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Cytokine-Associated Drug Toxicity in Human Hepatocytes Is Associated with Signaling Network Dysregulation

Benjamin D. Cosgrove^{1,2,3,6,9}, Leonidas G. Alexopoulos^{1,3,4,7,9}, Ta-chun Hang^{1,2,3}, Bart S. Hendriks^{5,8}, Peter K. Sorger^{1,3,4}, Linda G. Griffith^{1,2,3}, and Douglas A. Lauffenburger^{1,2,3}

¹Department of Biological Engineering, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA

²Biotechnology Process Engineering Center, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA

³Cell Decision Processes Center, Massachusetts Institute of Technology, Cambridge, Massachusetts, USA

⁴Department of Systems Biology, Harvard Medical School, Boston, Massachusetts, USA

⁵Pfizer Research Technology Center, Cambridge, Massachusetts, USA

Abstract

Idiosyncratic drug hepatotoxicity is a major problem in pharmaceutical development due to poor prediction capability of standard preclinical toxicity assessments and limited knowledge of its underlying mechanisms. Findings in animal models have shown that adverse effects of numerous drugs with idiosyncratic hepatotoxicity in humans can be reproduced in the presence of coincident inflammatory cytokine signaling. Following these observations, we have recently developed an *in vitro* drug/inflammatory cytokine co-treatment approach that can reproduce clinical drug hepatotoxicity signatures -- particularly for idiosyncratic drugs -- in cultured primary human hepatocytes. These observations have suggested that drug-induced stresses may interact with cytokine signaling to induce hepatic cytotoxicity, but the hepatocyte signaling mechanisms governing these interactions are poorly understood. Here, we collect high-throughput phosphoprotein signaling and cytotoxicity measurements in cultured hepatocytes, from multiple human donors, treated with combinations of hepatotoxic drugs (*e.g.* trovafloxacin, clarithromycin) and cytokines (tumor necrosis factor- α , interferon- γ , interleukin-1 α , and interleukin-6). We demonstrate, through orthogonal partial least-squares regression (OPLSR) modeling of these signal-response data, that drug/cytokine hepatic cytotoxicity is integratively controlled by four key signaling pathways: Akt, p70 S6 kinase, MEK-ERK, and p38-HSP27. This modeling predicted, and experimental studies confirmed, that the MEK-ERK and p38-HSP27 pathways contribute pro-death signaling influences in drug/cytokine hepatic cytotoxicity synergy. Further, our four-pathway OPLSR model produced successful prediction of drug/cytokine hepatic cytotoxicities across different human donors, even

Corresponding author Douglas A. Lauffenburger Building 16, Room 343 Massachusetts Institute of Technology 77 Massachusetts Avenue Cambridge, MA, 02139, USA 617-252-1629 (tel), 617-258-0204 (fax), lauffen@mit.edu.

⁶Present address: Molecular Imaging Program at Stanford, Stanford University School of Medicine, Stanford, California, USA

⁷Present address: Department of Mechanical Engineering, National Technical University of Athens, Athens, Greece

⁸Present address: Merrimack Pharmaceuticals, Cambridge, Massachusetts, USA

⁹These authors contributed equally to this work.

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though signaling and cytotoxicity responses were both highly donor-specific. Our findings highlight the critical role of kinase signaling in drug/cytokine hepatic cytotoxicity synergies and reveal that hepatic cytotoxicity responses are governed by multi-pathway signaling network balance.

Keywords

partial least-squares regression (PLSR); inflammation; xenobiotic; adverse drug reactions; signaling network; predictive data models

Introduction

Hepatotoxicity[†] is a major cause of failures in the pre-clinical, clinical and post-approval stages of drug development and thus represents a major challenge for the pharmaceutical industry^{1, 2}. Idiosyncratic drug hepatotoxicity -- a hepatotoxicity subset that occurs in a very small fraction of human patients (~1 in 10,000) and accounts for ~10% of acute liver failure cases -- is poorly predicted by standard preclinical models and in clinical trials and frequently leads to post-approval drug failure³. Following the establishment of bacterial liposaccharide (LPS)-administered rodent models to assess inflammation-associated idiosyncratic drug hepatotoxicity^{3, 4}, we have previously developed an drug/inflammatory cytokine co-treatment cell culture approach for studying inflammation-associated drug hepatic cytotoxicity *in vitro*⁵. Our approach demonstrated that numerous drugs with clinical findings of idiosyncratic chronic hepatotoxicity, but not their non-toxic control compounds, exhibit synergistic induction of acute hepatocellular death in multiple hepatocyte cell culture models co-dosed with drugs and cytokines at concentrations near their upper limits of clinical liver exposure. In that work, we showed that the drugs exhibiting inflammation-associated toxicities induce mild hepatocellular stresses that synergize with cytokine signaling to elicit hepatocellular death³. A novel concept speculated upon therein was that signaling network dysregulation underlies cooperative drug/cytokine hepatic cytotoxicity, and we now undertake direct examination of that hypothesis in this study.

The factors identified as potentiating inflammation-associated drug hepatic cytotoxicity (LPS and the cytokines tumor necrosis factor- α [TNF], interferon- γ [IFN- γ], interleukin-1 α [IL-1 α], and interleukin-6 [IL-6]) . in these animal and cell culture models stimulate a diversity of intracellular signaling pathways related to hepatocyte survival, stress, and apoptosis⁶⁻¹¹ (Fig. 1). For example, TNF activates the MEK-ERK, IKK-NF- κ B, JNK, and p38-HSP27 signaling pathways and the caspase cascade, whose integrated activities specify hepatocyte death responses to TNF^{6, 12}. Further, physiologically relevant growth factor co-stimuli such as insulin can provide activation of survival signaling pathways that antagonize cytokine-induced apoptosis^{13, 14} and modulate hepatocyte metabolism of xenobiotic compounds¹⁵. How these pathways regulate drug/cytokine toxicity outcomes in hepatocytes is not clear, but there is evidence that, in a LPS-administered mouse model, hepatotoxicity synergy between LPS-induced cytokine release and the idiosyncratic hepatotoxicant ranitidine is dependent on p38-mediated TNF release in the resident liver macrophages, Kupffer cells^{11, 16-18}. Drug-induced hepatocellular stresses can induce activation of many of the signaling pathways stimulated by cytokines and/or growth factors (Fig. 1). Glutathione depletion can perturb cellular nutrient levels, possibly leading to attenuation of mTOR signaling^{19, 20}, and can alter mitochondrial redox potentials, increasing susceptibility to mitochondria-mediated apoptosis²¹. Reactive oxygen species accumulation can activate the JNK and p38-HSP27 stress-response signaling pathways⁶. Consequently, we hypothesize that further identification of the signaling

[†]We define "hepatotoxicity" as *in vivo* liver toxicity, largely arising from hepatocyte injury and/or death. We define "hepatic cytotoxicity" as *in vitro* and *in vivo* hepatocyte injury and/or death.

mechanisms governing hepatocyte cell death responses to hepatotoxicants and inflammatory cytokines would benefit from a network-level examination of multiple intracellular signaling pathways.

Here, we use orthogonal partial least-squares regression (OPLSR) data modeling to analyze a hepatic cytotoxicity data compendium, spanning diverse drug/cytokine stimulation combinations and containing measurements of 17 phosphoproteins from multiple signaling pathways and a cell toxicity readout, in cultured human hepatocytes. OPLSR modeling indicated that the activation profiles of four signaling pathways -- Akt, p70 S6K, MEK-ERK, and p38-HSP27 -- are sufficient to accurately predict toxicity responses to drug/cytokine co-treatments. Model predictions of pro-death contributions from p38-HSP27 and, counter-intuitively, MEK-ERK signaling were confirmed using small molecule kinase inhibitors. This work highlights the importance of a multi-pathway signaling network state for characterizing hepatic cytotoxicity arising from drug/cytokine synergies.

