# Sequence and Pathway analysis

# FCGR1A and CBL are promising druggable targets for treating Hepatitis C that control activity of IRF7, SMAD5 and SPI1 transcription factor on promoters of differentially expressed genes in liver tissue

Demo User geneXplain GmbH info@genexplain.com Data received on 13/08/2019 ; Run on 11/06/2020 ; Report generated on 11/06/2020

Genome Enhancer release 2.0 (TRANSFAC®, TRANSPATH® and HumanPSD<sup>™</sup> release 2020.2)



## Abstract

In the present study we applied the software package "Genome Enhancer" to a data set that contains *transcriptomics* data obtained from *liver* tissue. The study is done in the context of *Hepatitis C*. 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: IRF7, SMAD5 and SPI1. The subsequent network analysis suggested

- FcgammaRI
- LPS:lbp:CD14:TLR4:MD-2:MyD88:IRAK-1{pS376}{pT387}
- IL-1beta-p17:IL-1RI:IL-1RAcP:MyD88:tollip:IRAK-1{pS376}{pT387}:IRAK-4:IRAK-2
- BGPI
- c-Cbl

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: Sorafenib, Abciximab and Alpha-Ketomalonic 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<sup>™</sup> 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 from HumanPSD<sup>™</sup> 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 used in the study

File name	Data type
E01_Transcriptomics_LogFC-Table	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 analyzed the following condition: Experiment.

# 3.1. Identification of target genes

In the first step of the analysis **target genes** were identified from the uploaded experimental data. Genes were ranked according to the expression value and 300 genes with highest value (see Table 2) and 300 genes with lowest value (see Table 3) were selected for further analysis.

Table 2. Top ten high expressed genes in Experiment. See full table  $\rightarrow$ 

ID	Gene description	Gene symbol	LogFoldChange
ENSG00000137959	interferon induced protein 44 like	IFI44L	6.19
ENSG00000169245	C-X-C motif chemokine ligand 10	CXCL10	6.02
ENSG0000134321	radical S-adenosyl methionine domain containing 2	RSAD2	5.97
ENSG00000137965	interferon induced protein 44	IFI44	3.78
ENSG0000133106	epithelial stromal interaction 1	EPSTI1	3.77
ENSG00000185745	interferon induced protein with tetratricopeptide repeats 1	IFIT1	3.71
ENSG00000187608	ISG15 ubiquitin like modifier	ISG15	3.63
ENSG00000185201	interferon induced transmembrane protein 2	IFITM2	3.54
ENSG0000185885	interferon induced transmembrane protein 1	IFITM1	3.54
ENSG00000135114	2'-5'-oligoadenylate synthetase like	OASL	3.48

ID	Gene description	Gene symbol	LogFoldChange
ENSG0000167910	cytochrome P450 family 7 subfamily A member 1	CYP7A1	-1.09
ENSG00000169282	potassium voltage-gated channel subfamily A member regulatory beta subunit 1	KCNAB1	-1.04
ENSG0000171560	fibrinogen alpha chain	FGA	-0.98
ENSG00000152133	G-patch domain containing 11	GPATCH11	-0.96
ENSG00000182372	CLN8 transmembrane ER and ERGIC protein	CLN8	-0.91
ENSG00000130649	cytochrome P450 family 2 subfamily E member 1	CYP2E1	-0.88
ENSG00000253327	RAD21 antisense RNA 1	RAD21- AS1	-0.88
ENSG00000170323	fatty acid binding protein 4	FABP4	-0.87
ENSG0000175390	eukaryotic translation initiation factor 3 subunit F	EIF3F	-0.86
ENSG00000261609	gigaxonin	GAN	-0.8

# 3.2. Functional classification of genes

A functional analysis of differentially expressed genes was done by mapping the top high expressed and top low expressed genes to several known ontologies, such as Gene Ontology (GO), disease ontology (based on HumanPSD<sup>™</sup> database) and the ontology of signal transduction and metabolic pathways from the TRANSPATH® database. Statistical significance was computed using a binomial test.

Figures 2-7 show the most significant categories.

## **High expressed genes in Experiment:**

300 top high expressed genes were taken for the mapping.

