. With the colored boxes we show those sub-categories of the data that are compared in our analysis.

3. Results

We have compared the following conditions: noRA_Dox versus noRA_noDox.

3.1. Identification of target genes

In the first step of the analysis **target genes** were identified from the uploaded experimental data. We applied the edgeR tool (R/Bioconductor package integrated into our pipeline) and compared gene expression in the following sets: "noRA_Dox" with "noRA_noDox". edgeR calculated the LogFC (the logarithm to the base 2 of the fold change between different conditions), the p-value and the adjusted p-value (corrected for multiple testing) of the observed fold change. As a result, we detected 8084 upregulated genes (LogFC>0) out of which 578 genes were found as significantly upregulated (p-value<0.1) and 8862 downregulated genes (LogFC<0) out of which 726 genes were significantly downregulated (p-value<0.1). See tables below for the top significantly up- and downregulated genes. Below we call **target genes** the full list of up- and downregulated genes revealed in our analysis (see tables in Supplementary section).

Table 2. Top ten significant **up-regulated** genes in noRA_Dox vs. noRA_noDox. See full table \rightarrow

ID	Gene symbol	Gene description	logFC	logCPM	PValue	FDR
ENSG00000145335	SNCA	synuclein alpha	4.38	10.36	2.71E-88	4.59E- 84
ENSG00000118785	SPP1	secreted phosphoprotein 1	2.98	-0.28	6.23E-7	2.11E- 3
ENSG00000169282	KCNAB1	potassium voltage-gated channel subfamily A member regulatory beta subunit 1	2.91	0.49	1.32E-7	5.61E- 4
ENSG00000162692	VCAM1	vascular cell adhesion molecule 1	2.67	-4.05E-2	1.1E-5	2.34E- 2
ENSG0000237280	AL136982.3	novel transcript	2.15	0.2	1.42E-4	0.16
ENSG00000214892	USP8P1	ubiquitin specific peptidase 8 pseudogene 1	1.81	0.19	4.4E-3	0.83
ENSG0000224837	GCSHP5	glycine cleavage system protein H pseudogene 5	1.72	-0.1	1.84E-3	0.6
ENSG00000243300	null	null	1.65	0.44	2.8E-3	0.68
ENSG00000229474	PATL2	PAT1 homolog 2	1.53	-0.14	7.97E-3	0.98
ENSG00000236813	BTF3P8	basic transcription factor 3 pseudogene 8	1.53	-0.48	9.95E-3	0.98

Table 3. Top ten significant **down-regulated** genes in noRA_Dox vs. noRA_noDox. See full table \rightarrow

ID	Gene symbol	Gene description	logFC	logCPM	PValue	FDR
ENSG00000186081	KRT5	keratin 5	-10.36	2.57	9.35E-7	2.64E- 3
ENSG00000257594	GALNT4	polypeptide N-acetylgalactosaminyltransferase 4	-2.37	1.39E-2	4.31E-5	6.08E- 2
ENSG0000167244	IGF2	insulin like growth factor 2	-2.13	-0.49	1.23E-3	0.47
ENSG00000255115	AP002812.4	family with sequence similarity 162, member A (FAM162A) pseudogene	-1.97	-0.38	2.22E-3	0.61
ENSG00000134955	SLC37A2	solute carrier family 37 member 2	-1.9	-0.42	9.67E-3	0.98
ENSG00000111679	PTPN6	protein tyrosine phosphatase non-receptor type 6	-1.9	-0.13	4.43E-3	0.83
ENSG00000179846	NKPD1	NTPase KAP family P-loop domain containing 1	-1.87	-0.33	3.5E-3	0.79
ENSG00000137801	THBS1	thrombospondin 1	-1.83	0.49	4.4E-3	0.83
ENSG0000269054	AC012313.6	novel transcript, antisense to ZNF497	-1.82	-9.51E-2	1.69E-3	0.56
ENSG00000224886	AL132656.1	novel pseudogene	-1.82	-0.35	6.76E-3	0.98

3.2. Functional classification of genes

A functional analysis of differentially expressed genes was done by mapping the significant up-regulated and significant downregulated genes to several known ontologies, such as Gene Ontology (GO), disease ontology (based on HumanPSD[™] database) and the ontology of signal transduction and metabolic pathways from the TRANSPATH® database. Statistical significance was computed using a binomial test.

Figures 3-8 show the most significant categories.

Heatmap of differentially expressed genes in noRA_Dox vs. noRA_noDox

A heatmap of all differentially expressed genes playing a potential regulatory role in the system (enriched in TRANSPATH® pathways) is presented in Figure 2.





Figure 2. Heatmap of genes enriched in Transpath categories. The colored bar at the top shows the types of the samples according to the legend in the upper right corner. See full diagram \rightarrow

Up-regulated genes in noRA_Dox vs. noRA_noDox:

578 significant up-regulated genes were taken for the mapping.

GO (biological process)

			golola	lical_proces	is Gene Untolog	y treemap			
positive regulation of centriole elongation	regulation of centriole elongation	L-fucose catabolic proc	fucos ess p	e catabolic rocess	membran membrane d	e to ocking	response to metal ion	cellular extravasation	astrocyte cell migration
positive regulation of centriole replication	positive regulation of centrosome duplication	L-fucose metabolic proc	ess fucos p	e metabolic rocess	membrane d membran membrane d	ocking ne to docking re	response to iron ion	T cell extravasation	glial cell migration
positive regulation	positive regulation of centrosome cycle of centriole elongation	L-fucose of carnitine t	catabolic p	quaternary ammonium group transport	regulation of leukocyte migratio	regulation on of cellular extravasatior	proteolysis	centriole elongation	cilium movement
DNA replication checkpoint	mitotic DNA replication checkpoint DNA damage checkpoint	amino-acid bet	aine transport	port	regulation of leuk defense response to virus	o <mark>cyte migration</mark> response to virus	macromolecule metabolic process	centriole elongation nitrogen compound metabolic process nitrogen compound	cilium movement negative regulation of G protein-coupled receptor signaling pathway negative regulation of G protein-coupled receptor
DNA integrity checkpoint	cell cycle checkpoint	G1/S transition of mitotic cell cycle	mitotic cell cycle phase transition	cell cycle G1/S phase transition	defense respo cellular response to ionizing	response to response to ionizing radiation	metabolic process metabolic process metabolic process	metabolic process sperm-egg recognition sperm-egg recognition	signaling pathway regulation of response to external stimulus regulation of response to external stimulus
AMP metabolic proce	ss nucleoside monophosphate phosphorylation	G1/S transition negative regulation of necroptotic process	negative regulation of programmed necrotic cell	cell cycle negative regulation of necrotic cell death	radiation cellular res ionizing r	sponse to adiation	response to cobalt ion response to cobalt ion oligosaccharide	double-strand break repair via break-induced replication	regulation of SNARE complex assembly regulation of SNARE complex assembly complement-dependent
purine ribonucleosid monophosphate metabolic process AMP meta	bolic process	perative regula	death	tic process	DNA rep	lication	metabolic process oligosaccharide metabolic process	metabolic process organic substance metabolic process	cytotoxicity

Figure 3. Enriched GO (biological process) of up-regulated genes in noRA_Dox vs. noRA_noDox. Full classification \rightarrow

biological_process Gene Ontology treemap



Figure 4. Enriched TRANSPATH® Pathways (2020.2) of up-regulated genes in noRA_Dox vs. noRA_noDox. Full classification \rightarrow

HumanPSD(TM) disease (2020.2)



🔳 Central Nervous System Diseases 📕 Brain Diseases 📕 Dry Eye Syndromes 🔳 Eosinophilia

Lacrimal Apparatus Diseases AIDS-Related Opportunistic Infections

🔳 Digestive System Abnormalities 🔳 Opportunistic Infections 📕 Fibromuscular Dysplasia

■ Giant Cell Tumor of Tendon Sheath ■ Synovitis ■ Synovitis, Pigmented Villonodular

🔳 Tendinopathy 📕 Tendon Injuries 📕 Sandhoff Disease

Figure 5. Enriched HumanPSD(TM) disease (2020.2) of up-regulated genes in noRA_Dox vs. noRA_noDox. The size of the bars correspond to the number of bio-markers of the given disease found among the input set. Full classification \rightarrow

Down-regulated genes in noRA_Dox vs. noRA_noDox:

726 significant down-regulated genes were taken for the mapping.