Results

Drug/cytokine co-treatments elicit shared regulation of multiple phosphoprotein signaling pathways

To examine the signaling mechanisms regulating inflammatory cytokine-associated drug hepatic cytotoxicity, we collected a 'cue-signal-response' (CSR) drug- and cytokine-induced hepatic cytotoxicity data compendium in primary human hepatocytes (Fig. 2). Human hepatocytes (donor #1) were treated with 66 different combinations of 11 'drug' conditions (six compounds with inflammation-associated hepatic cytotoxicity, four corresponding 'comparison'[‡] compounds, and a DMSO control) and six 'cytokine' conditions (no cytokine, TNF, IL-1 α , IL-6, LPS, and a mix containing all three cytokines and LPS). The hepatotoxicants (*e.g.* trovafloxacin, nefazodone, nimesulide, clarithromycin) and cytokines were selected based on identification of their role in inducing synergistic hepatic cytotoxicity synergy in our previous large-scale drug/cytokine co-treatment screen⁵. This study focused on single drug and cytokine dosing concentrations (drugs at 100*plasma C_{max} ⁵, 22) -- rather than examining drug and/or cytokine dose-response effects -- in order to enable a diversity of combinatorial drug/cytokine treatments within the constraints of limited human primary hepatocyte samples. Bead-based phosphoprotein and LDH release assays were used to quantitatively assess signaling pathway activation and cell death, respectively. Seventeen phosphoproteins, including signaling components of the MEK-ERK, mTOR-p70 S6K, PI3K-Akt, IKK-NF- κ B, JNK, p38-HSP27, STAT3, STAT6, cell cycle regulatory, and DNA damage signaling pathways, were measured at both early (0 and 20 min) and delayed time-points (4, 24, and 48 hr) following drug and/or cytokine stimulation to capture a diversity of intracellular signaling dynamics. These pathways are downstream of the cytokine and growth factor stimuli examined, but to our knowledge have not been reported to be directly regulated by the drugs used in this study (Fig. 1).

Across this CSR compendium, it was evident that cytokine-only treatments induced early-phase (20 min-4 hr) activation of multiple signals, including p-MEK1, p-STAT3, p-I κ B- α , and p-p38. Many toxic drug-only treatments induced a late-phase (24-48 hr) down-regulation of certain pro-survival signals (*e.g.* p-Akt, p-p70 S6K; see also Fig. 3D-E). This late-phase down-regulation was not evident for all phosphoproteins, as some survival (*e.g.* p-ERK1/2) and stress (*e.g.* p-HSP27) signals had sustained activation under toxic conditions (see also Fig. 4C,F). Multiple drug/cytokine co-treatments (*e.g.* trovafloxacin \pm the three-cytokine/LPS mix)

[‡]'Comparison' compounds are defined as paired control compounds to hepatotoxicants with similar molecular target(s), clinical indication, and chemical structure but are non-hepatotoxic.

synergistically induced both sustained p38–HSP27 signaling and cell death (Fig. 1 [donor #1] and Fig. 4B,F [donor #2]). This paired synergy raised the possibility that a single phosphoprotein signal might unilaterally yield a predictive readout of cell death across the entire data compendium. However, we calculated the correlation (Pearson coefficients) between observed cell death at 48 hours post-drug/cytokine co-treatment and each 17 phosphoprotein signal levels, and even the most positively (p-HSP27 at t = 4 hr, R = 0.78) and negatively (p-Akt at t = 24 hr, R = -0.74) correlated single phosphoprotein was poorly predictive of measured cell death (Fig. S3). This demonstrated need for multivariate analysis of signal-response relationships present in drug- and cytokine-induced hepatic cytotoxicity.

Multi-pathway OPLSR modeling identifies key molecular signals associated with drug- and cytokine-induced hepatic cytotoxicity

A spectrum of modeling approaches^{23, 24} are suited for analysis of signal transduction networks and their relationships to phenotypic cellular responses to diverse stimuli, but many of these approaches require *a priori* knowledge of how cellular stimuli are connected to signaling pathways and how those pathways mechanistically govern cellular responses of interest. The limited understanding of how idiosyncratic hepatotoxicants and inflammatory cytokine signals interact to synergistically induce hepatocyte cell death motivates the use of data-driven modeling approaches such as partial least-squares regression (PLSR), decision trees, and multiple linear regression, which can identify novel connectivities between measured signaling pathways and cellular responses without requiring an *a priori* network structure. From these approaches, we chose here to employ a variant of partial least-squares regression, which has been previously successful in establishing novel signal-response relationships from high-throughput signaling and response data^{13, 23, 25-28}. PLSR can make predictions of cellular responses across diverse cell types and experimental conditions and be used to reduce complexity in multi-variate protein signaling data to identify the most critical signaling nodes.

To generate a multi-pathway model relating the observed signaling activities and cell death responses, we subjected the initial CSR data compendium to orthogonal partial-least squares regression (OPLSR). OPLSR modeling relates the measured phosphoprotein signaling data matrix (X) and the observed hepatocellular death responses (Y), each arrayed across all 66 treatment conditions, through a linear regression of the relationship $Y = X \cdot B$, where B is a vector of regression coefficients that reflect how each phosphoprotein signal contributes to cell death. Following regression, the “fitness” of an OPLSR model can be assessed by comparing how the model “predicts” the cell death responses by multiplying the experimental signaling measurements with the model regression coefficients, and then calculating model fitness^{28, 29} by comparing the “predicted” and experimentally observed responses. We note that, in contrast to previous studies^{13, 25, 26, 28} implementing cell signaling-response PLSR modeling in cell line systems, our training data here comprises single biological replicates due to the severe limitations of human donor tissue samples; we chose to trade biological replication for a greater landscape of treatment conditions, within the constraints of this patient-oriented study.

Here, we found that an OPLSR model based of 102 phosphoprotein signaling metrics (17 phosphoproteins \times 6 time-dependent metrics; see *Experimental*) demonstrated very good model fitness ($R^2 = 0.92$) for cross-validated predictions of the observed cell death responses for all 66 drug/cytokine co-treatment conditions (Fig. 3A). Examination of the OPLSR model scores^{28, 30} projections (Fig. S2D) demonstrated the model's ability to distinguish between toxic and non-toxic treatment conditions by their scores in the first principal component (PC), which represents pro-death model contributions. The second PC of this OPLSR model captures remaining, orthogonal variation in the CSR data compendium irrelevant to response prediction.

OPLSR model loadings^{28, 30} ($w_1 \cdot c_1$; Fig. 3B and Fig. S4A) reflect the regression coefficients of the signaling metrics; positive model loadings imply signaling metrics with pro-death

contributions, and negative model loadings imply pro-survival contributions. Model variable-importance-of-projection (VIP) scores (Fig. 3B-C and Fig. S4B-C) reflect the relative importance of individual signaling metrics to the model fitness by capturing the loss in model fitness upon removal of their associated signaling metric^{28, 30, 31}. Metrics with VIP score >1 have significant importance in the model whereas metrics with a VIP score <<1 significantly lack unique information in the model. Inspection of model loadings and VIP scores identified four signaling pathways (MEK-ERK, Akt, p70 S6K, and p38-HSP27) with phosphoproteins having informative model contributions from four or more signaling metrics (Fig. 3B-C). p-ERK1/2, p-p38, and p-HSP27 were all identified as having significant pro-death contributions from multiple metrics, and p-Akt and p-p70 S6K were identified as having significant pro-survival contributions from multiple metrics. The identification of MEK-ERK signaling as having pro-death contributions by the model may seem surprising in light of this pathway's more commonly presumed pro-survival role, although this pathway has been recently found to similarly provide pro-death influence under inflammatory cytokine treatment in other human epithelial cell types³²; this effect is very clear in our study here given the sustained activation of MEK-ERK signaling evident under certain drug treatments (*e.g.* clarithromycin; Fig. 2 [donor #1] and Fig. 4C [donor #2]). For all 17 measured phosphoprotein, early time-point (20 min) metrics were uninformative as assessed by VIP scores (Fig. 2B and Fig. S4C), likely because these signaling activities were similarly activated by cytokines in both presence and absence of drug co-stimuli. In contrast, metrics from later time points (4-48 hr) and the average and integral metrics were, on average, more informative (Fig. 2B and Fig. S4C).

