A novel open access web portal for integrating mechanistic and toxicogenomic study results

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Running head: Open access web portal for toxicogenomics

Keywords: toxicogenomics, systems biology, gene expression analysis; mechanism inference, toxicity prediction

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Abstract

Applying toxicogenomics to improving the safety profile of drug candidates and crop protection molecules is most useful when it identifies relevant biological and mechanistic information that highlights risks and informs risk mitigation strategies. Pathway-based approaches, such as GSEA, integrate toxicogenomic data with known biological process and pathways. Network methods help define unknown biological processes and offer data reduction advantages. Integrating the two approaches would improve interpretation of toxicogenomic information. Barriers to the routine application of these methods in genome-wide transcriptomic studies include a need for “hands-on” computer programming experience, the selection of one or more analysis methods (e.g. pathway analysis methods), the sensitivity of results to algorithm parameters, and challenges in linking differential gene expression to variation in safety outcomes. To facilitate adoption and reproducibility of gene expression analysis in safety studies, we have developed Collaborative Toxicogenomics (CTox), an open-access integrated web portal using the Django web framework. The software, developed with the Python programming language, is modular, extensible and implements “best-practice” methods in computational biology. New study results are compared to over 4,000 rodent liver experiments from Drug Matrix and open TG-GATEs. A unique feature of the software is the ability to integrate clinical chemistry and histopathology-derived outcomes with results from gene expression studies, leading to relevant mechanistic conclusions. We describe its application by analyzing the effects of several toxicants on liver gene expression and exemplify application to predicting toxicity study outcomes upon chronic treatment from expression changes in acute-duration studies.
Introduction

Despite significant advances, determining the relevance of toxicity findings from animal safety studies to human health effects is challenging. Identifying the mechanistic underpinnings of a toxicity finding is key, and characterization of tissue samples using gene expression is often performed towards this end. Initiatives to study a broad range of toxicants and reference molecules has produced rich contextual information aiding the interpretation of gene expression changes produced by a molecule of interest (Ganter et al., 2005; Igarashi et al., 2015). However, these data are challenging to process, search and analyze. Further, many analysis algorithms are available, ranging from mature approaches such as overrepresentation analysis (widely used in the DAVID web application) (Huang et al., 2009), the related gene set enrichment analysis (GSEA) method (Subramanian et al., 2005) and Connectivity Map (CMAP) (Lamb et al., 2006; Smalley et al., 2010), to newer methods purporting to offer additional insights (Lee et al., 2014; Tawa et al., 2014; Te et al., 2016; Bell et al., 2016). These methods are generally not integrated and utility is often limited to computational specialists, who themselves may struggle to implement methods or reproduce results from the literature without having access to source code and detailed protocols. Finally, for those approaches that offer user-friendly interfaces, a user must navigate the process of producing “analysis-ready” fold change results for each gene and upload them in multiple applications.

Herein, we sought to improve the integration of standard methods with newer network-based methods to improve usability for toxicologists and mechanistic interpretation relevant to risk assessment. We describe the creation of an open-source, publicly-available platform for the analysis of gene expression results from toxicity studies. While the platform was developed using rat liver studies, it can be used to analyze data from any study comparing the effect of a treatment or other intervention to control samples. We have processed and loaded within the application results from Drug Matrix
(DM) (Ganter et al., 2005) and TG-GATEs (TG) (Igarashi et al., 2015), which between them provide results on 4182 liver experiments. This includes both expression results and the histology / clinical chemistry data within those repositories, analyzed using a shared lexicon and standardized between the sources. In addition to analyzing their uploaded data, users can query DM and TG experiments and analyze results at the level of individual genes and gene sets. CMAP (Lamb et al., 2006; Smalley et al., 2010) and related approaches attempt to infer properties of a treatment of interest by identifying other drugs or tool compounds having similar effects on the global transcriptome. Users can identify the experiments from DM/TG having the most similar transcriptional effects.

Previously, we described an approach linking perturbation of co-expression networks (or “modules”) to a variety of toxicity phenotypes (TXG-MAP analysis) (Sutherland et al., 2017). This allows the identification of networks perturbed by a molecule of interest that are also generally associated with its observed toxicity across a broad range of conditions. Here, we extend this approach by identifying a non-redundant collection of Gene Ontology (GO) and canonical pathways that can be modulated in liver toxicity studies.

**Methods**

**Web application and availability**

A web application was created using the Python and R programming languages. The application consists of a PostgreSQL database, Python and R backend computation scripts, celery queuing system for job management (http://www.celeryproject.org/; accessed Aug 29, 2018). The user interface was developed using the Django web framework and the HighCharts JavaScript library for visualizations (https://www.highcharts.com/; accessed Aug 29, 2018). All data files and source code are provided in the GitHub repository hosted at https://github.com/IndianaBiosciences/toxapp. References to file
names below refer to their location in the repository. The public instance of the application is hosted on Linux servers from Amazon Web Services. The application has been thoroughly tested on Ubuntu 16.04 LTS and Centos 7.4. A deployment script (setup_server.sh) allows private instances to be created with all requirements satisfied.