GO (biological process)

						biologic	al_proce	ess Gene	e Ontology treer	map						
regulation of cell developmer	neurog	ation of genesis	regulatio axonoge	on of re nesis m	gulation of cell orphogenesis	axonogenes	is m bo F mor	plasma nembrane ounded cel projection <u>phogenes</u>	negative regulation o phosphorylatio	negati f regulat on of prote phosphory	ve ion ein ylation	regulation of developmental process	regulation of cel differentiation	l cell jun organiz	ction ation	synapse organization
regulation of cel morphogenesis involved in differentiation regulation of	regula anato struc morphog regula neuron p develo	tion of mical cture genesis tion of rojection pment	negative regulation of nervous system evelopment negative regulation of neuron projection	positive regulatio of cell developm mathematical and marked mathematical	ent negative regulation of neurogenesis ent negative regulation of neuron differentiation	cell projectio morphogenes neuron project morphogenes	is mor	cell part phogenesi cellular omponent phogenesi	is negative regulation of phosphate metabolic process negative	negativ regulatio <u>MAPK cas</u> regulat	ve in of icade gulation of BdyAyrosine sphorylation	regulation of multicellular organismal development regula	positive regulation of developmental process	cell-substrat junction organizatior	e cell junctio assemb	Cell-substrate Junction assembly
development regulation of neur differentiation	regula ron axonog of cell morp	tion of ne	development positive regulation of ervous syste developmen positive regulation	f regulati m of axo t extensi regulati of dend	re negative regulation of cell projection organization on positive regulation of neurogenesis	axon axon development	ne proj deve	esis euron jection lopment	of phos	phorylat generation of neurons	ion	developme nervous system development	ntal process system der	cell jund	nult mult	icellular Ianism Iopment Cellular
regulation o cholesterol s biosynthetic process	f cell morp secondary alcohol iosynthetic process	biosynthe process	sinerentiatio pri tic loca to me	d in diffe rotein lization embrane	SRP-dependent cotranslational protein targeting to membrane	cell projection organization axon dev	plasma nembrane ounded cel projection rganization	dendrite development	generation embryonic morphogenesis	gastrulatio	ne c ar	ervous syste developmen natomical structu development	t system der re anatomical morphog	velopment structure genesis	org devel negative of nitroge metabo negative	opment e regulation n compound lic process e regulation
cholesterol ci biosynthetic bi process via p desmosterol la	holesterol osynthetic rocess via athosterol	alcohol biosynthet process	cotran tic pro targ to me	slational otein jeting mbrane	protein localization to endoplasmic reticulum	cell morphog involved differentiat	enesis in tion enesis	cell morphogenesis	embryonic orga morphogenesis embryonic mo cell developmen	t neuron developme	devi devi	developmental proce	ss neuron diffe	rentiation	cellular organ	in compound lic process component hization component
cholesterol metabolic process secondary alcohol	organic hydroxy compound metabolic process phenol-containin compound	alcohol metabol process g norepinephr metabolic	ic to me	targeting mbrane	protein targeting to ER	involved in n cell morphog in diffe multicellular organismal homeostasis	euron enesis i rentiatio er home	nvolved on nergy eostasis	cell deve		t dev neg	elopmental proce gative regulation iological process gative regulation iological process	of cellular comp organization biogenes cellular comp organization biogenesi	onent regu n or mult onent regu n or regu s mult	lation of icellular anismal fation of icellular	regulation of biological process
metabolic process sterol metabolic process cholesterol b	organic hydrox compound biosynthetic	small molecul	endopla e prot	simic reticulu tein loc o mem	calization	multi organismal	rena pro cellular homeo	l system ocess	differentiation	development process entiation	al tub	ube developmen	regulation biological qu regulation nt biological qu	of organizative of organizative of organizative of of uality me	regulat tabolic	biological process ion of process

Figure 6. Enriched GO (biological process) of down-regulated genes in noRA_Dox vs. noRA_noDox. Full classification \rightarrow

TRANSPATH® Pathways (2020.2)



Figure 7. Enriched TRANSPATH® Pathways (2020.2) of down-regulated genes in noRA_Dox vs. noRA_noDox. Full classification \rightarrow

HumanPSD(TM) disease (2020.2)



Skin and Connective Tissue Diseases

Female Urogenital Diseases and Pregnancy Complications

Female Urogenital Diseases Respiratory Tract Neoplasms Thoracic Neoplasms

Lung Neoplasms Neoplasms, Germ Cell and Embryonal Neoplasms, Nerve Tissue

Neuroectodermal Tumors Neuroendocrine Tumors Neuroepithelial

🔳 Kidney Neoplasms 📕 Glioma 📕 Leiomyoma

Figure 8. Enriched HumanPSD(TM) disease (2020.2) of down-regulated genes in noRA_Dox vs. noRA_noDox. The size of the bars correspond to the number of bio-markers of the given disease found among the input set. Full classification –

The result of overall Gene Ontology (GO) analysis of the differentially expressed genes of the studied pathology can be summarized by the following diagram, revealing the most significant functional categories overrepresented among the observed (differentially expressed genes):



Up-regulated genes hits Down-regulated genes hits 🗕 Up-regulated genes -log10(P-value) - Down-regulated genes -log10(P-value)

3.3. Analysis of enriched transcription factor binding sites and composite modules

In the next step a search for transcription factors binding sites (TFBS) was performed in the regulatory regions of the target genes by using the TF binding motif library of the TRANSFAC® database. We searched for so called composite modules that act as potential condition-specific enhancers of the target genes in their upstream regulatory regions (-1000 bp upstream of transcription start site (TSS)) and identify transcription factors regulating activity of the genes through such enhancers.

Classically, enhancers are defined as regions in the genome that increase transcription of one or several genes when inserted in either orientation at various distances upstream or downstream of the gene [8]. Enhancers typically have a length of several hundreds of nucleotides and are bound by multiple transcription factors in a cooperative manner [9].

We applied the Composite Module Analyst (CMA) [8] method to detect such potential enhancers, as targets of multiple TFs bound in a cooperative manner to the regulatory regions of the genes of interest. CMA applies a genetic algorithm to construct a generalized model of the enhancers by specifying combinations of TF motifs (from TRANSFAC®) whose sites are most frequently clustered together in the regulatory regions of the studied genes. CMA identifies the transcription factors that through their cooperation provide a synergistic effect and thus have a great influence on the gene regulation process.