The distinct importance of these four signaling pathways suggested that many of the phosphoproteins in the full CSR data compendium were neglectable for model prediction, permitting model reduction to more easily interpretable subset of signals. To test this, phosphoproteins (and all six of their associated signaling metrics) were removed from the model stepwise in order of the lowest average VIP score across all six metrics (Fig. 3D). Model fitness was maintained until the top ~5 phosphoproteins remained ($R^2 = \sim 0.87-0.91$); further removal of phosphoproteins resulted in the significant losses in model fitness. A similar trend in loss of model fitness was observed by model reduction by removal of each of the 102 individual signaling metrics in order of the lowest VIP scores (Fig. S4D). Thus, an equivalently predictive multi-pathway network model could be generated by focusing on representative phosphoproteins from four pathways -- MEK-ERK, Akt, p70 S6K, and p38-HSP27 -- and representative phosphoproteins (p-ERK1/2, p-Akt, p-p70 S6K, and p-HSP27, with $R^2 = 0.89$; Fig. 3A,D) from these pathways can serve as a signaling 'network gauge'²⁵ (Fig. S5), whose combined activities specify hepatocellular death responses to drug/cytokine treatment.

Multi-pathway OPLSR model successfully predicts hepatic cytotoxicity signal-response relationships across human hepatocyte donors

In this study, human hepatocyte donors were garnered based on availability of sufficient viable cell quantities for assay demands. Donors with known obesity, alcohol use, or liver disease were excluded; availability limitations precluded matching for more general potential variables such as gender or age (Table S1). To test ability of the OPLSR model to make accurate predictions of hepatic cytotoxicity cell death responses to across human hepatocyte donors, we collected a second CSR hepatic cytotoxicity data compendium (in cells from donor #2) containing drug/cytokine co-treatments that were both present in the CSR from donor #1 and new (Fig. 4A). In the second CSR, human hepatocytes were treated with 14 different combinations of seven drug conditions (two hepatotoxicants used in initial compendium, four hepatotoxicants not used in the initial compendium, and a DMSO control) and two cytokine conditions (no cytokine and the three-cytokine/LPS mix). This second CSR focused on six phosphoprotein signals (p-MEK1, p-ERK1/2, p-Akt, p-p70 S6K, p-p38, and p-HSP27) from the four 'network gauge' pathways (Fig. 4A), allowing direct evaluation of reduced six-

phosphoprotein OPLSR model generated from the first CSR data compendium. We note that human donor variation is evident in the widespread differences observed in both phosphoprotein activation and cell death observed between donors #1 and #2 in response to the same drug/cytokine co-treatments (Fig. S6).

An OPLSR model trained on data from these six phosphoproteins from donor #1 was evaluated for its ability to make cell death predictions based on signaling data in donor #2 (Fig. 5A). The six-phosphoprotein model produced good quantitative predictions of new donor cell death responses for test data on treatments present in the training compendium. Predictions for the yet more stringent challenge of new donor cell death test data on treatments not present in the training compendium were gratifyingly accurate for some of the novel conditions (chlorpromazine \pm cytokine mix) though less so for other of the novel conditions. These results demonstrate that the six-phosphoprotein OPLSR model can generate successful *a priori* predictions of drug/cytokine responses across different human hepatocyte donors for conditions present in the training data set and even for some conditions beyond. For some of the drugs (*e.g.*, clomipramine, riluzole) that were not present in the training set, it is likely that other signaling pathways play a more substantial role in regulating responses drug/cytokine co-stimulation as these drugs induced cell death without significant activation of the four 'network gauge' signaling pathways compared to control treatments (Fig. 4A). We thus expect that building a broader training compendium and a network gauge model including even just a few more signaling pathways could further widen the scope of successful *a priori* prediction.

The ability of our 'network gauge' modeling approach to predict hepatic cytotoxicity signaling-response relationships across human hepatocyte donors for similar drug/cytokine conditions suggests that hepatocytes from multiple human donors share a 'common effector processing' function²⁶. In cells from two different human donors, particular drug/cytokine treatment conditions (*e.g.* clarithromycin \pm cytokine mix) elicited significantly different signaling network activation profiles and induced different levels of cell death (Fig. S6), but the OPLSR model nonetheless quantitatively captured their common signal-response relationship (Fig. 5A-B).

Multi-pathway OPLSR model accurately predicts experimental observations that MEK and p38 kinase inhibitors antagonize drug- and cytokine-induced hepatic cytotoxicity

The initial 17-phosphoprotein OPLSR model suggested that sustained activation of both the MEK-ERK and p38-HSP27 signaling pathways positively contributes to drug- and cytokine-induced hepatocellular death. Whereas the p38-HSP27 pathway is generally considered pro-apoptotic, due to transcriptional regulation of effector caspases³³, the MEK-ERK pathway has been considered generally pro-survival in hepatocytes through activation of anti-apoptotic effectors such as Bad³⁴ (Fig. S7). To test this complementary set of expected (p38) and surprising (ERK) predictions, MEK and p38 inhibitors were selected from panel of candidate small molecules for their ability to potently inhibit signaling while themselves inducing minimal toxicity in human hepatocytes (from donor #3; Fig. S8). The selected MEK (U0126 and PD325901) and p38 (PHA-666859 and PHA-818637) inhibitors were used to evaluate (both through model-prediction and experiment) the control of MEK-ERK and p38-HSP27 signaling pathways in drug- and/or cytokine-induced hepatic cytotoxicity.

First, we tested the ability of the six-phosphoprotein OPLSR model to predict effects of MEK and p38 kinase inhibitor on drug/cytokine-induced hepatic cytotoxicity. We reasoned that a most practically implementable challenge would be to predict effects on new donor hepatocytes from a model constructed on a previously-generated database from other human hepatocyte donors using previously-measured signaling inhibition efficacies. That is, given the 'common effector processing' paradigm, a conceivable application could be to use a signal-response model established across multiple previous subjects to predict *a priori* expected effects of

interventions on a newly-arising subject. Accordingly, we calculated signaling time-courses for the DMSO ± three-cytokine/LPS mix and the hepatotoxicant nortriptyline ± cytokine mix conditions using uninhibited signaling data from all conditions examined in donors #1 and/or #2 in concert with data for MEK and p38 inhibitors from donor #3 (Fig. S8). This calculation entailed decreasing the phosphoprotein levels of both the inhibitor's targeted kinase (*e.g.* p-p38) and its downstream substrate (*e.g.* p-HSP27) by the experimentally-determined proportions (*e.g.* 93% decrease for the p38 inhibitor PHA-666859; see Supplementary Experimental Methods). Since these drug/cytokine ± inhibitor conditions produced very low levels of cell death (Fig. 6 and Fig. S9), we logarithmically scaled the response data to remove regression bias towards high-response conditions (see Fig. S9A-B).

Using a log-scale OPLSR model trained on CSR data from donors #1 and #2, we predicted the cell death responses to multiple MEK and p38 inhibitors based on the signaling time-course perturbation method described above (Fig. 6A,C). As expected, the model predicted significant decreases in hepatic cytotoxicity induced by cytokine mix-only and nortriptyline + cytokine mix treatments upon MEK or p38 kinase inhibitor pre-treatment. In human hepatocytes (from donor #4) treated with cytokine mix-only or the idiosyncratic hepatotoxicant nortriptyline + cytokine mix, one or both of MEK and p38 kinase inhibitors reduced the observed hepatotoxicities (LDH release at $t = 48$ hr; Fig. 6B,D). MEK or p38 inhibition did not perturb hepatic cytotoxicity in DMSO control or nortriptyline only conditions (Fig. S9C-D). Quantitative accuracy of the model predictions (based on data from donors #1-3) was assessed by comparing to these experimental observations in donor #4 (Fig. S9). The *log*-scaled OPLSR model demonstrated promising model fitness for the training data ($R^2 = 0.79$), but unsatisfactory predictions of the inhibitor test data ($R^2 = 0.08$). However, using a Pearson correlation metric, the model generated well-correlated predictions of both training ($R = 0.90$) and inhibitor test ($R = 0.63$) data. Importantly, the model correctly predicted the experimentally observed pro-survival effects of MEK and p38 inhibition under these treatment conditions, confirming that both MEK-ERK and p38-HSP27 signaling indeed both contribute positively to the induction of hepatocellular death by drug/cytokine co-treatment. Moreover, this demonstrates that the reduced six-phosphoprotein OPLSR model is capable of at least qualitatively correct predictions of kinase inhibitor perturbations to drug/cytokine hepatic cytotoxicity in a practically implementable protocol.