**Gene-level data preparation**

Throughout, an “experiment” denotes a comparison of treatment vs. control samples; a treatment consists of a combination of drug, dose, duration of dosing, route of administration. More broadly, an experiment (or intervention) can be used to compare animals harboring null alleles of a gene of interest (i.e. a knockout animal) vs. wild-type animals. In Drug Matrix (DM) (Ganter *et al.* 2005) and TG-GATEs (TG) (Igarashi *et al.*, 2015), experiments compare the effects of drug treatment in livers from 2-5 treated animals vs. 2-5 vehicle matched controls. The complete list of 3528 TG and 654 DM experiments viewable in the application is available in Table S1 of our prior work (Sutherland *et al.*, 2017) and the repository file data/experiments_DM_TG.txt

Affymetrix CEL files for individual rat liver samples analyzed with RG230-2 microarrays were obtained from Drug Matrix (https://ntp.niehs.nih.gov/drugmatrix/index.html, accessed Aug 29, 2018) and the TG-GATEs repositories (http://toxico.nibiohn.go.jp/english/; accessed Aug 29, 2018). Normalized log intensities were calculated using the University of Michigan “Brain Array” version 19 assignment of microarray probes to genes (http://brainarray.mbnf.med.umich.edu/Brainarray/Database/CustomCDF/19.0.0/entrezg.asp; accessed Aug 29, 2018) and the RMA algorithm (Irizarry *et al.*, 2003) as implemented in the Affy package in Bioconductor (scripts/NORM.R). For any given experiment, fold change values for each gene were obtained using the Bioconductor package Limma (scripts/Limma.R). Pre-processed fold-change results, consisting of log2 fold change (log2_fc), number of treated animals (n_trt), number of control animals
(n_ctl), average gene intensity in controls (expression_ctl0), unadjusted p-value (p) and Benjamini Hockberg-adjusted p-value (p_bh) are available in the data repository at location data/groupFC. For example, repeat-dose results for gemfibrozil (from TG-GATEs) are in the file data/groupFC/gemfibrozil.Rat.in_vivo.Liver.Repeat-groupFC.txt.

**Gene model and orthology**

The rat genome was selected as the reference organism, using the Entrez gene nomenclature. The application supports the upload of rat, mouse and human gene expression data. Mapping of human and mouse genes to rat was performed using orthology information from RGD (Shimoyama et al., 2015) as described in reference (Sutherland et al., 2016). The gene model includes all rat genes according to RGD, whether or not they are tested on a given measurement technology. However, human or mouse genes having no rat ortholog were excluded, and users uploading data for these organisms will not obtain fold-change results for such genes. At the level of pathways or modules where results are aggregated across several genes (see below), the impact is trivial. The file data/gene_info.txt includes 18004 rat Entrez gene ids, mapped to 17540 mouse and 17820 human genes.

**TXG-MAP co-expression analysis (modules)**

Co-expression network analysis seeks to identify genes that respond similarly to perturbations in a biological system of interest. We used the Bioconductor package WGCNA to identify groups of co-expressed genes (or “modules”) from the Drug Matrix rat liver data. Algorithm parameters and details on the creation the 415 modules were provided in reference (Sutherland et al., 2016). Because the modules are associated with several toxicity phenotypes, we refer to the method as “Toxicogenomic Module Association with Pathogenesis (TXG-MAP). A visual representation of the modules on a phylogenetic tree provides a consistent visual frame of reference for analysis. Module scores are
presented as Z-scores, such that a score of 2 is achieved by fewer than 4% of drug treatments, a score of 3 in fewer than 1%, etc. This indicates the magnitude of an effect in the context of other drug treatments.

Here, we updated the modules by replacing the original Affymetrix probe sets with Brain Array definitions (via rat Entrez gene IDs to which both were mapped 1:1). The assignment of genes to modules is provided in data/WGCNA_modules.txt, and module scores for a given experiment calculated from the method `score_modules` in the source package `src/computation.py`.

**Gene set analysis (GSA)**

Gene set analysis seeks to identify canonical pathways, gene ontology (GO) and other gene sets that are enriched among the most differentially expressed genes. We used the PAGE algorithm (Kim and Volsky, 2005) as implemented in the Bioconductor package Piano, given similar performance compared to the popular GSEA algorithm and significantly faster calculation (Varemo et al., 2013). Gene sets used in the analysis were taken from GO and MSigDB, as described in full in reference (Sutherland et al., 2016). The assignment of genes to gene sets is provided in `data/rgd_vs_GO_expansion.txt` and `data/MSigDB_and_TF_annotation.txt`. The method `score_gsa` in the source package `src/computation.py` performs the calculations in R. Gene sets from GO and MSigDB are numerous and overlapping. Limiting to GO and curated pathways from MSigDB, there are 11 581 gene sets for consideration. In prior work, we noted that 1839 of these gene sets were inducible or repressible in 1% or more of 3525 TG experiments (in brief, “inducible gene sets”). The vast majority of gene sets represent molecular processes that are either absent from liver or rarely affected by drug perturbation.

To eliminate redundancy among related gene sets, the pvclust R library was used to hierarchically cluster 1839 gene sets vs. GSA scores from all 3528 TG liver experiments (command in R: `pvclust(data, method.dist="cor", method.hclust="ward.D", nboot=1000, parallel = TRUE)`). Clustering
algorithms may create too many clusters, or divisions of samples that are no longer observed when the dataset is changed (here, the term sample from statistics is a liver experiment comparing treated vs. control liver expression, not a liver sample). The algorithm evaluates the significance of edges in the dendrogram while performing bootstrap resampling of the samples. The method reduce_tree in the src/treemap.py package takes as input the dendrogram exported from R, traces up the dendrogram from terminal to root nodes and finds the edge nearest the root that is non-significant. All gene sets below the edge are merged into a single cluster. The related method reduce_tree_pca uses principal components analysis on a table of GSA scores having clustered gene sets on columns and 3528 liver experiments on rows. The gene set with highest loading in the first principal component is selected as a representative for the cluster.