#### GO (biological process)

					biologica	al_process G	ene Ontolo	ogy treemap					
regulation of cytokine production	regulation of type I interferon production	positive regulation of cytokine production	negative regulation of cytokine production regulation of interleukin-1	regulation of Interleukin-1 beta production	cytokine-met signaling pa		rierkro-ganno-melakol signalog patroay	cellular response to type I response to type I inter		viral life cy	cle viral proces	interfe	ponse to ron-gamma
regulation of type I interferon regulation of type I production regulation of type regulation of type regulation of type regulation	ukin-6 tumor n fac produ tive regula ion of of	ecrosis lor interfero ton fion fion fion fion fion fion fion fi	pointive regulation of turner recreate factor speakering cytosize production n of regulation	positive egulation of terleukin-1 beta production positive egulation of nor necrosis	type I interferon sign		tumor necrosis factor-mediated signaling	cellular response to type 1		viral l	iotic process	interfe resp interfer	response to ron-gamma ponse to ron-gamma
superfamily cylokine interferon production produ regulation of interferon-gamma production produ regulation of positi	ction bet ion of regulation on-beta ction prostation	a secretion ton regulation perma tan of protein secretion positive regulation of	regulation of secretion secretion regulation of cytokine secretion regulation of peptide of proteir	regulation of Interferon-bata biosynthetic process	cytokine-mediate defense response to virus		g pathway se to virus	cellular response to organic substance response to	regula of defe respoi	nse of	of immune effector process	readaular readiant d to o non sequence f	itive regulation f NF-kappaB transcription actor activity regulation of DNA-binding
regulation of	tion of interteror ukin-6 ction ation positi pokine regulati	ton of secretion of secretion by cell on of	Interfeukin-1 secretion producti of inter	oduction	defense res	ponse t	o virus	organic substance response to organic substance regulation positive			regulation of process regulation of the effector pro	oduction of ular mediator une response t	transcription tive regulation f NF-kappaB ranscription actor activity response to interferon-alph
viral life cycle	-	iral life cycle	symbios encompas mutualism ti parasiti	sis, ssing hrough	response to biotic stimulus to st		n of se e to host ation of response patho patho patho	seregulation <sup>F kappaB</sup> of I-kappaB kinase/NF -kappaB signaling	response	to stress		riral genom replication regulation	response to interferon-alph
regulation of viral process	of	itive regulatio viral genome replication	of viral entry into host cell	by by symbiont of entry into host	regulation of	mune positive	response s by host regulation C Stimulus stem process	response to external stimulus response to	interfero respoi	on-beta to nse to	f response o stimulus	of response to stimulus regulation of response to stimulu	se response to
negative regulatic of viral process	geno	ulation of viral ome replicatio	n regulation by resultation to via the symbol of the symbo	negative guiation of rai release im host cell negative egulation of viral entry to host cell				external stimulus type I interferon-alpha production interferon-beta	cell respoi chen	ular nse to	I-kappaB nase/NF-kappaB signaling response to stimulus esponse to	regulatio	atory Se on of
regula	ation o	t viral l	ife cycle	gulation of ral release	Immune	e respoi	nse	type I interferon production	stim	ulus	stimulus	proce	SS signal transductio

Figure 2. Enriched GO (biological process) of high expressed genes in Experiment. **Full classification**  $\rightarrow$ 

# TRANSPATH® Pathways (2020.2)

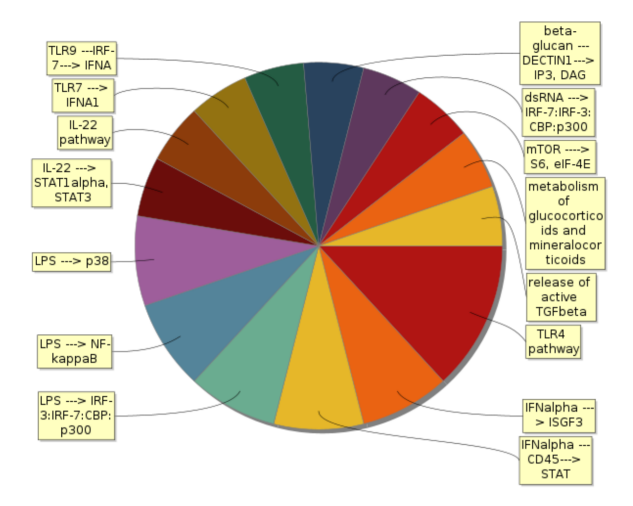
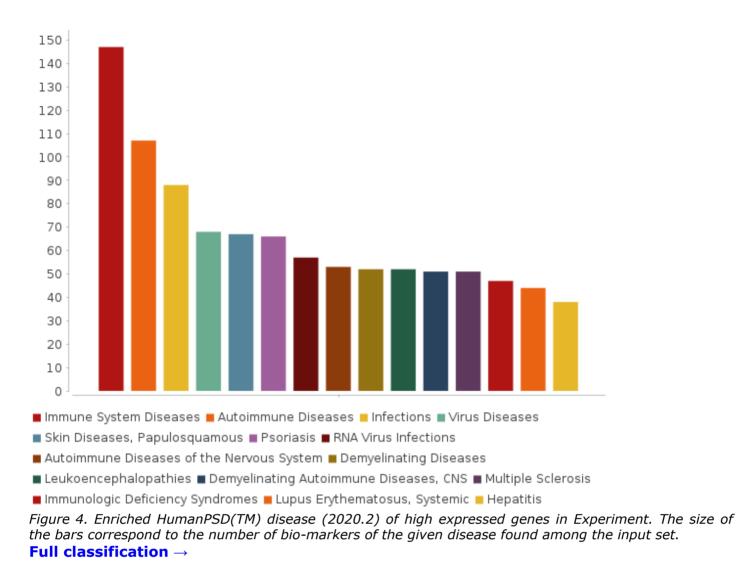