Enhancer model potentially involved in regulation of target genes (up-regulated genes in noRA_Dox vs. noRA_noDox).

To build the most specific composite modules we choose genes as the input of CMA algorithm. The obtained CMA model is then applied to compute CMA score for all up-regulated genes.

The model consists of 2 module(s). Below, for each module the following information is shown:

- PWMs producing matches,
- number of individual matches for each PWM,
- score of the best match.





See model visualization table \rightarrow

Table 4. List of top ten up-regulated genes in noRA_Dox vs. noRA_noDox with identified enhancers in their regulatory regions. **CMA score** - the score of the CMA model of the enhancer identified in the regulatory region. **See full table** \rightarrow

Ensembl IDs	Gene symbol	Gene description	CMA score	Factor names
ENSG00000234645	YWHAEP5	tyrosine 3-monooxygenase/tryptophan 5- monooxygenase activation protein epsilon pseudogene 5	12.74	ZEB1(h), POU6F1(h), PBX3(h),PREP- 1(h),Pbx1(h),Pbx2(h), TAFII250(h), POU2F1(h), GR(h),PR(h), HNF-3alpha(h)
ENSG00000256436	TAS2R31	taste 2 receptor member 31	12.42	TAFII250(h), HNF-3alpha(h), Tbx5(h), PBX3(h),PREP-1(h),Pbx1(h),Pbx2(h), ZEB1(h), POU6F1(h), POU2F1(h)
ENSG0000086300	SNX10	sorting nexin 10	12.31	GR(h),PR(h), JunD(h), POU6F1(h), PBX3(h),PREP- 1(h),Pbx1(h),Pbx2(h), ZEB1(h), Tbx5(h), HNF- 3alpha(h)
ENSG00000176857	GJA1P1	gap junction protein alpha 1 pseudogene 1	12.29	HNF-3alpha(h), Tbx5(h), ZEB1(h), PBX3(h),PREP- 1(h),Pbx1(h),Pbx2(h), GR(h),PR(h), POU6F1(h), JunD(h)
ENSG00000115548	KDM3A	lysine demethylase 3A	12.17	HNF-3alpha(h), JunD(h), GR(h),PR(h), POU2F1(h), ZEB1(h), PBX3(h),PREP-1(h),Pbx1(h),Pbx2(h), POU6F1(h)
ENSG00000253203	GUSBP3	GUSB pseudogene 3	11.9	PBX3(h),PREP-1(h),Pbx1(h),Pbx2(h), ZEB1(h), TAFII250(h), JunD(h), POU6F1(h), Tbx5(h), GR(h),PR(h)
ENSG00000235916	AC233279.1	mitochondrial ribosomal protein S18C (MRPS18C) pseudogene	11.8	GR(h),PR(h), ZEB1(h), Tbx5(h), JunD(h), HNF- 3alpha(h), TAFII250(h), PBX3(h),PREP- 1(h),Pbx1(h),Pbx2(h)
ENSG00000174827	PDZK1	PDZ domain containing 1	11.75	TAFII250(h), PBX3(h),PREP-1(h),Pbx1(h),Pbx2(h), ZEB1(h), POU2F1(h), JunD(h), GR(h),PR(h), Tbx5(h)
ENSG00000173889	PHC3	polyhomeotic homolog 3	11.73	JunD(h), Tbx5(h), GR(h),PR(h), ZEB1(h), HNF- 3alpha(h), POU2F1(h), POU6F1(h)
ENSG00000269743	SLC25A53	solute carrier family 25 member 53	11.6	HNF-3alpha(h), ZEB1(h), GR(h),PR(h), TAFII250(h), JunD(h), PBX3(h),PREP-1(h),Pbx1(h),Pbx2(h), POU6F1(h)

Enhancer model potentially involved in regulation of target genes (down-regulated genes in noRA_Dox vs. noRA_noDox).

To build the most specific composite modules we choose genes as the input of CMA algorithm. The obtained CMA model is then applied to compute CMA score for all down-regulated genes.

The model consists of 2 module(s). Below, for each module the following information is shown:

- PWMs producing matches,
- number of individual matches for each PWM,
- score of the best match.

Ν	Module 1:								
	V\$E2A_Q6_02	V\$SREBP_Q3 0.00; N=2	V\$YY1_02 0.00; N=2	V\$SMAD3_Q6 0.88; N=2	V\$CP2_02 0.81; N=3				
	Module width: 124								
Ν	1odule 2:								
	V\$RAD21_10 0.94; N=3	V\$OCT1_04 0.86; N=3	/\$RXRRAR_01	V\$REST_05 0.00; N=3	V\$GR_Q6_02 0.00; N=2				
	Module width: 147								

Model score (-p*log10(pval)): 19.33 Wilcoxon p-value (pval): 2.65e-39 Penalty (p): 0.501 Average yes-set score: 8.89 Average no-set score: 7.15 AUC: 0.78 Middle-point: 8.04 False-positive: 26.20% False-negative: 29.15%



See model visualization table \rightarrow

Table 5. List of top ten down-regulated genes in noRA_Dox vs. noRA_noDox with identified enhancers in their regulatory regions. **CMA score** - the score of the CMA model of the enhancer identified in the regulatory region. **See full table** \rightarrow

Ensembl IDs	Gene symbol	Gene description	CMA score	Factor names
ENSG00000128394	APOBEC3F	apolipoprotein B mRNA editing enzyme catalytic subunit 3F	15.59	Rad21(h), REST(h), GR(h), NR1B1(h),RXR-alpha(h), Smad3(h), CP2(h), SREBP-1(h),SREBP-2(h)
ENSG00000225746	MEG8	maternally expressed 8, small nucleolar RNA host gene	15.32	YY1(h), REST(h), Smad3(h), CP2(h), E2A(h), SREBP- 1(h),SREBP-2(h), Rad21(h)
ENSG00000154358	OBSCN	obscurin, cytoskeletal calmodulin and titin-interacting RhoGEF	15.27	NR1B1(h),RXR-alpha(h), REST(h), YY1(h), GR(h), Rad21(h), CP2(h), SREBP-1(h),SREBP-2(h)
ENSG00000236735	RPL31P63	ribosomal protein L31 pseudogene 63	14.88	Rad21(h), NR1B1(h),RXR-alpha(h), GR(h), REST(h), CP2(h), POU2F1(h), Smad3(h)
ENSG00000166402	TUB	TUB bipartite transcription factor	14.87	E2A(h), SREBP-1(h),SREBP-2(h), Smad3(h), YY1(h), CP2(h), Rad21(h), NR1B1(h),RXR-alpha(h)
ENSG00000214706	IFRD2	interferon related developmental regulator 2	14.83	Rad21(h), CP2(h), SREBP-1(h),SREBP-2(h), REST(h), YY1(h), Smad3(h), E2A(h)
ENSG00000102886	GDPD3	glycerophosphodiester phosphodiesterase domain containing 3	14.74	Smad3(h), SREBP-1(h),SREBP-2(h), YY1(h), CP2(h), Rad21(h), REST(h), E2A(h)
ENSG0000085978	ATG16L1	autophagy related 16 like 1	14.73	SREBP-1(h),SREBP-2(h), CP2(h), YY1(h), Rad21(h), E2A(h), Smad3(h), POU2F1(h)
ENSG00000049540	ELN	elastin	14.66	Smad3(h), GR(h), NR1B1(h),RXR-alpha(h), Rad21(h), REST(h), YY1(h), CP2(h)
ENSG00000128805	ARHGAP22	Rho GTPase activating protein 22	14.64	Rad21(h), NR1B1(h),RXR-alpha(h), E2A(h), GR(h), REST(h), YY1(h), SREBP-1(h),SREBP-2(h)

On the basis of the enhancer models we identified transcription factors potentially regulating the **target genes** of our interest. We found 13 and 12 transcription factors controlling expression of up- and down-regulated genes respectively (see Tables 6-7).