Discussion

To date, systems-level analyses of drug-associated hepatotoxicity have largely focused on gene expression³⁵ or metabolomic³⁶ data. Our new work here demonstrates the utility of a systems modeling approach for predicting hepatic cytotoxicity in human hepatocytes based on phosphoprotein signaling data. Using orthogonal partial least-squares regression modeling of multi-pathway phosphoprotein data, we showed that four key signaling pathways are instrumental to the cooperative induction of hepatocellular death by hepatotoxicants and physiologically relevant inflammatory cytokine co-stimuli. This informative subset contained two pathways (Akt and p70 S6K) with model-assigned pro-survival functions, in agreement with their accepted survival signaling roles^{19, 37} and two pathways (MEK-ERK and p38-HSP27) with model-assigned pro-death functions. These four model-identified signaling pathways represent a useful 'network gauge'²⁵ capturing the balance between pro-survival and pro-death signaling in drug- and cytokine-stimulated hepatocytes (Fig. S5). Thus, we propose that this kind of hepatic cytotoxicity may be understood in terms of a signaling network state, in which the integrated behavior of multiple pathways characterizes the effects of a candidate drug in the context of diverse, physiologically relevant co-stimuli (Fig. 7).

Though p38 and JNK signaling pathways have been individually identified as critical regulators of liver injury associated with the hepatotoxicants ranitidine^{11, 16, 18} and acetaminophen³⁸,

respectively, this work demonstrates that, for diverse hepatotoxicant and cytokine co-stimulatory conditions, specification of hepatic cytotoxicity responses depends on the pro-survival and pro-death signaling contributions from the Akt, p70 S6K, MEK-ERK, and p38-HSP27 pathways. It is likely that other cytokine signaling pathways, including IKK-NF- κ B and JNK, are indeed critical to inflammation-associated idiosyncratic drug hepatotoxicity, but they may be more associated with regulation of the inflammatory response in cells other than hepatocytes and/or are crucial for mechanisms of toxicity not shared by the hepatotoxicants in this study. We note that the MEK-ERK and p38-HSP27 pathways highlighted by our present work may influence a variety of downstream effector mechanisms involved in potentially diverse observed hepatocyte injury behaviors⁵. Future investigation into the mechanisms connecting drug-induced hepatocellular injuries and kinase signaling activation should lead to improved prediction and understanding of inflammatory cytokine-associated idiosyncratic drug hepatic cytotoxicity.

We submit that our findings here motivates complementary future studies -- perhaps using lower drug and cytokine concentrations -- focused on how hepatocyte signaling networks govern sub-lethal hepatic toxicity phenotypes (such as mitochondrial and drug transporter dysfunction^{22, 39}) measureable in high-throughput cell culture that underlie chronic liver toxicity. Moreover, this type of patient-specific data-driven toxicity modeling could be implemented to generate personalized predictions of idiosyncratic hepatotoxicity based on well-trained, multi-donor *in vitro* signaling-hepatic toxicity models that are further supplemented by focused *in vitro* studies on patient-specific hepatocytes using patient-tailored drug doses and cytokine levels.

Our data-driven modeling analysis indicated that the MEK-ERK and p38-HSP27 pathways are activated in more sustained manner by hepatotoxic drug and/or cytokine treatments to positively regulate the resulting cell death response. These predicted pro-death signaling contributions were confirmed using small molecule MEK and p38 kinase inhibitors to perturb the cooperative hepatic cytotoxicity induced by nortriptyline and a cytokine mix. While in agreement with the generally appreciated apoptotic signaling role of p38 signaling³³, this result is surprisingly counter to the general view of MEK-ERK contributing to survival signaling^{32, 34}. One conceivable mechanism for the surprisingly similar pro-death consequences of MEK-ERK and p38-HSP27 pathway activation is induction of the cellular metalloproteinase TACE. Activated ERK⁴⁰ and p38¹⁶ can both positively regulate TACE activity and its proteolytic release of TNF⁴¹. TNF is a crucial pro-death cytokine in inflammation-induced drug hepatotoxicity *in vivo*^{10, 11}, where its TACE-regulated release by Kupffer cells is p38-dependent^{16, 18}, and in our *in vitro* drug/cytokine co-treatment system⁵. We speculate that sustained ERK and p38 activation by the initial drug/cytokine stimuli may lead to enhanced hepatocyte TNF production and autocrine signaling, which could lead to further pro-death signaling (for example, through prolonged p38 activation; see Fig. 4F). This potential mechanism merits dedicated study in future work.

Experimental

Human hepatocyte cell culture and stimulation

Primary human hepatocytes were obtained from CellzDirect (Durham, NC). Human hepatocytes from multiple donors were used in this study, and donor identification is noted for each data set. Donor information is provided in Table S1. Human hepatocytes were seeded on collagen-coated 12- or 96-well plates (BD Biosciences, Franklin Lakes, NJ) at 1.5×10^5 cells/cm² in medium containing 5% FBS (Hyclone, Logan, UT) and 5 μ g/ml insulin (Sigma, St. Louis, MO), and, one day post-seeding, were overlaid with 0.25 mg/ml Matrigel (BD Biosciences) in medium containing insulin but not FBS, as described previously⁵. One day following Matrigel overlay, fresh culture medium containing insulin but not FBS was added.

For kinase inhibition studies, this medium contained inhibitors at $2\times$ final concentration; one hour later, an equal volume of medium containing drugs and/or cytokines at $2\times$ final concentration was added. All drugs were suspended in 0.25% final DMSO. Drugs with associated idiosyncratic hepatotoxicity and corresponding ‘comparison’ compounds were dosed at a $100\times C_{max}$ concentration, a therapeutically appropriate dosing level (see Supplementary Experimental Methods). Recombinant human cytokines were obtained from R&D Systems (Minneapolis, MN) and were used at the following final concentrations: 100 ng/ml TNF, 20 ng/ml IL-1 α , and 20 ng/ml IL-6. LPS serotype 1 from *E. coli* 0111:B4 (Sigma) was used at 10 μ g/ml. The MEK inhibitors PD98059, U0126, and MEK inhibitor-1 (MEKi-1) and the p38 inhibitors SB202190 and SB203580 were obtained from EMD Biosciences (San Diego, CA). The MEK inhibitor PD325901 and p38 inhibitors PHA-666859 and PHA-818637 were obtained from Pfizer's chemical sample bank (Groton, CT). All inhibitors were suspended in 0.1% final DMSO. See Supplementary Experimental Methods for details on selection of inhibitor concentrations.

Multiplexed phosphoprotein assays

Cell lysates were collected at 0, 20 min, 4, 24, and 48 hr following drug/cytokine stimulation, clarified, and normalized using a bicinchoninic assay (Pierce). Bio-Plex bead-based assays (Bio-Rad, Hercules, CA) were conducted using a Luminex 200 instrument (Austin, TX), according to manufacturer's instructions, to quantify the following 17 phosphoproteins in cell lysates: p-Akt (Ser⁴⁷³), p-CREB (Ser¹³³), p-c-Jun (Ser⁶³), p-GSK-3 α/β (Ser²¹/Ser⁹), p-I κ B- α (Ser³²/Ser³⁶), p-IRS-1 (Ser⁶³⁶/Ser⁶³⁹), p-ERK1/2 (Thr²⁰²/Tyr²⁰⁴, Thr¹⁸⁵/Tyr¹⁸⁷), p-Histone H3 (Ser¹⁰), p-HSP27 (Ser⁷⁸), p-JNK (Thr¹⁸³/Tyr¹⁸⁵), p-MEK1 (Ser²¹⁷/Ser²²¹), p-STAT3 (Ser⁷²⁷), p-STAT6 (Tyr⁶⁴¹), p-p38 (Thr¹⁸⁰/Tyr¹⁸²), p-p53 (Ser¹⁵), p-p70 S6 kinase (Thr⁴²¹/Ser⁴²⁴), and p-p90 RSK (Thr³⁵⁹/Ser³⁶³). Phosphoprotein assays were validated for high specificity to known activating conditions (Fig. S1). See Supplementary Experimental Methods for additional details.