Figure 3. Enriched TRANSPATH® Pathways (2020.2) of high expressed genes in Experiment. Full classification  $\rightarrow$ 

#### HumanPSD(TM) disease (2020.2)



#### Low expressed genes in Experiment:

300 top low expressed genes were taken for the mapping.

#### GO (biological process)

						iological_p	process G	ene Ontoic	gy treemap	'					
alpha-amino acid metabolic process	cellular amino acid metabolic process	cellular amino acio catabolic process	branched-chain amino acid catabolic process	branched-chain amino acid metabolic process	response to organonitroge compound	n nitro	nse to ogen oound	response to hormone	cellular glucuronida	tion meta	ronate ibolic cess	generation of precursor metabolites and energy	f energy derivation by oxidation of organic	cellular amide metabolic process	amide biosynthet process
carboxylic acid catabolic process	small molecule catabolic process	tyrosine metabolic process	cellular amino acid biosynthetic process	sulfur compound metabolic process	response to endogenous stimulus	cellular response to endogenou stimulus		to cellular response to peptide hormone stimulus	uronic acid metabolic process	glucuronidation <sup>a</sup>	arbohydrate metabolic	respiration r	energy glycogen reserve metabolic process	peptide trans biosynthetic process	lation peptid metabo proces
organic acid catabolic process	aromatic amino acid family biosynthetic	serine family amino acid metabolic process	acid family metabolic process	ing. harrocysteine	cellular response to organonitrogen compound cellular	cellular response nitrogen compound response	to d peptid	se to peptide	cellular cellular hormone metabolic	metabolic lucuronic androgen metabolic process	estroge metabo	metabolites en organio olic hydrox	y hydroxy cor compound blos	amide biosynt ganic droxy record ynhetic levels	to cellular
alpha-amino acid biosynthetic process	glycine metabolic process		amin cata pro	nily process	response to insulin stimulus cellular respo steroid		cellula nonitrogen esterol	r response to compound secondary	process	regulation re	etinoic ac	metabol proces	s alcohol a	icohol ymhelio	starvatio
sulfur amino acid metabolic process alpha-ar	neurotransmitter metabolic process		vocess interviewent	process	metabolic process	pro	abolic	alcohol metabolic process	metabolic process cellular horr	hormone	process terpenoio	metabol organic	catabolic	und resp	onse to ent level:
carboxylic ac metabolic proc	xid xess n	oxoacid netabolic process	orga	nic acid lic process	steroid catabolic process	met pro	abolic ca cess pri	tabolic process	regulation cellular resp to insulin stir	ODSE Insulin re	attway	ofactor coenzy etabolic metab rocess proce	onto RISC involved in SS SMIRNA foad	ing onto Ived in	tion-reduction process
					cellular amide	egulation re	esponse to mino acid	response to acid	response to	regulation of cellular insulin stimu	ar me	cofactor etabolic proce small molecule	e regulation o	f metabolic	tion-reduction process organonitrog
carboxylic acid biosynthetic process	monocarbo acid metat process	oolic fa	tty acid	onocarboxylic acid viosynthetic	metabolic tr process	anslation		chemical	3'-UTR-mediate mRNA stabilization	ed negative regu of mRNA cat process RNA stabilizat	abolic	small molecul stabolic proces	regulation of biological		metabolic organonitrog
	small mole	cule fatt	y acid long-d ynthetic fatty a	acid	regulation of r cellular amide	egulation of	cellular esponse to amino acid stimulus	response to thyroxine	mRNA s	-mediate tabilizati	d on	cellular proces	process	acid cycle	primar
organic acid biosynthetic process	process fatty aci	s pro	ygenase unsatur fatty a	ess ratedarachidonic	process positive positive regulation cellular am		response to phenylalanin derivative		drug catabolic process	drug metabo process	Ce	cellular proce	venobiot	acid cycle organic	proces
carboxv	metabolic pr	ocess F	2450 metab thway proce	olic metabolic	of metaboli- translational process integulation of amide metabolic	translation	cellular esponse to	response to L-glutamate		exogenous o	irug	cellular etabolic proce	metaboli	c metabolic	organomuog

Figure 5. Enriched GO (biological process) of low expressed genes in Experiment. Full classification  $\rightarrow$ 