Table 6. Transcription factors of the predicted enhancer model potentially regulating the differentially expressed genes (up-regulated genes in noRA_Dox vs. noRA_noDox). **Yes-No ratio** is the ratio between frequencies of the sites in Yes sequences versus No sequences. It describes the level of the enrichment of binding sites for the indicated TF in the regulatory target regions. **Regulatory score** is the measure of involvement of the given TF in the controlling of expression of genes that encode master regulators presented below (through positive feedback loops). **See full table** \rightarrow

ID	Gene symbol	Gene description	Regulatory score	Yes-No ratio
MO000054297	PGR	progesterone receptor	4.92	1.31
MO000031266	NR3C1	nuclear receptor subfamily 3 group C member 1	4.63	1.31
MO000139677	ZEB1	zinc finger E-box binding homeobox 1	4.56	1.54
MO000025003	POU2F1	POU class 2 homeobox 1	4.53	1.54
MO00007834	JUND	JunD proto-oncogene, AP-1 transcription factor subunit	4.08	1.41
MO000028320	null	null	3.17	1.46
MO000081793	TAF1	TATA-box binding protein associated factor 1	3	1.23
MO000026492	FOXA1	forkhead box A1	0	1.44
MO000028025	PKNOX1	PBX/knotted 1 homeobox 1	0	1.45
MO000028026	PBX2	PBX homeobox 2	0	1.55

Table 7. Transcription factors of the predicted enhancer model potentially regulating the differentially expressed genes (down-regulated genes in noRA_Dox vs. noRA_noDox). **Yes-No ratio** is the ratio between frequencies of the sites in Yes sequences versus No sequences. It describes the level of the enrichment of binding sites for the indicated TF in the regulatory target regions. **Regulatory score** is the measure of involvement of the given TF in the controlling of expression of genes that encode master regulators presented below (through positive feedback loops). **See full table** \rightarrow

ID	Gene symbol	Gene description	Regulatory score	Yes-No ratio
MO000042938	RAD21	RAD21 cohesin complex component	5.32	3.22
MO000025003	POU2F1	POU class 2 homeobox 1	4.99	
MO000078913	YY1	YY1 transcription factor	4.86	1.1
MO000057832	SMAD3	SMAD family member 3	4.8	9.32
MO000041817	REST	RE1 silencing transcription factor	4.55	1.53
MO000019619	RXRA	retinoid X receptor alpha	4.34	1.48
MO000031266	NR3C1	nuclear receptor subfamily 3 group C member 1	4.3	1.87
MO000025765	SREBF2	sterol regulatory element binding transcription factor 2	4.29	5.08
MO000117988	TFCP2	transcription factor CP2	4.24	1.4
MO000056029	SREBF1	sterol regulatory element binding transcription factor 1	4.2	3.39

The following diagram represents the key transcription factors, which were predicted to be potentially regulating differentially expressed genes in the analyzed pathology: PGR, NR3C1, ZEB1, RAD21, POU2F1 and YY1.



3.4. Finding master regulators in networks

In the second step of the upstream analysis common regulators of the revealed TFs were identified. These master regulators appear to be the key candidates for therapeutic targets as they have a master effect on regulation of intracellular pathways that activate the pathological process of our study. The identified master regulators are shown in Tables 8-9. Table 8. Master regulators that may govern the regulation of **up-regulated** genes in noRA_Dox vs. noRA_noDox. **Total rank** is the sum of the ranks of the master molecules sorted by keynode score, CMA score, transcriptomics data. See full table \rightarrow

ID	Master molecule name	Gene symbol	Gene description	logFC	Total rank
MO000021902	TFIIH-CAK(h)	CCNH, CDK7, MNAT1	MNAT1 component of CDK activating kinase, cyclin H, cyclin dependent kinase 7	0.23	115
MO000023445	Cdc25A(h)	CDC25A	cell division cycle 25A	0.44	167
MO000021981	brca1(h)	BRCA1	BRCA1 DNA repair associated	0.31	171
MO000281381	(angiotensin II)2:(AT2 receptor)2:(ATIP- isoform3)2:SHP-1	AGT, AGTR2, MTUS1, PTPN6	angiotensin II receptor type 2, angiotensinogen, microtubule associated scaffold protein 1, protein	0.38	180
MO000117508	TC-PTP(h)	PTPN2	protein tyrosine phosphatase non-receptor type 2	0.3	183
MO000030911	PIAS1(h)	PIAS1	protein inhibitor of activated STAT 1	0.22	190
MO000043863	prlr(h):tec(h):Vav(h)	PRLR, TEC, VAV1	prolactin receptor, tec protein tyrosine kinase, vav guanine nucleotide exchange factor 1	0.52	216
MO000085337	Cdc25A1(h)	CDC25A	cell division cycle 25A	0.44	216
MO000085339	Cdc25A2(h)	CDC25A	cell division cycle 25A	0.44	217
MO000104136	cyclosome(h):Fzr1(h)	ANAPC1, ANAPC10, ANAPC11, ANAPC2, ANAPC4, ANAPC5, ANAPC7, CDC16, CDC23, CDC26, CDC27, FZR1	anaphase promoting complex subunit 1, anaphase promoting complex subunit 10, anaphase promoting comp	0.23	239

Table 9. Master regulators that may govern the regulation of **down-regulated** genes in noRA_Dox vs. noRA_noDox. **Total rank** is the sum of the ranks of the master molecules sorted by keynode score, CMA score, transcriptomics data.

ID	Master molecule name	Gene symbol	Gene description	logFC	Total rank
MO000031101	plk3(h)	PLK3	polo like kinase 3	-0.54	230
MO000281381	(angiotensin II)2:(AT2 receptor)2:(ATIP-isoform3)2:SHP-1	AGT, AGTR2, MTUS1, PTPN6	angiotensin II receptor type 2, angiotensinogen, microtubule associated scaffold protein 1, protein	-1.9	260
MO000165201	mTOR(h):rictor(h):mLST8(h):SIN1(h):Protor-1(h)	MAPKAP1, MLST8, MTOR, PRR5, RICTOR	MAPK associated protein 1, MTOR associated protein, LST8 homolog, RPTOR independent companion of MTO	-0.66	269
MO000138699	plk3(h)	PLK3	polo like kinase 3	-0.54	281
MO000280531	rictor-isoform1(h):mTOR(h):SIN1(h):mLST8(h)	MAPKAP1, MLST8, MTOR, RICTOR	MAPK associated protein 1, MTOR associated protein, LST8 homolog, RPTOR independent companion of MTO	-0.55	295
MO000161481	Cytochrome b- 558:p22phox:p40phox{p}:p67phox:Rac1:GTP:JFC1:PtdIns(3,4)P2:PA:p47phox	CYBA, CYBB, NCF1, NCF2, NCF4, RAC1, SYTL1	Rac family small GTPase 1, cytochrome b- 245 alpha chain, cytochrome b- 245 beta chain, neutrophil cyt	-0.53	302
MO000104136	cyclosome(h):Fzr1(h)	ANAPC1, ANAPC10, ANAPC11, ANAPC2, ANAPC4, ANAPC5, ANAPC7, CDC16, CDC23, CDC23, CDC26, CDC27, FZR1	anaphase promoting complex subunit 1, anaphase promoting complex subunit 10, anaphase promoting comp	-0.54	303
MO000043414	cyclosome(h)	ANAPC1, ANAPC10, ANAPC11, ANAPC2, ANAPC4, ANAPC5, ANAPC7, CDC16, CDC23, CDC26, CDC27	anaphase promoting complex subunit 1, anaphase promoting complex subunit 10, anaphase promoting comp	-0.54	341
MO000017421	AKT-1(h)	AKT1	AKT serine/threonine kinase 1	-0.41	342
MO000031189	PKCdelta(h)	PRKCD	protein kinase C delta	-0.34	342