A visual representation of the non-redundant gene sets was created from the pruned dendrogram returned by the reduce_tree_pca method. The tree was truncated at a height of 15, a threshold selected to separate large branches near the tree root. Visually, we found this to improve the aesthetics and interpretability of results by making better use of 2D plane (i.e. web browser screen space). The truncated dendrogram was rendered using the yFiles circular layout algorithm in Cytoscape.

Clinical chemistry and histology results

Clinical chemistry and histology results from DM and TG experiments are available from the respective repositories. A simplified view of these results are provided within our application, by focusing on a limited number of non-redundant endpoints encountered 10 or more times across the combined DM and TG datasets; details on the curation process and lexicon creation are given in reference (Sutherland et al., 2017). The following histology endpoints are included: Apoptosis/SingleCellNecrosis:Hepatocellular; Congestion/Hemorrhage/Edema:Vascular;
Degeneration/Necrosis:Hepatocellular; Dilation/Dilatation/Ectasia/Distension:Vascular; Fibrosis;
Glycogen_Increased:Hepatocellular; Hematopoiesis; Hyperplasia:Biliary; Hypertrophy:Hepatocellular;
Infiltration/Inflammation; Mitosis_Increased:Hepatocellular; Vacuolation:Hepatocellular. The following
clinical chemistry endpoints are albumin (ALB), alkaline phosphatase (ALP), alanine aminotransferase
(ALT), aspartate aminotransferase (AST), cholesterol (chol), gamma-glutamyl transpeptidase (GGT),
glucose (Glu), total bilirubin (T Bili), total protein (TP), triglycerides (Trig). The source file containing
these results for each experiment is data/toxicology_results_DM_TG.txt.

Association of gene sets with toxicity phenotypes

Previously, we described the assignment of DM and TG experiments to 13 representative
toxicity phenotypes (Table S4 in reference (Sutherland et al., 2017)). A toxicity phenotype denotes the
occurrence of one or more dominant histological or clinical chemistry changes. For example, the
phenotype “Necrosis, 1) no other findings” denotes the occurrence of 2 minimal grades, one slight
grade, or greater severity of hepatocellular necrosis among 3 animals in a dose group, without other
significant co-occurring histological changes (“pure” necrosis). In addition, we defined an additional
composite phenotype of “adverse at 29 days” which included histological changes such as necrosis,
single cell necrosis, bile duct hyperplasia that are deemed adverse by pathologists in studies of 29-day
duration. The 14 toxicity phenotypes were “adverse at 29 days”, “bile duct hyperplasia”, “cholesterol
decrease”, “cholesterol increase”, “fibrosis”, “hematopoiesis”, “hypertrophy”, “increased glycogen”,
“increased mitosis”, “necrosis”, “single cell necrosis”, “trigs (triglyceride) decrease”, “trigs increase”,
“vacuolation”. These abbreviated phenotype terms correspond to the detailed curated histology labels
above: “necrosis” aligns to “Degeneration/Necrosis:Hepatocellular”, meaning that a lesion denoted as
necrosis, degeneration, or degeneration and necrosis of hepatocytes would be assigned to the
“necrosis” toxicity phenotype.
DM and TG experiments were labelled as positive or negative for each of the 14 toxicity phenotypes. Repeating our earlier analysis with modules (Sutherland et al., 2017), the statistical association between induction or repression of each gene set (in the 1839 “inducible” set) and the occurrence of the liver phenotype was calculated using logistic regression. The models in the R statistical language were computed as: 

\[
\text{glm(}\text{toxic} \sim \text{avgAbsEG} + \text{pathway\_GSA\_score, family=“binomial”}).
\]

As described in reference (Sutherland et al., 2017), avgAbsEG is the average absolute module score, a measure which is equivalent to computing the percentage of genes that are differentially expressed. This covariate removes the variation in the odds of observing toxicity that is explained by overall effects of the treatment on gene expression, such that attributing significance to a particular gene set genuinely reflects its role in explaining toxicity, not surrogacy with overall gene expression changes. Finally, we report separately the association between gene sets in toxicity for concurrent injury (expression analysis performed on samples where the lesion is present) vs. predictive (expression analysis performed on samples collected at 3, 6, 9 or 24 hours of dosing, before the lesion was noted). The source file containing these association results is \textit{data\_geneset\_vs\_tox\_association.txt}.

Expression similarity from microarray vs. RNA-seq measurement technologies

To support the validity of comparing experiments analyzed using RNA-seq to experiments from DM and TG using Affymetrix microarrays, datasets were obtained from the Gene Expression Omnibus repository (GEO; \url{https://www.ncbi.nlm.nih.gov/geo/}, accessed 03/22/2019) and analyzed with the CTox application.

The first comparison evaluated RNA-seq results from the SEQC consortium (Wang et al., 2014). The file

\textit{GSE55347\_TGxSEQC\_GeneExpressionIndex\_Magic\_20120831\_116samplesGSE47875\_series\_matrix.txt}
was obtained for GEO series GSE55347. The authors’ mapping of Aceview/Refseq transcripts to NCBI Entrez Gene IDs was used. Since our RNA-seq workflow uses Ensemble identifiers, we mapped Entrez Gene IDs to Ensemble Rat version 6.0 (Rnor_6.0) using the NCBI mapping file ([ftp://ftp.ncbi.nih.gov/gene/DATA/gene2ensembl.gz](ftp://ftp.ncbi.nih.gov/gene/DATA/gene2ensembl.gz); accessed 3/18/2019). Where multiple Aceview/Refseq transcripts mapped to a single Ensemble gene, only the highest abundance transcript was retained, as assessed by the average abundance across the 116 samples. Finally, since the authors’ pipeline produced continuous abundance estimates with quasi-normal distribution, these were analyzed directly using the R package Limma, producing fold-change and p-value of each gene. Computation of pathway, module and experiment similarity were performed in the same manner as all other results processed within the CTox application. Because the SEQC study used DM microarray results, there was no need to process Affymetrix CEL files from GSE47875, as they were already processed and available in the application.