# TRANSPATH® Pathways (2020.2)

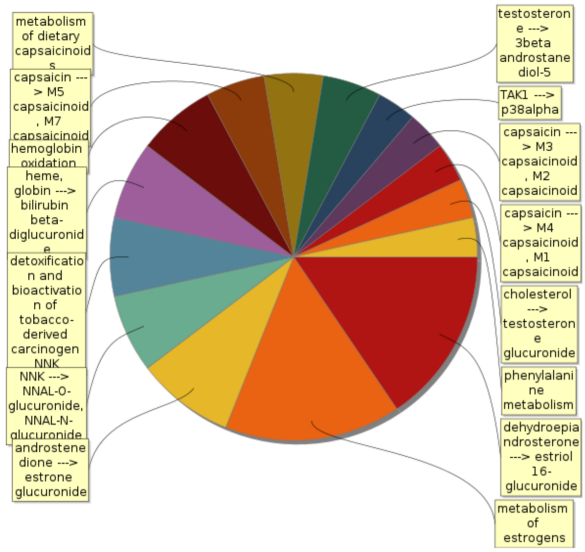
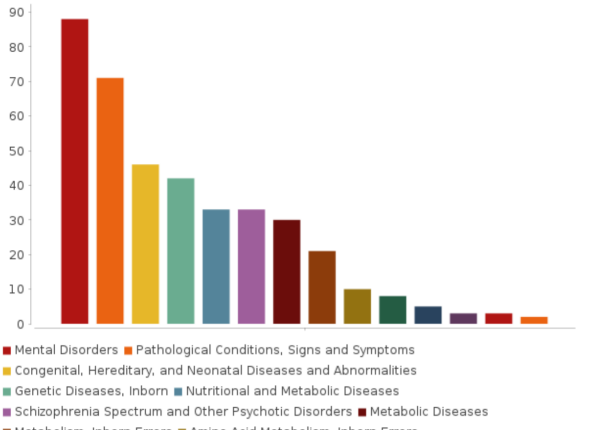


Figure 6. Enriched TRANSPATH® Pathways (2020.2) of low expressed genes in Experiment. Full classification  $\rightarrow$ 

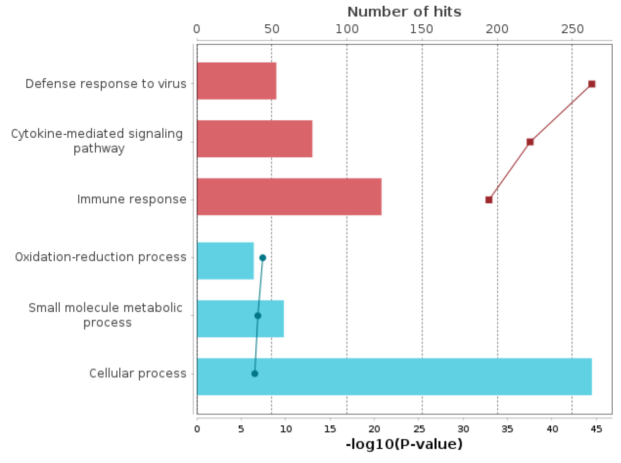
HumanPSD(TM) disease (2020.2)



- 🔳 Metabolism, Inborn Errors 🔳 Amino Acid Metabolism, Inborn Errors
- 🔳 Brain Diseases, Metabolic, Inborn 🔳 Signs and Symptoms, Respiratory 🔳 Chondrosarcoma
- Geographic Atrophy Maple Syrup Urine Disease

Figure 7. Enriched HumanPSD(TM) disease (2020.2) of low expressed genes in Experiment. The size of the bars correspond to the number of bio-markers of the given disease found among the input set. **Full classification**  $\rightarrow$ 

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):



High expressed genes hits Low expressed genes hits -- High expressed genes -log10(P-value)

- Low expressed 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].

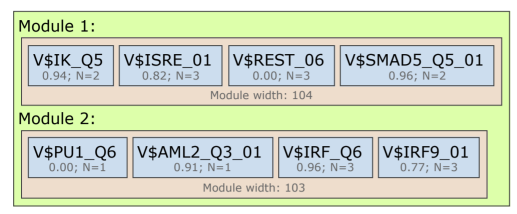
We applied the Composite Module Analyst (CMA) [8] method to detect such potential enhancers, as targets of multiple TFs bound in a cooperative manner to the regulatory regions of the genes of interest. CMA applies a genetic algorithm to construct a generalized model of the enhancers by specifying combinations 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 (high expressed genes in Experiment).