Lactate dehydrogenase cell death assay

At 20 min, 4, 24, and 48 hr post-stimulation, conditioned medium samples were collected to assay lactate dehydrogenase (LDH) release using a CytoTox-ONE Homogeneous Membrane Integrity Assay (Promega, Madison, WI) according to manufacturer's recommendations.

Orthogonal partial least-squares modeling of cue-signal-response data compendia

‘Cue-signal-response’ (CSR) data compendia containing phosphoprotein time-course ($t = 0$ –48 hr) and LDH release end-point ($t = 48$ hr) data were constructed separately for each human hepatocyte donor. Phosphoprotein data were fold-change normalized to untreated samples (at 0 min) for each phosphoprotein assay and separately for each hepatocyte donor. LDH release data were fold-change normalized to untreated samples at 48 hr post-stimulation separately for each hepatocyte donor. These normalized data are available in Supplementary Data. CSR data compendia normalization and fusion was performed using the DataRail toolbox⁴³ for Matlab (The Mathworks, Inc., Natick, MA). For each phosphoprotein signaling time-course, two time-dependent signaling metrics¹³ were extracted: (i) integral for the entire time-course, and (ii) the average of the late time-points (4–48 hr), reflecting the steady-state signaling level. These were added to the four time-points (20 min, 4, 24, and 48 hr) to yield six signaling metrics for each assayed phosphoprotein. For each compendium, the signaling metrics from all measured phosphoproteins were then fused into a signaling network data matrix (X), and the toxicity response data were cast into a vector (Y), with both X and Y arrayed across all treatment conditions.

The relationship between the signaling data matrix (X) and the toxicity response vector (Y) was modeled using partial least-squares regression (PLSR). Briefly, we implemented PLSR

using the NIPALS algorithm in SIMCA-P software (Umetrics, Inc., Kinnelon, NJ) following standard methods^{13, 25, 26, 28-30, 44}, with some modifications. Data matrices and vectors were mean-centered and unit-variance scaled. All models were generated using four principal components under standard optimization criteria³⁰. Model calibration was conducted using leave-one-out cross-validation, and model uncertainties were calculated by jack-knifing⁴². Calibrated models were subjected to a principal-component space linear transformation by rotating the projection of the single cell death response variable completely into the first principal-component, yielding an orthogonal PLSR (OPLSR) model⁴⁵ (see Fig. S2), to allow for simplified interpretation of model loadings and scores. Model loadings were calculated using the mean-centered regression coefficients $w_a \cdot c_a$ from the a -th OPLSR principal-component¹³. The information content of each signaling metric was assessed by its variable importance of projection (VIP) score, which captures the loss in model fitness upon removal of the individual signaling metric^{28, 30, 31}. The accuracy of model predictions for both training and test data were assessed using a model fitness parameter^{28, 29} (R^2) and Pearson correlation coefficient²⁵ (R). See Supplementary Experimental Methods for additional details.

Statistical analysis

For comparing two means, a Student's t test was used. All tests were performed at a significance level of $\alpha = 0.05$.

Statement for Graphical Abstract

Certain pharmaceutical compounds exhibit rare hepatocyte toxicities in humans that may be dependent on coincident inflammatory cytokine signaling but are poorly understood mechanistically. We employ a systems biology approach to identify a set of four kinase signaling pathways that govern these toxicities across human hepatocyte donors.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

Akt	serine-threonine protein kinase B/Akt
ASP	aspirin
BUS	bupirone
CHL	chlorpromazine
CIM	cimetidine
CLA	clarithromycin
CLO	clomipramine
C_{\max}	average plasma maximum drug concentration

CSR	cue-signal-response
DMSO	dimethyl sulfoxide
ERK	extracellular signal-regulated kinase
FBS	fetal bovine serum
GSH	glutathione
HSP27	heat shock protein 27
IFN- γ	interferon- γ
IKK	inhibitor of nuclear factor- κ B kinase
IL-1 α	interleukin-1 α
IL-6	interleukin-6
JNK	c-Jun N-terminal kinase
LDH	lactate dehydrogenase
LEV	levofloxacin
LPS	lipopolysaccharide
MEK	MAPK-ERK kinase
mTOR	mammalian target of rapamycin
NEF	nefazodone
NF- κ B	nuclear factor- κ B
NIM	nimesulide
NIPALS	non-linear iterative partial least-squares
NOR	nortriptyline
OPLSR	orthogonal partial least-squares regression
p38	p38 mitogen-activated protein kinase
p70 S6K	p70 S6 protein kinase
PC	principal component
PI3K	phosphoinositide 3-kinase
RAN	ranitidine
RIL	riluzole
ROS	reactive oxygen species
TACE	tumor necrosis factor- α -converting enzyme
TEL	telithromycin
TNF	tumor necrosis factor- α
TRO	trovafloxacin
VIP	variable-importance-of-projection

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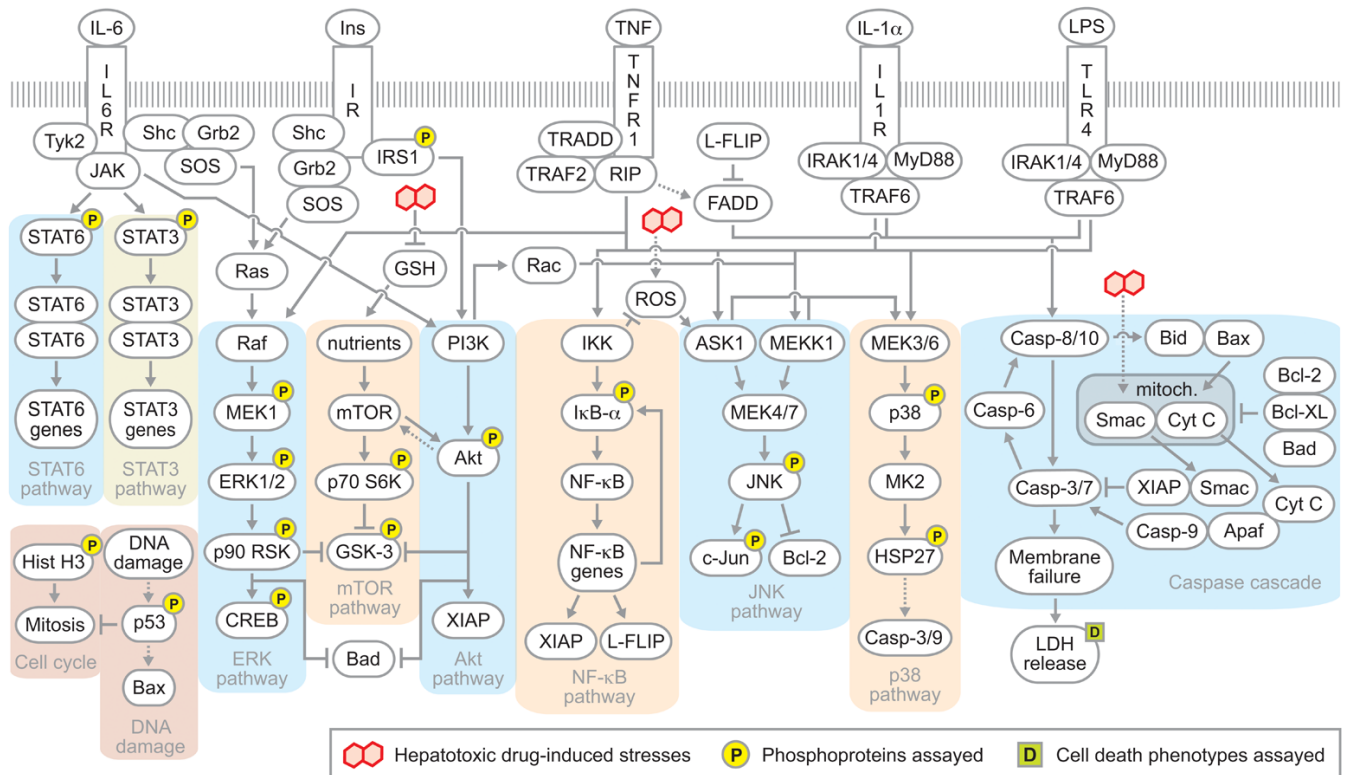


Figure 1.