A second comparison evaluated results from a recent study, referred to as “Abvie study” in results (Rao et al., 2019). As RNA-seq pipelines have become increasingly standardized since the SEQC study, this afforded the opportunity to evaluate results from a representative “industry-standard” approach. The file GSE122315_Raw_counts.txt was obtained from GSE122315. Except for rounding the counts to the nearest integer, this file was processed as-is by the CTox application. Affymetrix RG230-2 CEL files were obtained from the file GSE122184_RAW.tar for series GSE122184, and uploaded using the standard workflow.

The above datasets produced pairs of treatment vs. control experiments that differed only in the measurement technology employed. We evaluated the similarity of experiment pairs by calculating the Pearson R correlation coefficient using the 382 non-redundant pathways or the 405 TXG MAP modules. The experiments in Dataset S6 were sorted by the average absolute module score (avgAbsEG), calculated as described in reference (Sutherland et al., 2017).
Results

Web application

A web application for analysis of gene expression study data was created using the Django web framework. The use of Python, the R statistical language and PostgreSQL database, technologies with a large user base among scientific developers, will facilitate its extension by others. The collaborative toxicogenomics (CTox) application allows users to describe the purpose of a study, treatments/perturbations under investigation and analyze sample results in the form of Affymetrix CEL files or RNA-seq count data. Expression results from rat, mouse and human are currently supported. Representative application dialogs are shown in Figure 1.

Available gene expression and histology / clinical chemistry results

DrugMatrix (DM) (Ganter et al., 2005) and the open TG-GATEs (TG) (Igarashi et al., 2015) repositories are rich resources describing the effects of 308 drugs, toxicants and other perturbations across 4182 liver experiments (an experiment denotes the administration of a drug for at a given dose and duration compared to a control group). We reprocessed all Affymetrix CEL files using updated probe set definitions (Dai et al., 2005) and robust p-value estimation using Limma (Irizarry et al., 2003).

Several result types are available when viewing one or more experiments (Table 1). Gene-level analysis, whereby the expression change of each gene is reported as a fold change and p-value, can be used to identify the most induced or repressed genes. Gene set analysis (GSA), implemented using the PAGE algorithm (Kim and Volsky, 2005), is applied to gene sets from Gene Ontology (GO) terms, canonical pathways and others from the Molecular Signatures Database (MSigDB) (Subramanian et al., 2005). Co-expression networks summarize results across 415 co-expressed gene sets (i.e. “modules”)
obtained by analysis of Drug Matrix rat liver experiments (Sutherland et al., 2017). When an experiment of interest strongly resembles the global transcriptional profile of a reference compound, they may share a similar mode of action. To that end, comparison of a new experiment to all TG and DM rat liver experiments is easily achieved. Finally, a simplified view of clinical chemistry and histology results across the two repositories allows the identification of treatments that lead to a toxicity phenotype (or apical endpoint) of interest and their further characterization in the application.

Table 1. Analysis methods available within the CTox web application

<table>
<thead>
<tr>
<th>Method</th>
<th>Summary</th>
<th>Strengths/weaknesses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene-level analysis</td>
<td>A list of all genes probed by the measurement technology, with fold change and associated p-value for treatment vs. control</td>
<td>Ease of interpretation, but high risk of false positives due to large number of hypotheses being tested*. High risk of “confirmation bias”, i.e. focusing on the induction/repression of genes of interest in experiments where many (unrelated) genes are more significantly perturbed by the experimental conditions</td>
</tr>
<tr>
<td>Gene Set Analysis (GSA)</td>
<td>A list of gene sets (GO terms, pathways, and others) for which the constituent genes are disproportionally induced or repressed compared to all measured genes; returns a score and p-value for treatment vs. control</td>
<td>Simplifies by summarizing effects aggregated across related genes. Ease of interpretation depends on the perturbed gene set; e.g. “cholesterol biosynthesis” is clear in the context of liver, but “generation of neurons” is not. In theory, lower risk of false positives, however the MSigDB collection now approaches 18,000 gene sets.</td>
</tr>
<tr>
<td>TXG-MAP module analysis</td>
<td>A list of scores, one for each of 415 modules in the TXG-Map. Modules consist of co-expressed genes.</td>
<td>Simplifies by summarizing across co-expressed genes. Unlike other methods, provides context on magnitude of effect compared to other liver perturbations. Captures co-regulated biology not described by canonical pathways, but interpretation requires study of constituent genes. Results depends on quality / comprehensiveness of training datasets. Low risk of false positives.</td>
</tr>
<tr>
<td>Similar experiments analysis</td>
<td>A list of most similar experiments, compared to user’s experiment. Global transcriptional similarity, assessed by comparing ~400</td>
<td>The fastest way to help understand mechanism, in those cases where a compound of interest is similar to compounds / drugs with well-understood mechanistic effects. Intermediate risk of false positives (ca. 4000 comparisons performed). Requires global (i.e. expression</td>
</tr>
</tbody>
</table>
Non-redundant gene sets for liver analysis

The identification of gene sets for which the constituent genes are disproportionally induced or repressed is a common analysis method for transcriptomic studies. The popular GSEA approach can be applied to gene sets from GO, KEGG, REACTOME, and the MSigDB compilation of curated pathways and other sources. MSigDB currently approaches 18,000 gene sets, resulting in as many hypothesis tests as analysis of all individual genes in the genome (gene-level analysis). As such, large adjustments to p-values occur when correcting for multiple hypothesis testing, introducing many false negative results. Several approaches for reducing the number of gene sets have been proposed (Liberzon et al., 2015; Cantini et al., 2018). Minimal, non-redundant gene sets will vary by tissue, owing to varying expression of the underlying genes and tissue-specific functions. Previously, we described a subset of GO terms (biological process and cellular component) and curated pathways (KEGG, REACTOME, Biocarta and others) that are perturbed in 1% or more of TG liver experiments (Sutherland et al., 2016). This reduced the number of gene sets for consideration from 11581 to 1839 (henceforth, the “inducible/repressible” set, or “inducible pathways” for conciseness).