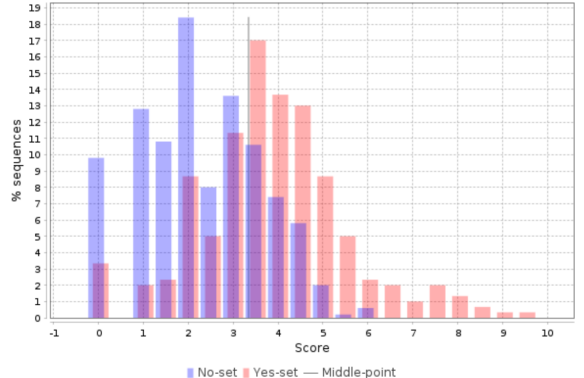
To build the most specific composite modules we choose top high expressed genes as the input of CMA algorithm.

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.



Model score (-p\*log10(pval)): 19.30 Wilcoxon p-value (pval): 9.83e-37 Penalty (p): 0.536 Average yes-set score: 3.84 Average no-set score: 2.33 AUC: 0.77 Middle-point: 3.34 False-positive: 24.00% False-negative: 33.67%



See model visualization table  $\rightarrow$ 

Table 4. List of top ten high expressed genes in Experiment 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
ENSG00000136514	RTP4	receptor transporter protein 4	10.26	REST(h), Ikaros(h), Smad5(h), IRF-9(h), IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), PU.1(h)
ENSG00000126709	IFI6	interferon alpha inducible protein 6	9.96	IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), IRF- 9(h), PU.1(h), Ikaros(h)
ENSG00000185507	IRF7	interferon regulatory factor 7	9.57	REST(h), Ikaros(h), PU.1(h), IRF-9(h), IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), AML2(h)
ENSG00000111331	OAS3	2'-5'- oligoadenylate synthetase 3	9.34	Smad5(h), REST(h), Ikaros(h), IRF-9(h), IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), PU.1(h), AML2(h)
ENSG00000183486	MX2	MX dynamin like GTPase 2	9.31	Smad5(h), REST(h), Ikaros(h), IRF- 1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), IRF- 9(h), AML2(h), PU.1(h)
ENSG00000120217	CD274	CD274 molecule	8.83	IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), IRF- 9(h), PU.1(h), AML2(h), REST(h)
ENSG00000152778	IFIT5	interferon induced protein with tetratricopeptide repeats 5	8.69	Ikaros(h), REST(h), Smad5(h), PU.1(h), IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), IRF- 9(h)
ENSG00000185885	IFITM1	interferon induced transmembrane protein 1	8.67	Smad5(h), Ikaros(h), REST(h), AML2(h), IRF-9(h), IRF-1(h),IRF-2(h),IRF- 3(h),IRF-4(h),IRF-5(h),IRF-6(h),IRF- 7(h),IRF-8(h), PU.1(h)
ENSG00000119917	IFIT3	interferon induced protein with tetratricopeptide repeats 3	8.58	IRF-1(h),IRF-2(h),IRF-3(h),IRF-4(h),IRF- 5(h),IRF-6(h),IRF-7(h),IRF-8(h), IRF- 9(h), Ikaros(h), PU.1(h), Smad5(h)
ENSG00000178685	PARP10	poly(ADP-ribose) polymerase family member 10	8.35	AML2(h), IRF-9(h), IRF-1(h),IRF- 2(h),IRF-3(h),IRF-4(h),IRF-5(h),IRF- 6(h),IRF-7(h),IRF-8(h), PU.1(h), REST(h), Smad5(h), Ikaros(h)

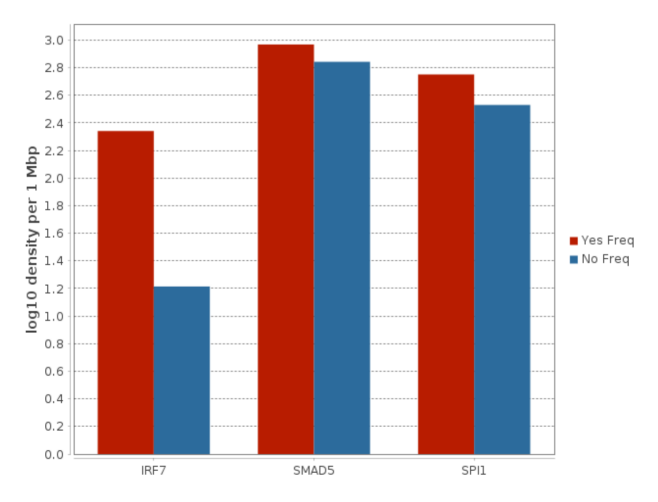
On the basis of the enhancer models we identified transcription factors potentially regulating the **target genes** of our interest. We found 14 transcription factors controlling expression of high expressed genes (see Table 5).