A schematic of the drug- and cytokine-induced hepatocellular death signaling network, curated from literature references^{6-14, 19-21}. Inflammatory cytokines bind to their cognate receptors and activate shared downstream phosphoprotein signaling pathways regulating cell death. Hepatotoxic drugs (hepatotoxicants) lead to various cellular stresses (*e.g.* ROS accumulation) that are integrated into stress-related phosphoprotein signaling (*e.g.* JNK pathway) and mitochondrial control of apoptosis. Phosphoproteins and cell death phenotypes experimentally assayed in this study are indicated.

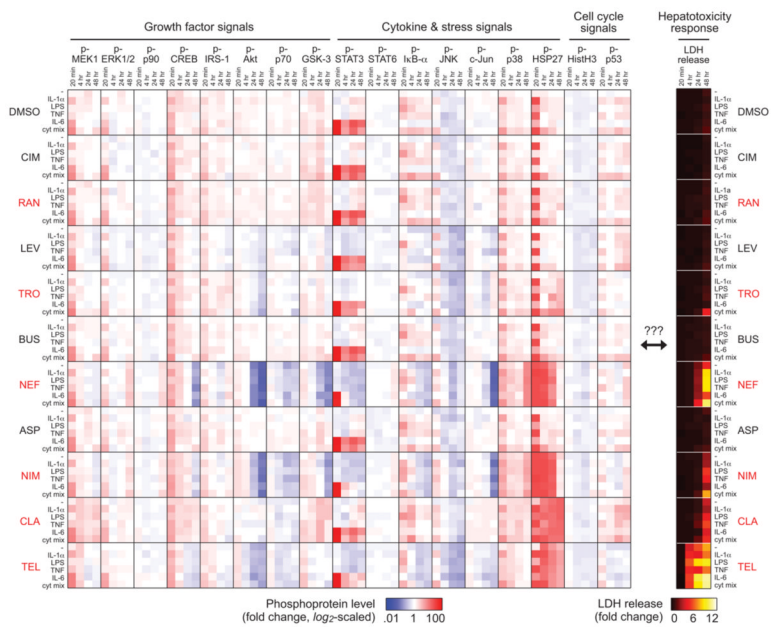
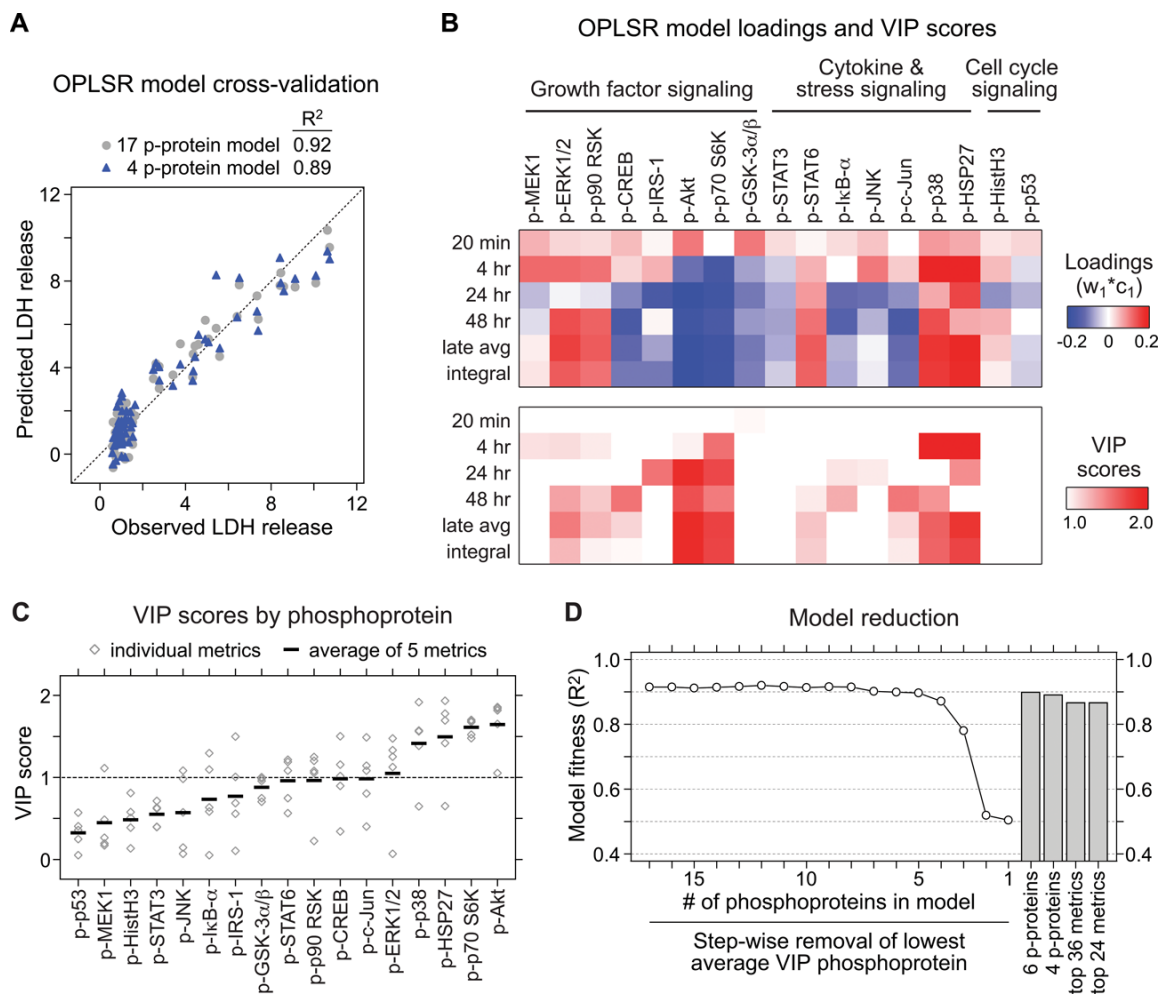


Figure 2. A cue-signal-response (CSR) drug/cytokine-induced hepatic cytotoxicity data compendium for model training. Cultured primary human hepatocytes (donor #1) were treated with 66 combinations of 11 ‘drug’ conditions (six idiosyncratic hepatotoxicants, four corresponding ‘comparison’ compounds, and a DMSO control) and six ‘cytokine’ conditions (no cytokine, 20 ng/ml IL-1 α , 10 μ g/ml LPS, 100 ng/ml TNF, 20 ng/ml IL-6, and a mix containing all three cytokines plus LPS). All drugs were dosed at 100* C_{max} concentrations (see *Experimental*): 1.5 mM cimetidine (CIM), 142 μ M ranitidine (RAN), 1.6 mM levofloxacin (LEV), 770 μ M trovafloxacin (TRO), 0.46 μ M buspirone (BUS), 86 μ M nefazodone (NEF), 552 μ M aspirin (ASP), 2.1 mM nimesulide (NIM), 334 μ M clarithromycin (CLA), and 277 μ M telithromycin (TEL). Idiosyncratic hepatotoxicants (red font) and corresponding ‘comparison’ compounds are presented in vertical juxtaposition. Seventeen phosphoproteins (left) and LDH release (right) were quantitatively assayed at both early (0 and 20 minutes) and delayed time-points (4, 24, and 48 hours) following drug and/or cytokine stimulation. Phosphoprotein levels were fold-change normalized to untreated samples at 0 minutes and are plotted on a log_2 -scaled colormap to capture both up- and down-regulated signaling levels. LDH release levels were fold-change normalized to untreated samples at 48 hours post-stimulation and are plotted on a linearly scaled colormap. Single biological replicates were used for both phosphoprotein and LDH assays.