Within the 1839 inducible set, inclusion of gene sets from different sources suggested a significant level of redundancy (e.g. cholesterol biosynthesis from GO, KEGG, REACTOME). Using scores for each gene set vs. 3528 TG liver experiments, hierarchical clustering was used to organize the gene sets into clusters
of gene sets that are induced/repressed in the same experiments. By testing the stability of clusters upon repeated sampling of experiments (methods), we obtained 382 clusters containing one or more gene sets (Dataset S1). Within each cluster, a single gene set that best captures the variation within the cluster was identified, serving as a surrogate for the others (no consideration of whether the gene set in question best summarizes biological themes represented in the cluster). The 382 non-redundant gene sets were graphically organized to facilitate visual interpretation of GSA results for liver gene expression studies (Figure 2). The example in Figure 2 shows a cluster of 34 terms for which the REACTOME term “Genes involved in Destabilization of mRNA by AUF1” captured the maximum variation for the 34 clustered terms. Inclusion in the cluster of 4 GO cellular component (CC) terms relating to the proteasome and inspection of other pathway annotations suggests degradation of cell signaling factors by the proteasome, a common method of regulation (Rousseau and Bertolotti, 2018).

Association of inducible gene sets with toxicity phenotypes

Gene expression profiling can be used to explain putative mechanisms responsible for drug-induced morphologic changes in the liver. Since many differentially-expressed genes or gene sets will have no relation to the observed morphologic changes, identifying the subset of expression changes linked to injury is difficult. Previously, we described the statistical association between induction or repression of networks of co-expressed genes sets, TXG-map modules, and the occurrence of various histology-anchored toxicity phenotypes (Sutherland et al., 2017). Here, we performed similar analyses to establish the relationship between perturbation of inducible gene sets and the occurrence of liver toxicity phenotypes. We distinguish predictive relationships (i.e. expression profiling from tissue collected 24 hours after a single dose, before the appearance of injury) from concurrent relationships (i.e. expression profiling from the injured tissue). Mirroring our findings with liver modules, we found a larger number of statistically-significant relationships for concurrent injury, compared to single dose...
exposures (Dataset S2). For each combination of a time point and toxicity phenotype, pathways were ranked from most to least associated with a given phenotype. For any experiment, the odds of observing a toxicity phenotype can be computed from pathway scores (Table 2). This allows for the prediction of toxicity in longer duration studies when performing expression studies of ≤ 24 hour treatment duration.
Table 2. Top three non-redundant pathways for association with toxicity phenotype

<table>
<thead>
<tr>
<th>Tox phenotypea</th>
<th>Timeb</th>
<th>Top-ranked pathwaysc</th>
</tr>
</thead>
<tbody>
<tr>
<td>adverse at 29 days</td>
<td>1d</td>
<td>GO:cellular amino acid metabolic process (-0.7), GO:drug metabolic process (-0.6), GO:unsaturated fatty acid metabolic process (0.8)</td>
</tr>
<tr>
<td>bile duct hyperplasia</td>
<td>1d</td>
<td>GO:positive regulation of leukocyte apoptotic process (1.2), GO:cofactor metabolic process (-0.6), KEGG:Cell adhesion molecules (CAMs) (0.6)</td>
</tr>
<tr>
<td>cholesterol decrease</td>
<td>1d</td>
<td>C REACTOME:Genes involved in Apoptotic execution phase (1.8), GO:positive regulation of extrinsic apoptotic signaling pathway in absence of ligand (1.7), GO:regulation of triglyceride metabolic process (-1.7)</td>
</tr>
<tr>
<td>cholesterol increase</td>
<td>C</td>
<td>REACTOME:Genes involved in Ethanol oxidation (0.7), GO:ethanol catabolic process (0.6), GO:canicular bile acid transport (0.5)</td>
</tr>
<tr>
<td>hematopoeisis</td>
<td>1d</td>
<td>GO:negative regulation of programmed cell death (1.3), REACTOME:Genes involved in Sulfur amino acid metabolism (-1.4), GO:dicarboxylic acid metabolic process (-1.1)</td>
</tr>
<tr>
<td>hypertrophy</td>
<td>1d</td>
<td>C REACTOME:Genes involved in Orc1 removal from chromatin (0.6), REACTOME:Genes involved in SCF(Skp2)-mediated degradation of p27/p21 (0.7), GO:ribosomal large subunit export from nucleus (1.5)</td>
</tr>
<tr>
<td>increased glycogen</td>
<td>C</td>
<td>PID:TNF receptor signaling pathway PMID:18832364 (-1.6), GO:cellular response to lipoprotein particle stimulus (-0.9), GO:intrinsic apoptotic signaling pathway in response to DNA damage (-1.5)</td>
</tr>
<tr>
<td>increased mitosis</td>
<td>1d</td>
<td>KEGG:Pyruvate metabolism (1.4), KEGG:Propanoate metabolism (1), GO:single-organism catabolic process (0.6)</td>
</tr>
<tr>
<td>increased mitosis</td>
<td>C</td>
<td>GO:nuclear ubiquitin ligase complex (1.6), PID:Aurora A signaling PMID:18832364 (1.1), GO:homologous chromosome segregation (1)</td>
</tr>
<tr>
<td>necrosis</td>
<td>C</td>
<td>GO:macrophage chemotaxis (0.8), GO:myeloid leukocyte migration (0.5), REACTOME:Genes involved in Activation of Genes by ATF4 (1), GO:endoplasmic reticulum unfolded protein response (1.1), PID:ATF-2 transcription factor network PMID:18832364 (0.6)</td>
</tr>
<tr>
<td>single cell necrosis</td>
<td>C</td>
<td>BIOCARTA:Apoptotic DNA fragmentation and tissue homeostasis (-1.3), GO:pronucleus (-1), GO:protein depolymerization (-0.9)</td>
</tr>
<tr>
<td>trigs decrease</td>
<td>C</td>
<td>KEGG:Steroid biosynthesis (-0.3), GO:sterol esterification (0.7), GO:response to drug (-0.4)</td>
</tr>
<tr>
<td>trigs increase</td>
<td>C</td>
<td>KEGG:Steroid biosynthesis (0.7), GO:isoprenoid biosynthetic process (0.7), GO:extracellular matrix assembly (-0.8)</td>
</tr>
</tbody>
</table>