Table 5. Transcription factors of the predicted enhancer model potentially regulating the differentially expressed genes (high expressed genes in Experiment). **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
MO000007703	IRF7	interferon regulatory factor 7	7.61	13.38
MO000020635	SMAD5	SMAD family member 5	6.57	1.34
MO000085616	SPI1	Spi-1 proto-oncogene	6.55	1.66
MO000285816	IRF3	interferon regulatory factor 3	6.44	7.02
MO000007686	IRF1	interferon regulatory factor 1	6.3	7.02
MO000041817	REST	RE1 silencing transcription factor	6.08	1.46
MO000026678	IKZF1	IKAROS family zinc finger 1	5.96	1.17
MO000023424	IRF8	interferon regulatory factor 8	5.89	7.02
MO000007691	IRF2	interferon regulatory factor 2	5.73	7.02
MO000026238	RUNX3	RUNX family transcription factor 3	5.33	1.17

The following diagram represents the key transcription factors, which were predicted to be potentially regulating differentially expressed genes in the analyzed pathology: IRF7, SMAD5 and SPI1.



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 Table 6.

Table 6. Master regulators that may govern the regulation of high expressed genes in Experiment. **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	Total rank	LogFoldChange
MO000038322	LPS:lbp:CD14:TLR4:MD- 2:MyD88:IRAK-1{pS376} {pT387}	CD14, IRAK1, LBP, LY96, MYD88, TLR4	CD14 molecule, MYD88 innate immune signal transduction adaptor, interleukin 1 receptor associated ki	68	0.62
MO000039099	IL-1beta-p17:IL-1RI:IL- 1RAcP:MyD88:tollip:IRAK- 1{pS376}{pT387}:IRAK- 4:IRAK-2	AC093012.1, IL1B, IL1R1, IL1RAP, IRAK1, IRAK2, MYD88, TOLLIP	MYD88 innate immune signal transduction adaptor, interleukin 1 beta, interleukin 1 receptor accessor	69	0.62
MO000041437	dsRNA:TLR3:TRIF	TICAM1, TLR3	toll like receptor 3, toll like receptor adaptor molecule 1	144	0.75
MO000038316	LPS:lbp:CD14:TLR4:MD- 2:TIRAP:IRAK-2	CD14, IRAK2, LBP, LY96, TIRAP, TLR4	CD14 molecule, TIR domain containing adaptor protein, interleukin 1 receptor associated kinase 2, li	159	0.61
MO000019259	c-Cbl(h)	CBL	Cbl proto- oncogene	174	0.37
MO000032632	PKCepsilon(h)	PRKCE	protein kinase C epsilon	185	0.35
MO000019312	IKK-i(h)	ІКВКЕ	inhibitor of nuclear factor kappa B kinase subunit epsilon	197	0.45
MO000079043	PML-4(h)	PML	promyelocytic leukemia	211	1.35
MO000057630	c-Cbl(h)	CBL	Cbl proto- oncogene	225	0.37
MO000082806	IKK-i-isoform1(h)	ІКВКЕ	inhibitor of nuclear factor kappa B kinase subunit epsilon	237	0.45

The intracellular regulatory pathways controlled by the above-mentioned master regulators are depicted in Figure 8. This diagram displays 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.

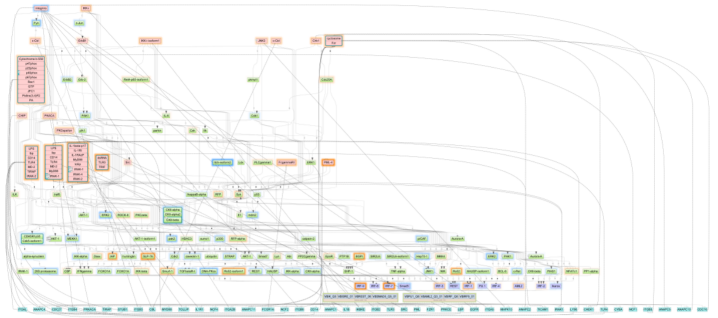


Figure 8. Diagram of intracellular regulatory signal transduction pathways of high expressed genes in Experiment. Master regulators are indicated by red rectangles, transcription factors are blue rectangles, and green rectangles are intermediate molecules, which have been added to the network during 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 HumanPSD<sup>TM</sup> [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 HumanPSD<sup>TM</sup> 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<sup>TM</sup> 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 7. Prospective drug targets selected from full list of identified master regulators filtered by druggability score from HumanPSD<sup>™</sup> 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.

#### See full table $\rightarrow$

Gene symbol	Gene Description	Druggability score	Total rank	LogFoldChange
FCGR1A	Fc fragment of IgG receptor Ia	21	253	0.55
CEACAM1	CEA cell adhesion molecule 1	1	283	0.85
MAPKAPK2	MAPK activated protein kinase 2	7	408	0.31
PSMA7	proteasome 20S subunit alpha 7	3	449	0.2
ROCK2	Rho associated coiled-coil containing protein kinase 2	2	462	0.25
ITGAL	integrin subunit alpha L	8	479	0.33

Table 8. 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.