**Figure 3.**

OPLSR modeling identifies key molecular signals regulating drug/cytokine-induced hepatic cytotoxicity. An OPLSR model was trained on the 66-condition CSR data compendium from human hepatocyte donor #1 to relate the 102 time-dependent phosphoprotein signaling metrics (17 phosphoproteins \times 6 metrics) to the cell death measurement (LDH release at $t = 48$ hr). (A) A correlation plot relating the observed and OPLSR model-predicted LDH release responses for all 66 drug/cytokine co-treatment conditions for the 17-phosphoprotein and a reduced 4-phosphoprotein (p-ERK1/2, p-Akt, p-p70 S6K, and p-HSP27) model both demonstrate good model fitness (R^2)^{28, 29}. (B-C) OPLSR model loadings and VIP scores from the 17-phosphoprotein model. Positive loadings contribute to and negative loadings antagonize the model's cell death prediction. VIP scores identify the relative importance of individual phosphoprotein signaling metrics, with the threshold value of 1 for informative VIP scores^{28, 31} used as the lower-limit of the color scale in (B) and is plotted in (C). In (D), VIP scores from individual metrics and averages for each phosphoprotein signal are plotted, with the 20-min metrics were omitted, as these metrics were uniformly uninformative. (D) Model fitness sensitivity to the removal of phosphoprotein signaling metrics. Model complexity was reduced using: (i) step-wise removal of the lowest average VIP phosphoprotein; (ii) by selecting four (p-ERK1/2, p-Akt, p-p70 S6K, and p-HSP27) or six (those four plus p-MEK1 and p-p38) phosphoproteins; or (iii) using an equivalent number of signaling metrics as the four- and six-phosphoprotein models (24 and 36 metrics, respectively) but selected from the top VIP scores across all phosphoprotein metrics.

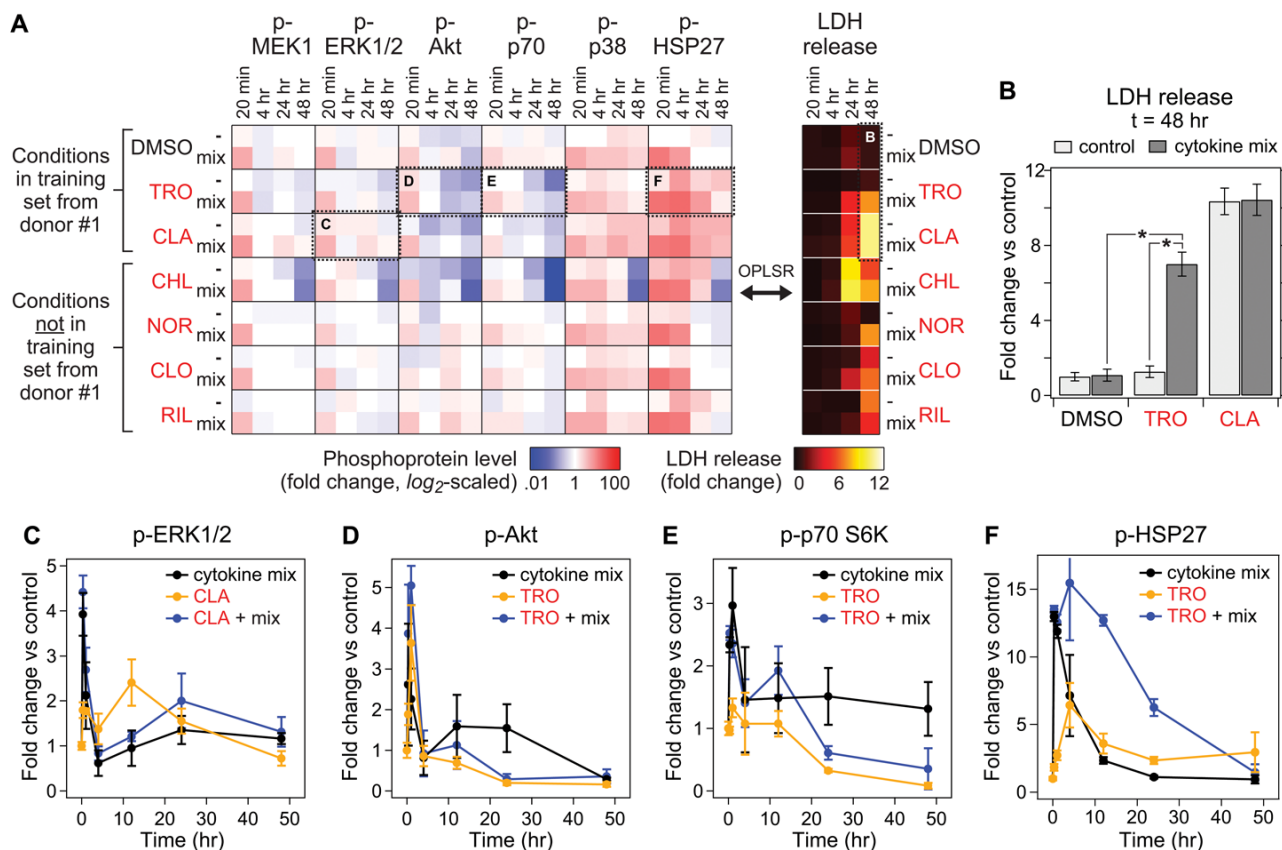
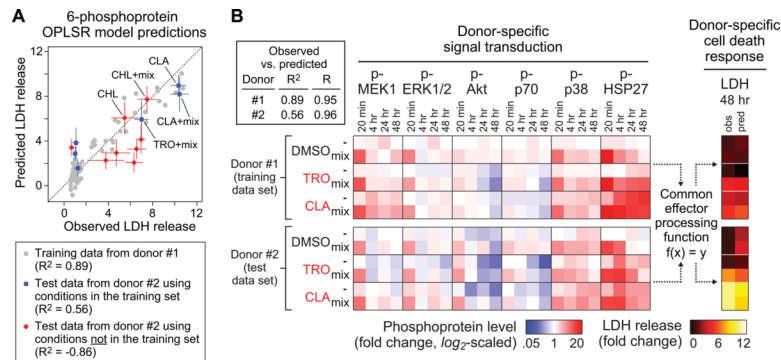


Figure 4.

A CSR drug/cytokine-induced hepatic cytotoxicity data compendium for model training. (A) Cultured primary human hepatocytes (donor #2) were treated with 14 combinations of seven ‘drug’ conditions (two idiosyncratic hepatotoxicants used in initial compendium, four idiosyncratic hepatotoxicants not used in the initial compendium, and a DMSO control) and two ‘cytokine’ conditions (no cytokine and the three-cytokine/LPS mix). Idiosyncratic hepatotoxicants (red font) were dosed at $100 \times C_{max}$ concentrations: 770 μ M trovafloxacin (TRO), 334 μ M clarithromycin (CLA), 111 μ M chlorpromazine (CHL), 10 μ M nortriptyline (NOR), 13 μ M clomipramine (CLO), and 107 μ M riluzole (RIL). Six phosphoproteins (p-MEK1, p-ERK1/2, p-Akt, p-70 S6K, p-p38, p-HSP27) and LDH release were assayed at both early (0 and 20 min) and delayed time-points (4, 24, and 48 hr) following stimulation. Some conditions were also assayed at 1 and 12 hr post-stimulation (see (C-F)). Biological triplicates were used for both phosphoprotein and LDH assays. Phosphoprotein and LDH release levels were normalized and plotted as in Fig. 2. (B) Trovafloxacin, but not clarithromycin, synergizes with cytokine mix co-treatment to induce hepatic cytotoxicity. Differences between drug/cytokine co-treatment and either drug-only or cytokine mix-only conditions are labeled (*) if $P < 0.05$ by a Student's t test. (C-F) Phosphoprotein signaling time-courses for idiosyncratic drug (TRO or CLA) and/or cytokine mix treatment demonstrate sustained p-ERK1/2 (C) and p-HSP27 (F) activation and late p-Akt (D) and p-p70 S6K (E) down-regulation under hepatotoxic conditions compared to cytokine mix-only controls. In (C-F), DMSO-only control treatments are not shown but are in (A). In (B-F), data is presented as mean \pm s.e.m. of three biological replicates.