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a Representative toxicity phenotype as defined in methods; “adverse at 29 days” is an aggregate endpoint and included any treatments that produced one or more adverse morphologic changes at 29 days of dosing; only relationships at 1 day (1d) were evaluated. b Whether expression analysis is performed from samples collected after 1 day (1d) or concurrent with the observed toxicity phenotype (C). c Showing the top 3-ranked pathways for the toxicity phenotype and time, ranked using p-adj per (Sutherland et al., 2017), selecting only the most highly ranked pathway within each cluster. The value in parentheses is the pathway’s coefficient in the logistic regression model, interpreted as the natural
log of odds-ratio for observing toxicity given a one unit increase in the pathway score. The sign indicates whether induction (positive) or repression (negative) associates with increased odds of toxicity. A coefficient value of 0.69 corresponds to 2x odds of toxicity, 1.1 corresponds to 3x, etc. The full results are provided in Dataset S2.

**Omeprazole-induced liver hypertrophy**

In addition to their use for predictive applications, the relationship between pathway perturbation and toxicity phenotypes can be used to identify putative mechanisms linked to an observed toxicity phenotype. When studying a compound that causes a given toxicity phenotype (e.g. hepatocellular hypertrophy), this allows a user to observe whether pathways highly ranked by expression are also highly ranked in their association with the toxicity phenotype: “1) this new compound causes hypertrophy and induces pathway \(x\); 2) pathway \(x\) associates with hypertrophy across many other compounds; 3) therefore pathway \(x\) may contribute to the occurrence of hypertrophy for this new compound.”

Hepatocellular hypertrophy is frequently observed in toxicity studies and is generally considered an adaptive response to molecule exposure. When hypertrophy is observed, understanding its putative causes may help rationalize its relationship to the compound’s pharmacological effects and/or relevance in other species. Administration of omeprazole at 300 and 1000 mg/kg caused hypertrophy at 4 days or more of exposure (TG histology results). To identify potential mechanisms linked to hypertrophy, we identified 46 non-redundant gene sets having an adjusted \(p\)-value of < 0.001 at both doses, 24 hours after administration of a single dose (Figure 3; Dataset S3). The significantly-perturbed gene sets encompass a wide range of biological functions, as underscored by their distribution throughout the pathway map (recall that gene sets that tend to be perturbed by the same experiments are nearby on the map). We subsequently filtered these to the top 50 most strongly associated with hypertrophy across the entire TG database (i.e. exhibit a statistically significant relationship vs. the phenotype across a wide range of conditions; Dataset S2, \(p\)-adj). This led to the identification of 8 significant gene sets,
including glutathione metabolism, proteasome complex, DNA replication and mitotic cell cycle gene sets. These results are consistent with reports indicating that omeprazole induces Nrf2 mediated response to oxidative stress. Thus, filtering a larger list of differentially expressed gene sets against those associated with the observed morphologic changes facilitates the process of identifying molecular processes causally linked to injury.

Comparison of non-redundant gene sets vs. TXG-map modules

Gene sets from GO terms and pathways represent biological processes as they are understood and curated from the biomedical literature (henceforth, “pathways”). Notably, whether or not a gene is transcriptionally active in response to stress is not a consideration in associating genes with terms or pathways. Several approaches have been described for identifying putative biological networks from gene expression data, with no reference to pathways. TXG-MAP modules (Sutherland et al., 2017) were obtained by analyzing co-expression patterns of genes across the DM liver database. Biological processes driving co-expression of genes in response to treatments that are not represented in pathways are identified, but those drivers may be difficult to discern from the constituent genes and interpretation is more challenging than pathway analysis.

The CTox web application allows users to analyze their experiments using pathways and co-expression networks. To study the extent to which pathways recapitulate analysis using co-expression modules, and vice-versa, each TG liver experiment was analyzed using the non-redundant pathways and TXG-MAP modules. When comparing module and pathway scores across 3528 TG experiments, a small proportion of modules are observed to be highly correlated with pathways and therefore summarize similar effects (e.g. module 42m and KEGG glutathione metabolism; Figure 4A). Overall, only 16% of TXG-MAP modules capture co-expression behavior well-represented by pathways (defined as having similarity ≥ 0.7; Figure 4B). Conversely, only 20% of non-redundant pathways are well-represented by
modules. Modules and pathways are similarly predictive of concurrent toxicity phenotypes, whereas modules outperform for predictive applications (i.e. analyzing samples at 1 day to predict later toxicity; Figure 5). Hence, orthogonal approaches are more likely to lead to mechanistic insights for a treatment of interest.