The intracellular regulatory pathways controlled by the above-mentioned master regulators are depicted in Figures 9 and 10. These diagrams display the connections between identified transcription factors, which play important roles in the regulation of differentially expressed genes, and selected master regulators, which are responsible for the regulation of these TFs.



Figure 9. Diagram of intracellular regulatory signal transduction pathways of up-regulated genes in noRA_Dox vs. noRA_noDox. Master regulators are indicated by red rectangles, transcription factors are blue rectangles, and green rectangles are intermediate molecules, which have been added to the network during the search for master regulators from selected TFs. Orange and blue frames highlight molecules that are encoded by up- and downregulated genes, resp.





Figure 10. Diagram of intracellular regulatory signal transduction pathways of down-regulated genes in noRA_Dox vs. noRA_noDox. Master regulators are indicated by red rectangles, transcription factors are blue rectangles, and green rectangles are intermediate molecules, which have been added to the network during the search for master regulators from selected TFs. Orange and blue frames highlight molecules that are encoded by up- and downregulated genes, resp. See full diagram \rightarrow

4. Finding prospective drug targets

The identified master regulators that may govern pathology associated genes were checked for druggability potential using HumanPSDTM [5] database of gene-disease-drug assignments and PASS [11-13] software for prediction of biological activities of chemical compounds on the basis of a (Q)SAR approach. Respectively, for each master regulator protein we have computed two druggability scores: HumanPSD druggability score and PASS druggability score. Where druggability score represents the number of drugs that are potentially suitable for inhibition (or activation) of the corresponding target either according to the information extracted from medical literature (from HumanPSDTM database) or according to cheminformatics predictions of compounds activity against the examined target (from PASS software).

The cheminformatics druggability check is done using a pre-computed database of spectra of biological activities of chemical compounds from a library of all small molecular drugs from HumanPSD[™] database, 2507 pharmaceutically active known chemical compounds in total. The spectra of biological activities has been computed using the program PASS [11-13] on the basis of a (Q)SAR approach.

If both druggability scores were below defined thresholds (see Method section for the details) such master regulator proteins were not used in further analysis of drug prediction.

As a result we created the following two tables of prospective drug targets (top targets are shown here):

Table 10. Prospective drug targets selected from full list of identified master regulators filtered by druggability score from HumanPSD[™] database. **Druggability score** contains the number of drugs that are potentially suitable for inhibition (or activation) of the target. The drug targets are sorted according to the **Total rank** which is the sum of three ranks computed on the basis of the three scores: keynode score, CMA score and expression change score (logFC, if present). See Methods section for details.

Gene symbol	Gene Description	Druggability score	logFC	Total rank
PRLR	prolactin receptor	2	0.52	276
PRKCQ	protein kinase C theta	3	0.32	314
ITGA4	integrin subunit alpha 4	8	0.22	420
ITGAV	integrin subunit alpha V	1	0.22	420
KAT2B	lysine acetyltransferase 2B	3	0.41	482
SETD7	SET domain containing 7, histone lysine methyltransferase	1	0.17	492

Table 11. Prospective drug targets selected from full list of identified master regulators filtered by druggability score predicted by PASS software. Here, the **druggability score** for master regulator proteins is computed as a sum of PASS calculated probabilities to be active as a target for various small molecular compounds. The drug targets are sorted according to the **Total rank** which is the sum of three ranks computed on the basis of the three scores: keynode score, CMA score and expression change score (logFC, if present). See Methods section for details.

See full table \rightarrow

Gene symbol	Gene Description	Druggability score	logFC	Total rank
MNAT1	MNAT1 component of CDK activating kinase	469.58	0.23	115
CCNH	cyclin H	15.76	0.23	115
NEK10	NIMA related kinase 10	22.89	0.2	242
PRKCQ	protein kinase C theta	196.29	0.32	314
MAP2K4	mitogen-activated protein kinase kinase 4	123.27	0.17	388
CLK4	CDC like kinase 4	27.7	0.38	392

Below we represent schematically the main mechanism of the studied pathology. In the schema we considered the top two drug targets of each of the two categories computed above. In addition we have added two top identified master regulators for which no drugs may be identified yet, but that are playing the crucial role in the molecular mechanism of the studied pathology. Thus the molecular mechanism of the studied pathology was predicted to be mainly based on the following key master regulators:

- TFIIH-CAK
- PKCtheta
- prlr:tec:Vav
- Cdc25A

This result allows us to suggest the following schema of affecting the molecular mechanism of the studied pathology:



Drugs which are shown on this schema: Phosphonothreonine, D-Myo-Inositol-Hexasulphate, 2,4-Dihydroxybenzoic Acid and Fluoxymesterone, should be considered as a prospective research initiative for further drug repurposing and drug development. These drugs were selected as top matching treatments to the most prospective drug targets of the studied pathology, however, these results should be considered with special caution and are to be used for research purposes only, as there is not enough clinical information for adapting these results towards immediate treatment of patients.

The drugs given in dark red color on the schema are FDA approved drugs or drugs which have gone through various phases of clinical trials as active treatments against the selected targets.

The drugs given in pink color on the schema are drugs, which were cheminformatically predicted to be active against the selected targets.

5. Identification of potential drugs

In the last step of the analysis we strived to identify known activities as well as drugs with cheminformatically predicted activities that are potentially suitable for inhibition (or activation) of the identified molecular targets in the context of specified human diseases(s).

Proposed drugs are top ranked drug candidates, that were found to be active on the identified targets and were selected from 4 categories:

- 1. FDA approved drugs or used in clinical trials drugs for the studied pathology;
- 2. Repurposing drugs used in clinical trials for other pathologies;
- 3. Drugs, predicted by PASS to be active against identified drug targets and against the studied pathology;
- 4. Drugs, predicted by PASS to be active against identified drug targets but for other pathologies.

Proposed drugs were selected on the basis of drug rank which was computed from two scores:

- target activity score (depends on ranks of all targets that were found for the selected drug);
- disease activity score (weighted sum of number of clinical trials on disease(s) under study where the selected drug is known to be applied or PASS disease activity score - cheminformatically predicted property of the compound to be active against the studied disease(s)).

You can refer to the Methods section for more details on drug ranking procedure.