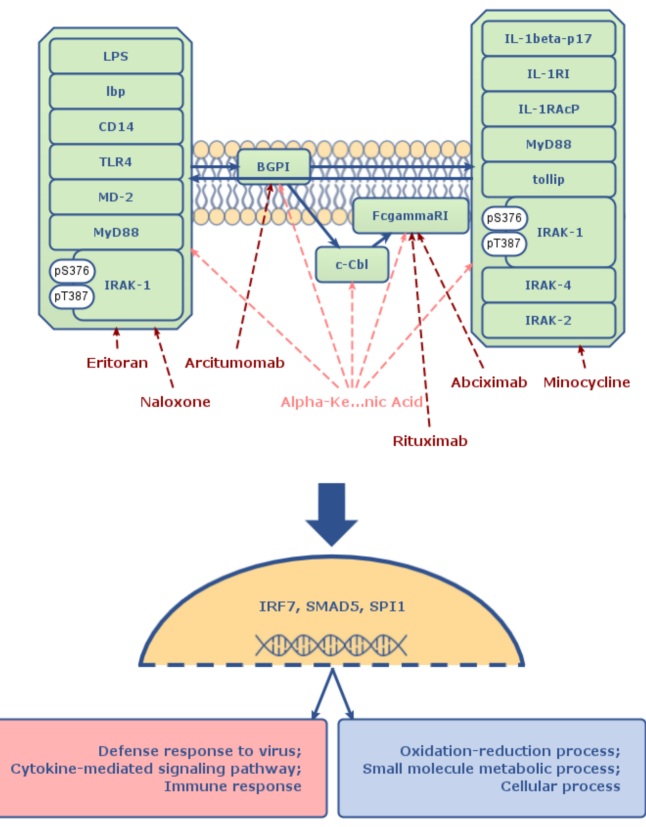
#### See full table $\rightarrow$

Gene Description	Druggability score	Total rank	LogFoldChange
Cbl proto-oncogene	131.54	225	0.37
Fc fragment of IgG receptor Ia	9.41	253	0.55
CEA cell adhesion molecule 1	9.41	283	0.85
microtubule associated serine/threonine kinase like	2.98	296	0.93
dual specificity phosphatase 4	63.51	332	0.33
midline 2	131.54	384	0.61
	Cbl proto-oncogene Fc fragment of IgG receptor Ia CEA cell adhesion molecule 1 microtubule associated serine/threonine kinase like dual specificity phosphatase 4	Gene DescriptionscoreScore131.54Cbl proto-oncogene131.54Fc fragment of IgG receptor Ia9.41CEA cell adhesion molecule 19.41microtubule associated serine/threonine kinase like2.98dual specificity phosphatase 463.51	Gene DescriptionscorerankCbl proto-oncogene131.54225Fc fragment of IgG receptor Ia9.41253CEA cell adhesion molecule 19.41283microtubule associated serine/threonine kinase like2.98296dual specificity phosphatase 463.51332

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:

- FcgammaRI
- LPS:lbp:CD14:TLR4:MD-2:MyD88:IRAK-1{pS376}{pT387}
- IL-1beta-p17:IL-1RI:IL-1RAcP:MyD88:tollip:IRAK-1{pS376}{pT387}:IRAK-4:IRAK-2
- BGPI
- c-Cbl

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: Arcitumomab, Naloxone, Abciximab, Eritoran, Alpha-Ketomalonic Acid, Rituximab and Minocycline, 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 9. 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<sup>m</sup> database) See full table  $\rightarrow$ 

Name	Target names	Drug rank	Disease activity score	Phase 4	Status (provided by Drugbank)
Sorafenib	BRAF, RET	135	1	Carcinoma, Hepatocellular, Carcinoma, Renal Cell, Liver Neoplasms, Neoplasms, Noma, Thrombosis	small molecule,approved,investigational
IDN-6556	CASP7, CASP1	140	2		small molecule, investigational
Rituximab	FCGR1A	162	2	Anti-Neutrophil Cytoplasmic Antibody-Associated Vasculitis, Arthritis, Arthritis, Rheumatoid, Autoimmune Diseases, Brain Abscess, Burkitt Lymphoma, Diabetes Mellitus	biotech,approved
Peginterferon alfa-2a	IFNAR2	221	11	Hepatitis C, HIV Infections, Hemophilia A, Hepatitis, Hepatitis B, Hepatitis B, Chronic, Hepatitis C, Chronic	biotech,approved,investigational
Peginterferon alfa-2b	IFNAR2	221	11	Hepatitis C, Hepatitis, Hepatitis B, Hepatitis B, Chronic, Hepatitis C, Chronic, Hepatitis, Chronic	biotech,approved