**Identification of similar experiments**

Compounds that exhibit similar effects on the transcriptome may share pharmacological and/or mechanistic effects, an observation that underpins CMap and related approaches (Lamb et al., 2006; Smalley et al., 2010). Similarity of transcriptional responses can be assessed by describing the effects of each compound as a vector of pathway (GSA) or module scores (e.g. a vector of 382 pathway scores for the 382 non-redundant pathways). Taking each pair of TG liver experiments, we compared their similarity using the non-redundant pathways vs. TXG-MAP modules. Experiment similarities between pathway and modules scores were modestly correlated (Figure 6A; Pearson R = 0.59). Using the average similarity from non-redundant pathways and TXG-MAP modules, most pairs of highly similar experiments involved the same compound at different doses or time points, or comparisons of different compounds in the same pharmacological class (e.g. NSAIDs, PPAR modulator; Figure 6B). High similarity pairs of experiments involving compounds in different pharmacological classes may indicate that they share previously unrecognized pharmacological activity (or similar mechanisms of toxicity).

**Uricosuric agents benzbromarone and benziodarone are PPAR modulators**

Among TG liver experiment pairs having similarity > 0.9, a high proportion were PPAR modulators (clofibrate, fenofibrate, gemfibrozil, pirinixic acid, rosiglitazone). In addition, we noted 5 pairs of experiments where one agent was a PPAR modulator and the other was one of the uricosuric agents benzbromarone or benziodarone, commonly used to treat gout. A further 95 such pairs were
found in the next lower similarity range from 0.8 to 0.9. Compared to the lower similarity ranges, this represented a significant enrichment in prevalence of PPAR-uricosuric agents (4.6% or 100 of 2887 pairs with ≥ 0.8 similarity, vs. 0.2% of 2.6M pairs with < 0.8 similarity; p = <2e-16, χ² test). The 100 pairs included 12 benzbromarone or benziodarone experiments, with treatment duration ranging from 9 hours to 29 days (Dataset S4). This suggested that benzbromarone and benziodarone are PPAR modulators.

Clinical chemistry and histology changes for the 12 benzbromarone/benziodarone experiments included 6 causing >50% decrease in triglyceride levels and 7 causing hepatocellular hypertrophy. The 39 PPAR experiments that comprise the 100 high similarity pairs also lower triglyceride levels, without causing hepatocellular hypertrophy. However, most caused hepatocellular necrosis, not observed with benzbromarone/benziodarone (Dataset S4).

To further validate triglyceride-lowering mechanistic effects, we evaluated pathway and module scores for the 6 uricosuric agents and 26 PPAR modulator experiments that caused >50% decreases in triglyceride levels, ranking them from most to least perturbed across agents of each class (Table 3). As expected from global similarity of gene expression profiles, the most highly induced pathways and modules for uricosuric agent experiments were also highly ranked for PPAR modulators. In addition, most of the gene sets and modules had strong association with triglyceride lowering in general but not hypertrophy. The utility of considering induction/repression in the experiments of interest and the general association with the tox phenotype can be seen for module 285; it is induced across the uricosuric agents and PPAR modulators but ranks poorly in associating with triglyceride lowering in general. Hence, it may not play a role in the observed phenotype.
Table 3. Most perturbed modules and pathways by the uricosuric agents benzbromarone and benziodarone, and relationship to observed toxicity phenotypes

<table>
<thead>
<tr>
<th>gene set</th>
<th>urico agent expression rank</th>
<th>PPAR mod expression rank</th>
<th>q-adj decreased trigs</th>
<th>rank decreased trigs</th>
<th>q-adj hypertrophy</th>
<th>Rank hypertrophy</th>
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<tr>
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<td>1</td>
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<td>20</td>
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<td>195</td>
</tr>
<tr>
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<td>0.0000064</td>
<td>21</td>
<td>9.2E-08</td>
<td>130</td>
</tr>
<tr>
<td>GO negative regulation of oxidative stress-induced intrinsic apoptotic signaling pathway</td>
<td>3</td>
<td>9</td>
<td>5.06E-08</td>
<td>26</td>
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<tr>
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<tr>
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<td>0.0000429</td>
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<td>51</td>
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<td>REACTOME Genes involved in Metabolism of lipids and lipoproteins</td>
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<td>6.16E-13</td>
<td>7</td>
<td>0.00000173</td>
<td>92</td>
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</table>

*a each pathway or module was ranked from most to least perturbed (absolute value of pathway/module score) within each of 6 uricosuric agent and 26 PPAR modulator experiments, and the rank averaged across the 6/26 experiments; b the adjusted q-value indicating the pathway or module’s general association with the phenotype across all TG-GATEs experiments, and c the corresponding rank for the phenotype. These are “concurrent” associations, since expression results were taken from samples where the phenotype is present. Trigs = triglycerides. Full results are provided in Dataset S5.