Top drugs of each category are given in the tables below:

Drugs approved in clinical trials



Table 12. FDA approved drugs or drugs used in clinical trials for the studied pathology (most promising treatment candidates selected for the identified drug targets on the basis of literature curation in HumanPSD^m database) See full table \rightarrow

Name	Target names	Drug rank	Disease activity score	Phase 4	Status (provided by Drugbank)
Acetylcysteine	ІКВКВ, СНИК	98	3	Acute Kidney Injury, Alcoholism, Anemia, Atherosclerosis, Atrophy, Bipolar Disorder, Bronchiectasis	small molecule, approved
Hydrocortisone	NR3C1	113	3	Adrenal Insufficiency, Asthma, Bites and Stings, Burns, Candidiasis, Cicatrix, Cicatrix, Hypertrophic	small molecule, approved
Lovastatin	HDAC2	198	2	Dyslipidemias, Fragile X Syndrome, Genetic Diseases, Inborn, Hyperlipidemias	small molecule,approved,investigational
Fludrocortisone	NR3C1	242	1	Hypertension, Syncope, Syncope, Vasovagal	small molecule, approved
Naloxone	TLR4	258	3	Angina Pectoris, Angina, Unstable, Arthritis, Bursitis, Constipation, Cysts, Depression	small molecule, approved

Repurposing drugs

Table 13. Repurposed drugs used in clinical trials for other pathologies (prospective drugs against the identified drug targets on the basis of literature curation in HumanPSD[™] database)

Name	Target names	Drug rank	Phase 4	Status (provided by Drugbank)
Ruxolitinib	JAK2, JAK1	19	Splenomegaly	small molecule, approved
Vedolizumab	ITGA4	20	Colitis, Colitis, Ulcerative, Crohn Disease, Ulcer	biotech, approved
Peginterferon alfa-2a	IFNAR1, IFNAR2	26	HIV Infections, Hemophilia A, Hepatitis, Hepatitis B, Hepatitis B, Chronic, Hepatitis C, Hepatitis C, Chronic	biotech,approved,investigational
Peginterferon alfa-2b	IFNAR1, IFNAR2	26	Hepatitis, Hepatitis B, Hepatitis B, Chronic, Hepatitis C, Hepatitis C, Chronic, Hepatitis, Chronic	biotech,approved
Interferon beta-1a	IFNAR1, IFNAR2	26	Brain Abscess, Multiple Sclerosis, Multiple Sclerosis, Relapsing- Remitting	biotech,approved,investigational



Table 14. Prospective drugs, predicted by PASS software to be active against the identified drug targets with predicted activity against the studied disease(s) (drug candidates predicted with the cheminformatics tool PASS) See full table \rightarrow

Name	Target names	Drug rank	Target activity score
D-Myo-Inositol-Hexasulphate	TEC, ADAM17, PTPRO, PTPRJ, TRIM36, MNAT1, USP44	418	0.84
6-hydroxydopa quinone	TEC, PTPRO, PTPRJ, TRIM36, USP44, TRAF6, TXN	431	0.88
2,5-DI-(TERT- BUTYL)-1,4,BENZOHYDROQUINONE	TEC, PTPRO, PTPRJ, HGF, TRIM36, USP44, TRAF6	486	0.88
2-tert-butylbenzene-1,4-diol	TEC, PTPRO, PTPRJ, HGF, TRIM36, USP44, TRAF6	509	0.88
Alpha-L-Fucose	TEC, ADAM17, HGF, MNAT1, TRIM36, USP44, CBLB	509	0.78



Table 15. Prospective drugs, predicted by PASS software to be active against the identified drug targets, though without cheminformatically predicted activity against the studied disease(s) (drug candidates predicted with the cheminformatics tool PASS) See full table \rightarrow

Name	Target names	Drug rank	Target activity score
2,4-Dihydroxybenzoic Acid	TEC, PTPRO, PTPRJ, HGF, TRIM36, USP44, TRAF6	457	1.19
L-Aspartic Acid	TEC, PTPRO, PTPRJ, TRIM36, USP44, TRAF6, CD72	461	0.97
D-Aspartic Acid	TEC, PTPRO, PTPRJ, TRIM36, USP44, TRAF6, CD72	461	0.97
2-(3,6-DIHYDROXYPHENYL)ACETIC ACID	TEC, PTPRO, PTPRJ, TRIM36, USP44, TRAF6, TXN	472	1.08
Tricarballylic Acid	TEC, PTPRO, PTPRJ, TRIM36, USP44, TRAF6, CD72	495	1

As the result of drug search we propose the following drugs as most promising candidates for treating the pathology under study: Acetylcysteine, Ruxolitinib, D-Myo-Inositol-Hexasulphate and 2,4-Dihydroxybenzoic Acid. These drugs were selected for acting on the following targets: IKBKB, JAK2 and MNAT1, which were predicted to be active in the molecular mechanism of the studied pathology.

The selected drugs are top ranked drug candidates from each of the four categories of drugs: (1) FDA approved drugs or used in clinical trials drugs for the studied pathology; (2) repurposing drugs used in clinical trials for other pathologies; (3) drugs, predicted by PASS software to be active against the studied pathology; (4) drugs, predicted by PASS software to be repurposed from other pathologies.

6. Conclusion

We applied the software package "Genome Enhancer" to a data set that contains *transcriptomics* data. The study is done in the context of *Parkinson Disease*. The data were pre-processed, statistically analyzed and differentially expressed genes were identified. Also checked was the enrichment of GO or disease categories among the studied gene sets.

We propose the following drugs as most promising candidates for treating the pathology under study:



These drugs were selected for acting on the following targets: IKBKB, JAK2 and MNAT1, which were predicted to be involved in the molecular mechanism of the pathology under study.

The identified molecular mechanism of the studied pathology was predicted to be mainly based on the following key drug targets:



These potential drug targets should be considered as a prospective research initiative for further drug repurposing and drug development purposes. The following drugs were predicted as, matching those drug targets: Phosphonothreonine, D-Myo-Inositol-Hexasulphate, 2,4-Dihydroxybenzoic Acid and Fluoxymesterone. These drugs should be considered with special caution for research purposes only.

In this study, we came up with a detailed signal transduction network regulating differentially expressed genes in the studied pathology. In this network we have revealed the following top master regulators (signaling proteins and their complexes) that play a crucial role in the molecular mechanism of the studied pathology, which can be proposed as the most promising molecular targets for further drug repurposing and drug development initiatives.

- TFIIH-CAK
- PKCtheta
- prlr:tec:Vav
- Cdc25A

Potential drug compounds which can be affecting these targets can be found in the "Finding prospective drug targets" section.

7. Methods

Databases used in the study

Transcription factor binding sites in promoters and enhancers of differentially expressed genes were analyzed using known DNAbinding motifs described in the TRANSFAC® library, release 2020.2 (geneXplain GmbH, Wolfenbüttel, Germany) (https://genexplain.com/transfac).

The master regulator search uses the TRANSPATH® database (BIOBASE), release 2020.2 (geneXplain GmbH, Wolfenbüttel, Germany) (https://genexplain.com/transpath). A comprehensive signal transduction network of human cells is built by the software on the basis of reactions annotated in TRANSPATH®.

The information about drugs corresponding to identified drug targets and clinical trials references were extracted from HumanPSD[™] database, release 2020.2 (https://genexplain.com/humanpsd).

The Ensembl database release Human99.38 (hg38) (http://www.ensembl.org) was used for gene IDs representation and Gene Ontology (GO) (http://geneontology.org) was used for functional classification of the studied gene set.