# <u>Repurposing drugs</u>



Table 10. Repurposed drugs used in clinical trials for other pathologies (prospective drugs against the identified drug targets on the basis of literature curation in HumanPSD<sup>TM</sup> database) See full table  $\rightarrow$ 

Name	Target names	Drug rank	Phase 4	Status (provided by Drugbank)
Abciximab	ITGB3, FCGR1A, ITGA2B	25	Acute Coronary Syndrome, Angina, Unstable, Coronary Artery Disease, Coronary Disease, Diabetes Mellitus, Ischemia, Myocardial Infarction	biotech,approved
Efalizumab	ITGAL, FCGR1A	27	Psoriasis, ST Elevation Myocardial Infarction	biotech, approved, investigational
Porfimer	FCGR1A	28	Carcinoma, Non-Small-Cell Lung, Carcinoma, Small Cell, Lung Neoplasms, Neoplasms, Small Cell Lung Carcinoma	small molecule,approved,investigational
Bosutinib	SRC, HCK, LYN	29	Leukemia, Myeloid	small molecule, approved
Tirofiban	ITGB3, ITGA2B	35	Acute Coronary Syndrome, Coronary Artery Disease, Coronary Disease, Myocardial Infarction, No- Reflow Phenomenon, ST Elevation Myocardial Infarction	small molecule, approved



No prospective drugs were found, which would be predicted by PASS software to be active against the identified drug targets and would be predicted to have biological activity against the studied disease(s).



Table 11. 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
Alpha-Ketomalonic Acid	TEC, PTPRO, PTPN5, IL4R, PRKACA, JAK3, HTRA2	94	2.1
Myo-Inositol	TEC, PTPRO, PTPN5, IL4R, PRKACA, JAK3, HTRA2	115	2.64
4,7-Dioxosebacic Acid	TEC, PTPRO, PTPN5, PRKACA, JAK3, HTRA2, TRAF6	131	1.98
2-Oxo-4- Methylpentanoic Acid	TEC, PTPRO, PTPN5, IL4R, JAK3, PRKACA, HTRA2	139	1.73
Trifluoroalanine	TEC, PTPRO, PTPN5, IL4R, JAK3, HTRA2, TRAF6	143	1.84

As the result of drug search we propose the following drugs as most promising candidates for treating the pathology under study: Sorafenib, Abciximab and Alpha-Ketomalonic Acid. These drugs were selected for acting on the following targets: RET, FCGR1A and CBL, 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 obtained from *liver* tissue. The study is done in the context of *Hepatitis C*. 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: RET, FCGR1A and CBL, 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:



#### FcgammaRI, LPS:lbp:CD14:TLR4:MD-2:MyD88:IRAK-1{pS376} {pT387}, IL-1beta-p17:IL-1RI:IL-1RAcP:MyD88:tollip:IRAK-1{pS376}{pT387}:IRAK-4:IRAK-2, BGPI and c-Cbl

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: Arcitumomab, Naloxone, Abciximab, Eritoran, Alpha-Ketomalonic Acid, Rituximab and Minocycline. 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.

- FcgammaRI
- LPS:lbp:CD14:TLR4:MD-2:MyD88:IRAK-1{pS376}{pT387}
- IL-1beta-p17:IL-1RI:IL-1RAcP:MyD88:tollip:IRAK-1{pS376}{pT387}:IRAK-4:IRAK-2
- BGPI
- c-Cbl

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 DNA-binding 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<sup>™</sup> 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 DNA-binding 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<sup>™</sup> and predicting potential drugs using PASS program.

#### Method for analysis of known pharmaceutical compounds

We selected compounds from HumanPSD<sup>M</sup> 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<sub>PSD</sub>),

2. ranking by "Disease activity score" (*D*-score<sub>PSD</sub>).

"Target activity score" (*T*-score<sub>PSD</sub>) is calculated as follows:

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

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<sub>PSD</sub>):

$$D\text{-}score_{\scriptscriptstyle PSD} = \begin{cases} \sum\limits_{d \in D} \sum\limits_{p \in P} phase(d, p) \\ 0, \ D = \varnothing \end{cases},$$

where *D* is the set of selected diseases, and if *D* is empty set, D-score<sub>PSD</sub>=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 precalculated 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:

$$D$$
-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.

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In case of any questions please contact us at <a href="mailto:support@genexplain.com">support@genexplain.com</a>

#### Supplementary material

1. Supplementary table 1 - Detailed report. Composite modules and master regulators (high expressed genes in Experiment).

#### 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<sup>™</sup> 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.

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