Our results strongly suggest that benzbromarone and benziodarone are PPAR modulators in rodent liver. Others have investigated the PPAR activity of benzbromarone, confirming our findings in rodents (Kunishima et al., 2007; Lee et al., 2016) and human patients being treated for gout (Inokuchi et al., 2009).
Comparing results from RNA-seq and microarray studies

Users of the CTox application may upload results obtained by microarray or RNA-seq analysis. Several studies support the validity of comparing results from different measurement technologies (Wang et al., 2014; Rao et al., 2019; Black et al., 2014). To further corroborate these conclusions, we loaded into the application results from two studies: 1) RNA-seq from the Sequencing Quality Control (SEQC) consortium (Wang et al., 2014), and 2) recently published Affymetrix microarray and RNA-seq results generated concurrently from the same samples (Abvie study) (Rao et al., 2019) (Dataset S6). For the SEQC comparison, the median Pearson R expression similarity was 0.80 and 0.86 for non-redundant pathways and TXG-MAP modules, respectively. For the Abvie study, the median expression similarity was 0.82 and 0.86 for pathways and modules, respectively. These results describe concordance across measurement technologies (i.e. same tissue samples, different technology). By comparison, expression similarity assessed from samples generated by different laboratories using the same measurement technology and treatment conditions are notably lower (Sutherland et al., 2016). Consistent with our prior results (Sutherland et al., 2016), the level of agreement between microarrays and RNA-seq increases as the level of expression perturbation increases (Dataset S6). These results support the validity of comparing results obtained using microarray and RNA-seq in the CTox application.

Discussion and conclusion

Numerous reports and algorithms have been described for predicting and understanding mechanism of liver injury using transcriptomic results from nonclinical safety studies as a means to identify relevant mechanistic information. However, nearly two decades after transcript profiling was widely available for use by toxicity researchers in academia and industry, barriers to fuller adoption of
Toxicogenomic approaches in risk assessment remain. A notable technical barrier is the need for computational skills held by a minority of researchers in the field. Other technical challenges include the difficulty of implementing methods described in scientific reports without having access to computer source code or detailed protocols. In addition, analysis usually relies on one method rather than an integrated approach using multiple analysis methods. These (and other) barriers contribute to challenges in assuring the reproducibility of results. Finally, gaining greater acceptance of the significance of particular findings (e.g. induction of the glutathione metabolism pathway) requires an understanding of results for one compound in a broad context and rapid interpretation, preferably by the toxicologist doing the risk assessment analysis.

To facilitate progress towards these goals, we have created a collaborative toxicogenomics application (CTox) that allows non-computational specialists to evaluate liver gene expression results for their studies. Users can describe their experiments, upload the corresponding samples, evaluate their results using a variety of established and emerging systems biology analysis methods, compare them to the extensive DM and TG repositories, and share them with other researchers. Users can also search for and analyze data from those repositories using the same analysis methods. Analyses in other public applications are enabled, via exports of suitably formatted data. The CTox software is hosted in the Amazon Web Services cloud for use by the scientific community, or can be downloaded and installed within organizations’ private networks. Finally, the source code is publicly available on the GitHub.com repository, and all software components are available via open source licenses. This enables the extension and improvement of the application by others without licensing barriers.

To illustrate the utility of the CTox application, we described its use for identifying pathways predictive of an adaptive response in liver after treatment with omeprazole. In this case there was good overlap between modules and GO terms and canonical pathways that are predictive of hypertrophy. We also elucidated a putative role of PPAR modulation by the uricosuric agents benzbromarone and
benziodarone, in explaining their effects in rat liver. We demonstrated the importance of combining multiple lines of investigation to arrive at a small number of gene sets and/or co-expression modules (“features”) putatively linked to the observed phenotype: 1) identify features consistently affected by the treatment across doses and/or timepoints, 2) identify the subset of those features robustly associated with the phenotype across a variety of treatments, and, 3) corroborate if possible in molecules showing overall similar expression effects compared to the treatment of interest. In the absence of known toxicity study outcomes for a compound of interest, the subset of affected features with statistically significant relation to one or more toxicity phenotypes can be used to understand the odds of toxicity for the molecule of interest. These analyzes are possible either in the application itself, or by exporting small subsets of the data and further manipulating them in Excel.

We seek to further develop the application in collaboration with the toxicology community. Improvements to the suite of visualization tools will permit more detailed analysis within the application itself, such as exploring the connection between co-expression networks, canonical gene sets and their linkage to gene-level results. Also, we are actively implementing other analysis methods, such as the benchmark dose (BMD) methodology to identify points of departure for multi-dose expression studies. Finally, while the application can be used to analyze data from any system, we are developing additional tissue-specific non-redundant gene set collections and co-expression modules, and annotating their association to tissue injury (heart and kidney).

In summary, we have developed an open-source toxicogenomics analysis application which helps to increase accessibility, transparency and collaboration between researchers in the field. The application is available to the scientific community at http://ctox.indianabiosciences.org.
Data availability

Supporting Datasets S1-S6 are available at doi:10.5061/dryad.159h65k

Funding

This work was supported by financial contributions of Corteva Agriscience and Eli Lilly and Company to the Indiana Biosciences Research Institute.

Acknowledgements

The authors wish to thank Steve Evans, Bino John, Jon Klinginsmith, Derek Marren and Terry Wright for helpful discussions during the course of this work, and Meeta Pradhan for contributions towards coding microarray and RNA-seq fold change calculation scripts.

Conflicts of interest

The authors declare no conflicts of interest

References


**Figure legends**

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Figure 4. Comparison of TG liver experiment scores via non-redundant pathways and TXG-MAP modules. A) When comparing scores for 3528 TG experiments, the most similar pathway for module 42m was KEGG glutathione metabolism, and vice versa, with Pearson R = 0.82. B) The analysis in A) was repeated for each module, identifying the most similar pathway, and assigning the calculated Pearson R values to ranges as shown; stacked bar graphs show the cumulative distribution of similarities across the 415 modules. Likewise, each pathway was compared to modules, the most similar identified and assigned to similarity ranges in the same manner.

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