Genomic data processing

When analyzing a list of genomic variations (from vcf file or computed by Genome Enhancer from fastq files), first of all, we compute a specific mutation weight (w) for each variation depending on it's location in gene body and gene flanking regions (-1000 upstream and +1000 downstream of the gene body).

- w = 0.7 for variations in exon area
- w = 1.3 for variations in promoter region (-1000bp upstream and 100bp downstream of TSS),
- w = 1.0 for variations in other locations.

Total Gene mutation weight is the sum of the weights w of all variations located inside the gene body and in the gene flanking regions.

Next, a weighted score is calculated for all genes with the following formula: Weighted score = $In_disease * In_transpath * Gene mutation weight, where$

- In_disease = 1.5 for genes assigned to selected diseases,
- In_transpath = 2.0 for genes mapped to Transpath pathways,
- and In_disease = In_transpath = 1.0 in all other cases.

At the next step, 300 genes with highest weighted score are selected for further CMA model search.

The mutation weights (w) are also used to find the regulatory regions of the genes most affected by the variations. A sliding window of 1100 bp is used to scan through the intronic, 5' and 3' regions of the genes and a region is selected with the highest sum of the mutation weights.

Methods for the analysis of enriched transcription factor binding sites and composite modules

Transcription factor binding sites in promoters and enhancers of differentially expressed genes were analyzed using known DNAbinding motifs. The motifs are specified using position weight matrices (PWMs) that give weights to each nucleotide in each position of the DNA binding motif for a transcription factor or a group of them.

We search for transcription factor binding sites (TFBS) that are enriched in the promoters and enhancers under study as compared to a background sequence set such as promoters of genes that were not differentially regulated under the condition of the experiment. We denote study and background sets briefly as Yes and No sets. In the current work we used a workflow considering promoter sequences of a standard length of 1100 bp (-1000 to +100). The error rate in this part of the pipeline is controlled by estimating the adjusted p-value (using the Benjamini-Hochberg procedure) in comparison to the TFBS frequency found in randomly selected regions of the human genome (adj.p-value < 0.01).

We have applied the CMA algorithm (Composite Module Analyst) for searching composite modules [7] in the promoters and enhancers of the Yes and No sets. We searched for a composite module consisting of a cluster of 10 TFs in a sliding window of 200-300 bp that statistically significantly separates sequences in the Yes and No sets (minimizing Wilcoxon p-value).

Methods for finding master regulators in networks

We searched for master regulator molecules in signal transduction pathways upstream of the identified transcription factors. The master regulator search uses a comprehensive signal transduction network of human cells. The main algorithm of the master regulator search has been described earlier [3,4]. The goal of the algorithm is to find nodes in the global signal transduction network that may potentially regulate the activity of a set of transcription factors found at the previous step of the analysis. Such nodes are considered as most promising drug targets, since any influence on such a node may switch the transcriptional programs of hundreds of genes that are regulated by the respective TFs. In our analysis, we have run the algorithm with a maximum radius of 12 steps upstream of each TF in the input set. The error rate of this algorithm is controlled by applying it 10000 times to randomly generated sets of input transcription factors of the same set-size. Z-score and FDR value of ranks are calculated then for each potential master regulator node on the basis of such random runs (see detailed description in [9]). We control the error rate by the FDR threshold 0.05.

Methods for analysis of pharmaceutical compounds

We seek for the optimal combination of molecular targets (key elements of the regulatory network of the cell) that potentially interact with pharmaceutical compounds from a library of known drugs and biologically active chemical compounds, using information about known drugs from HumanPSD[™] and predicting potential drugs using PASS program.

Method for analysis of known pharmaceutical compounds

We selected compounds from HumanPSDTM database that have at least one target. Next, we sort compounds using "*Drug rank*" that is sum of two other ranks:

1. ranking by "Target activity score" (*T*-score_{PSD}),

2. ranking by "Disease activity score" (*D*-score_{PSD}).

"Target activity score" (*T*-score_{PSD}) is calculated as follows:

$$T\text{-}score_{PSD} = -\frac{|T|}{|T| + w(|AT| - |T|))} \sum_{t \in T} log_{10} \left(\frac{rank(t)}{1 + maxRank(T)}\right),$$

where *T* is set of all targets related to the compound intersected with input list, |T| is number of elements in *T*, *AT* and |AT| are set set of all targets related to the compound and number of elements in it, *w* is weight multiplier, rank(t) is rank of given target, maxRank(T) equals max(rank(t)) for all targets *t* in *T*.

We use following formula to calculate "Disease activity score" (D-score_{PSD}):

$$D\text{-}score_{PSD} = \begin{cases} \sum_{d \in D} \sum_{p \in P} phase(d, p) \\ 0, D = \emptyset \end{cases}$$

where *D* is the set of selected diseases, and if *D* is empty set, D-score_{*PSD*}=0. *P* is a set of all known phases for each disease, *phase*(*p*,*d*) equals to the phase number if there are known clinical trials for the selected disease on this phase and zero otherwise.

Method for prediction of pharmaceutical compounds

In this study, the focus was put on compounds with high pharmacological efficiency and low toxicity. For this purpose, comprehensive library of chemical compounds and drugs was subjected to a SAR/QSAR analysis. This library contains 13040 compounds along with their pre-calculated potential pharmacological activities of those substances, their possible side and toxic effects, as well as the possible mechanisms of action. All biological activities are expressed as probability values for a substance to exert this activity (*Pa*). We selected compounds that satisfied the following conditions:

- 1. Toxicity below a chosen toxicity threshold (defines as Pa, probability to be active as toxic substance).
- 2. For all predicted pharmacological effects that correspond to a set of user selected disease(s) *Pa* is greater than a chosen effect threshold.
- 3. There are at least 2 targets (corresponding to the predicted activity-mechanisms) with predicted *Pa* greater than a chosen target threshold.

The maximum *Pa* value for all toxicities corresponding to the given compound is selected as the "Toxicity score". The maximum *Pa* value for all activities corresponding to the selected diseases for the given compound is used as the "Disease activity score". "Target activity score" (T-score) is calculated as follows:

$$T\text{-}score(s) = \frac{|T|}{|T| + w(|AT| - |T|))} \sum_{m \in M(s)} \left(pa(m) \sum_{g \in G(m)} IAP(g)optWeight(g) \right),$$

where M(s) is the set of activity-mechanisms for the given structure (which passed the chosen threshold for activity-mechanisms

Pa); *G*(*m*) is the set of targets (converted to genes) that corresponds to the given activity-mechanism (m) for the given compound; pa(m) is the probability to be active of the activity-mechanism (m), IAP(g) is the invariant accuracy of prediction for gene from G(m); optWeight(g) is the additional weight multiplier for gene. *T* is set of all targets related to the compound intersected with input list, |T| is number of elements in *T*, *AT* and |AT| are set set of all targets related to the compound and number of elements in it, *w* is weight multiplier.

"Druggability score" (D-score) is calculated as follows:

$$D\text{-}score(g) = IAP(g) \sum_{s \in S(g)} \sum_{m \in M(s,g)} pa(m),$$

where S(g) is the set of structures for which target list contains given target, M(s,g) is the set of activity-mechanisms (for the given structure) that corresponds to the given gene, pa(m) is the probability to be active of the activity-mechanism (m), IAP(g) is the invariant accuracy of prediction for the given gene.