**Figure 5.**

OPLSR model testing on CSR drug/cytokine-induced hepatic cytotoxicity data compendia in different human hepatocyte donors. An OPLSR model accurately predicts cell death responses across human hepatocyte donors for drug/cytokine treatments present in the training data compendium, even though donor-specific signaling network activation profiles and cell death responses were observed. (A) An OPLSR model of LDH release was trained on the time-dependent signaling metrics from six phosphoproteins (p-MEK1, p-ERK1/2, p-Akt, p-70 S6K, p-p38, p-HSP27) using data from donor #1. This trained model was used to predict LDH release responses based on the same signaling metrics measured in hepatocytes from donor #2. A correlation plot relating the observed and predicted LDH release responses for the both training data (donor #1) and the test data (donor #2), which contained six treatments present and eight treatments not present in the training data. Test conditions (CHL \pm cytokine mix) not present in the training data but nonetheless predicted accurately are noted. The model-predicted responses are presented as the mean prediction \pm cross-validation standard error, calculated by jack-knifing⁴². The test data is presented as mean \pm s.e.m. of three biological replicates. Experimental and prediction uncertainties are not shown for the training data. (B) The predictive accuracy of this OPLSR model suggests that a ‘common effector processing’ mechanism ($f(x) = y$), encompassing the integration of multiple signaling pathways (x) yields quantitatively concerted cell death responses (y), exists and is shared between hepatocytes from different human donors given similar drug/cytokine treatments. The inset tabulates model fitness (R^2) and Pearson correlation coefficient (R) values for the model predictions of these shared conditions. See Fig. S6 for expanded line plots comparing the phosphoprotein and LDH release data in donors #1 and #2.

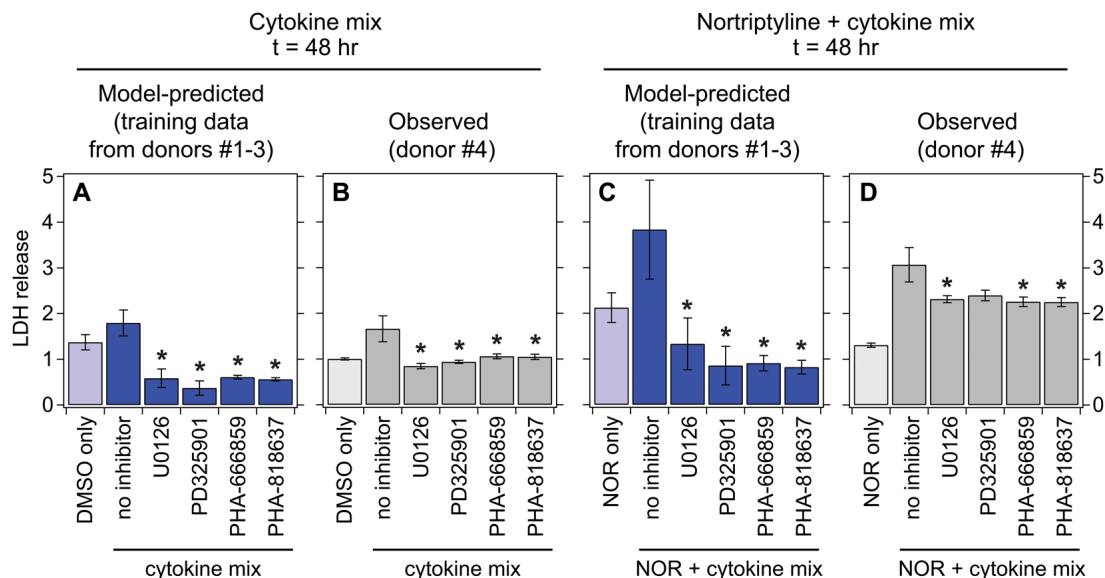


Figure 6.

Kinase inhibitor perturbation of MEK and p38 signaling confirms their model-predicted pro-death signaling regulation of drug/cytokine-induced hepatic cytotoxicity. (**A,C**) OPLSR model predictions of the effects of MEK and p38 inhibitors in perturbing drug/cytokine-induced hepatic cytotoxicity. To make predictions of the effects of kinase inhibitor perturbation on hepatocellular death responses, uninhibited signaling time-courses measured in cells from donors #1 and/or #2 were ‘computationally inhibited’ based on inhibitor activities measured in cells from donor #3 (Fig. S8). An OPLSR model trained on data from donors #1 and #2 was used to make cell death response predictions from these ‘computationally inhibited’ time-courses (Fig. S9A-B). Model predictions are presented as the mean \pm cross-validation standard error, calculated by jack-knifing⁴². (**B,D**) Experimental inhibition of MEK and p38 kinase activities demonstrates their pro-death regulation of cytokine mix- and nortriptyline (NOR) + cytokine mix-induced hepatic cytotoxicity. Cultured human hepatocytes (donor #4) were treated with DMSO control (**B**) or 10 μ M nortriptyline (**D**) \pm a three-cytokine/LPS mix and were assayed for LDH release 48 hr post-treatment. Cells were pretreated with a MEK inhibitor (10 μ M U0126 or 1 μ M PD325901) or a p38 inhibitor (1 μ M PHA-666859 or 1 nM PHA-818637) one hour before stimulation. Data are presented as the mean \pm s.e.m. of eight biological replicates. Kinase inhibition did not significantly perturb cell death responses in the absence of cytokine mix treatment (Fig. S9C-D). In (**A**) to (**D**), differences between to control and inhibitor conditions are labeled as significant (*) if $P < 0.05$ by a Student's t test.

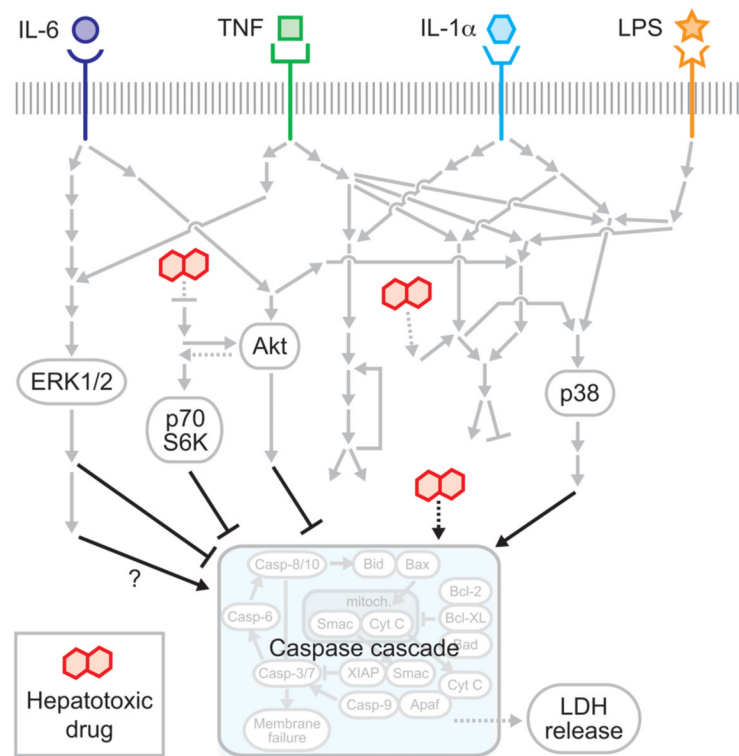


Figure 7. Inflammatory cytokine-associated idiosyncratic drug hepatic cytotoxicity as a ‘network toxicity’. The multi-pathway modeling approach presented here suggests that an integration of the activities of multiple intracellular signaling pathways -- namely ERK, Akt, p70 S6K, and p38 -- is necessary for hepatocytes to fully specify death responses to hepatotoxic drug/ cytokine co-treatment conditions. This provides motivation of the network-level consideration of multiple survival, stress, and apoptosis signaling pathways in evaluating the hepatic cytotoxicity mechanisms of context-dependent hepatotoxic drugs.