8. References

- 1. Kel A, Voss N, Jauregui R, Kel-Margoulis O, Wingender E. Beyond microarrays: Finding key transcription factors controlling signal transduction pathways. *BMC Bioinformatics*. **2006**;7(S2), S13. doi:10.1186/1471-2105-7-s2-s13
- Stegmaier P, Voss N, Meier T, Kel A, Wingender E, Borlak J. Advanced Computational Biology Methods Identify Molecular Switches for Malignancy in an EGF Mouse Model of Liver Cancer. *PLoS ONE.* 2011;6(3):e17738. doi:10.1371/journal.pone.0017738
- 3. Koschmann J, Bhar A, Stegmaier P, Kel A, Wingender E. "Upstream Analysis": An Integrated Promoter-Pathway Analysis Approach to Causal Interpretation of Microarray Data. *Microarrays.* **2015**;4(2):270-286. doi:10.3390/microarrays4020270.
- 4. Kel A, Stegmaier P, Valeev T, Koschmann J, Poroikov V, Kel-Margoulis OV, and Wingender E. Multi-omics "upstream analysis" of regulatory genomic regions helps identifying targets against methotrexate resistance of colon cancer. *EuPA Open Proteom.* **2016**;13:1-13. doi:10.1016/j.euprot.2016.09.002
- 5. Michael H, Hogan J, Kel A et al. Building a knowledge base for systems pathology. *Brief Bioinformatics*. **2008**;9(6):518-531. doi:10.1093/bib/bbn038
- Matys V, Kel-Margoulis OV, Fricke E, Liebich I, Land S, Barre-Dirrie A, Reuter I, Chekmenev D, Krull M, Hornischer K, Voss N, Stegmaier P, Lewicki-Potapov B, Saxel H, Kel AE, Wingender E. TRANSFAC and its module TRANSCompel: transcriptional gene regulation in eukaryotes. *Nucleic Acids Res.* 2006;34(90001):D108-D110. doi:10.1093/nar/gkj143
- Kel AE, Gössling E, Reuter I, Cheremushkin E, Kel-Margoulis OV, Wingender E. MATCH: A tool for searching transcription factor binding sites in DNA sequences. *Nucleic Acids Res.* 2003;31(13):3576-3579. doi:10.1093/nar/gkg585
- Waleev T, Shtokalo D, Konovalova T, Voss N, Cheremushkin E, Stegmaier P, Kel-Margoulis O, Wingender E, Kel A. Composite Module Analyst: identification of transcription factor binding site combinations using genetic algorithm. *Nucleic Acids Res.* 2006;34(Web Server issue):W541-5.
- Krull M, Pistor S, Voss N, Kel A, Reuter I, Kronenberg D, Michael H, Schwarzer K, Potapov A, Choi C, Kel-Margoulis O, Wingender E. TRANSPATH: an information resource for storing and visualizing signaling pathways and their pathological aberrations. *Nucleic Acids Res.* 2006;34(90001):D546-D551. doi:10.1093/nar/gkj107
- Boyarskikh U, Pintus S, Mandrik N, Stelmashenko D, Kiselev I, Evshin I, Sharipov R, Stegmaier P, Kolpakov F, Filipenko M, Kel A. Computational master-regulator search reveals mTOR and PI3K pathways responsible for low sensitivity of NCI-H292 and A427 lung cancer cell lines to cytotoxic action of p53 activator Nutlin-3. *BMC Med Genomics.* **2018**;11(1):12. doi:10.1186/1471-2105-7-s2-s13
- 1. Filimonov D, Poroikov V. Probabilistic Approaches in Activity Prediction. Varnek A, Tropsha A. *Cheminformatics Approaches to Virtual Screening.* Cambridge (UK): RSC Publishing. **2008**;:182-216.
- 2. Filimonov DA, Poroikov VV. Prognosis of specters of biological activity of organic molecules. *Russian chemical journal.* **2006**;50(2):66-75 (russ)
- 3. Filimonov D, Poroikov V, Borodina Y, Gloriozova T. Chemical Similarity Assessment Through Multilevel Neighborhoods of Atoms: Definition and Comparison with the Other Descriptors. *ChemInform.* **1999**;39(4):666-670. doi:10.1002/chin.199940210

Thank you for using the Genome Enhancer!

In case of any questions please contact us at support@genexplain.com

Supplementary material

- 1. Supplementary table 1 Up-regulated genes
- 2. Supplementary table 2 Down-regulated genes
- **3.** Supplementary table 3 Detailed report. Composite modules and master regulators (up-regulated genes in noRA_Dox vs. noRA_noDox).
- 4. Supplementary table 4 Detailed report. Composite modules and master regulators (down-regulated genes in noRA_Dox vs. noRA_noDox).

Disclaimer

Decisions regarding care and treatment of patients should be fully made by attending doctors. The predicted chemical compounds listed in the report are given only for doctor's consideration and they cannot be treated as prescribed medication. It is the physician's responsibility to independently decide whether any, none or all of the predicted compounds can be used solely or in combination for patient treatment purposes, taking into account all applicable information regarding FDA prescribing recommendations for any therapeutic and the patient's condition, including, but not limited to, the patient's and family's medical history, physical examinations, information from various diagnostic tests, and patient preferences in accordance with the current standard of care. Whether or not a particular patient will benefit from a selected therapy is based on many factors and can vary significantly.

The compounds predicted to be active against the identified drug targets in the report are not guaranteed to be active against any particular patient's condition. GeneXplain GmbH does not give any assurances or guarantees regarding the treatment information and conclusions given in the report. There is no guarantee that any third party will provide a refund for any of the treatment decisions made based on these results. None of the listed compounds was checked by Genome Enhancer for adverse side-effects or even toxic effects.

The analysis report contains information about chemical drug compounds, clinical trials and disease biomarkers retrieved from the HumanPSD[™] database of gene-disease assignments maintained and exclusively distributed worldwide by geneXplain GmbH. The information contained in this database is collected from scientific literature and public clinical trials resources. It is updated to the best of geneXplain's knowledge however we do not guarantee completeness and reliability of this information leaving the final checkup and consideration of the predicted therapies to the medical doctor.

The scientific analysis underlying the Genome Enhancer report employs a complex analysis pipeline which uses geneXplain's proprietary Upstream Analysis approach, integrated with TRANSFAC® and TRANSPATH® databases maintained and exclusively distributed worldwide by geneXplain GmbH. The pipeline and the databases are updated to the best of geneXplain's knowledge and belief, however, geneXplain GmbH shall not give a warranty as to the characteristics or to the content and any of the results produced by Genome Enhancer. Moreover, any warranty concerning the completeness, up-to-dateness, correctness and usability of Genome Enhancer information and results produced by it, shall be excluded.

The results produced by Genome Enhancer, including the analysis report, severely depend on the quality of input data used for the analysis. It is the responsibility of Genome Enhancer users to check the input data quality and parameters used for running the Genome Enhancer pipeline.

Note that the text given in the report is not unique and can be fully or partially repeated in other Genome Enhancer analysis reports, including reports of other users. This should be considered when publishing any results or excerpts from the report. This restriction refers only to the general description of analysis methods used for generating the report. All data and graphics referring to the concrete set of input data, including lists of mutated genes, differentially expressed genes/proteins/metabolites, functional classifications, identified transcription factors and master regulators, constructed molecular networks, lists of chemical compounds and reconstructed model of molecular mechanisms of the studied pathology are unique in respect to the used input data set and Genome Enhancer pipeline parameters used for the current run.