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Pharmacogenomic and clinical data link non-pharmacokinetic metabolic dysregulation to drug side effect pathogenesis

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Drug side effects cause a significant clinical and economic burden. However, mechanisms of drug action underlying side effect pathogenesis remain largely unknown. Here, we integrate pharmacogenomic and clinical data with a human metabolic network and find that non-pharmacokinetic metabolic pathways dysregulated by drugs are linked to the development of side effects. We show such dysregulated metabolic pathways contain genes with sequence variants affecting side effect incidence, play established roles in pathophysiology, have significantly altered activity in corresponding diseases, are susceptible to metabolic inhibitors and are effective targets for therapeutic nutrient supplementation. Our results indicate that metabolic dysregulation represents a common mechanism underlying side effect pathogenesis that is distinct from the role of metabolism in drug clearance. We suggest that elucidating the relationships between the cellular response to drugs, genetic variation of patients and cell metabolism may help managing side effects by personalizing drug prescriptions and nutritional intervention strategies.

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A dverse drug reactions, commonly known as side effects, are thought to be responsible for as much as 11% of hospital admissions\(^1\), a fifth of both phase II\(^1\) and III\(^4\) clinical trial failures, high-profile market withdrawals (for example, Vioxx, Lipobay), and a large fraction of patient therapeutic non-compliance incidents\(^5\). Risk factors associated with side effects have been identified, including number of drugs prescribed\(^6\), patient age\(^7\) and genetic variants\(^8\). Side effect-linked genetic variants identified so far are predominantly associated with drug pharmacokinetics, thereby affecting exposure of the body to a particular drug, but these variants do not give any indication of the mechanism by which pathogenesis is initiated. A recent study suggests that as many as half of drug side effects are related to known drug–protein-binding events\(^9\), and progress has been made towards systematically identifying drug-binding events\(^10\). However, only modest progress has been made towards elucidating specific drug-induced changes downstream of binding events for the majority of drugs (Fig. 1a)\(^11\). These downstream effects in many cases may be most directly tied to side effect pathogenesis as well as patient genetic and environmental background.

Recent literature suggests that altered gene expression induced by drug treatment may be one mechanism by which drugs induce systemic off-target effects\(^12\)-\(^15\). Unfortunately, the lack of clinical data has impeded the determination of causality of particular gene expression changes in side effect pathogenesis\(^16\). Recent studies have successfully utilized in vitro drug-treated gene expression profiles to predict clinical drug effectiveness\(^17,18\), suggesting that in vitro data may contain features that are clinically conserved. However, demonstrating the relevance of in vitro drug response features to clinical side effect pathogenesis presents a significant challenge, due largely to the lack of suitable validating data sets and difficulty of clinical experimentation.

To address this challenge, we develop a network-based data analysis workflow built upon the use of in vitro drug treatment data to identify candidate side effect-linked features and a large collection of historical clinical and disease model data as a source of validation (Fig. 1). First, we identify in vitro gene expression changes preferentially induced by drugs with clinically defined side effects to identify candidate side effect-linked expression features. Then, we cross-reference these side effect-linked features with independent legacy clinical data found in the literature to corroborate their relevance in terms of five causal relationships. We implement this strategy within the context of the reconstructed global human metabolic network\(^19,20\), which provides a biologically coherent structure for data integration due to the high degree of network annotation and clear functional connectivity between genes via metabolic pathways\(^20,21\).

**Results**

**Calculation of drug-induced metabolite perturbations.** We first identified drug-induced metabolic gene expression changes within 6,040 gene expression profiles in the Connectivity Map (CMap) data set, representing three human cell lines exposed to 1,221 drug compounds\(^22\) (Fig. 1a). We analysed the expression profiles using the reconstructed global human metabolic network Recon 1 (ref. 19) with a novel metabolic pathway analysis algorithm, termed MetChange (Metabolite-Centered Hotspots of Altered Network Gene Expression). MetChange is a constraint-based modelling\(^23\) algorithm that computes a score for each metabolite summarizing the drug-induced gene expression changes along calculated production pathways for the metabolite (Fig. 2). A MetChange score for a metabolite defines how expression has changed in a pathway containing these metabolite production reactions. Production in this case does not necessarily indicate secretion, as the majority of metabolites produced by one metabolic pathway are consumed in other metabolic pathways. We also note that gene expression is not the sole determinant of pathway activity, as gene and protein
Validation of calculated metabolic perturbations. To compare the MetChange method against existing approaches that predict a metabolic outcome based on gene expression data, a published gene expression data from carbon and nitrogen starvation in *S. cerevisiae* was analysed23,26. A previously generated metabolic reconstruction of *S. cerevisiae*, iMM904 (ref. 27), was used to compute MetChange scores for each condition. Scores were compared with metabolomics data generated for the same conditions using the relative changes from the initial time point. A total of 60 metabolites across five time points for each carbon and nitrogen starvation were compared for both absolute (that is, magnitude) correlation and directional (that is, increased or decreased) correlation with MetChange scores. The MetChange algorithm compared favourably both with other network-based expression analysis methods and with the use of gene expression alone in predicting metabolic perturbations (Fig. 3). Reassuringly, k-means clustering and principle component analysis of MetChange scores, mapped metabolomics data, and mapped expression data suggest that MetChange scores maintain functional relationships between time points that are present in expression and metabolomics data (Fig. 3).

To further validate the metabolic perturbations predicted by the MetChange method using the CMap data set, we performed a number of high-throughput computational analyses comparing MetChange perturbations with drug response properties (Table 1). First, metabolite scores were compared with co-occurrences of drug-metabolite text terms in the PubMed database. A bootstrap analysis of Z-score permutations showed that PubMed drug-metabolite associations are recalled in a statistically significant manner (non-parametric perturbation *P*-value *P* < 10<sup>−3</sup> for 1,000 perturbations of MetChange scores). Second, it was found that known metabolic drug targets as found in the DrugBank database28,29 are significantly closer in reaction proximity to known highly perturbed metabolites than less perturbed metabolites for these drugs (median Wilcoxon rank sum *P* < 1.65 × 10<sup>−10</sup> for the highest scoring bin). Third, MetChange scores were found to be able to predict drug–protein literature co-associations within the PubMed database in a statistically significant manner (non-parametric perturbation *P* < 0.01).

Finally, we validated perturbation predictions in targeted experimental measurements in MCF-7 cell culture under treatment by the drugs metformin, haloperidol and genistein. These drugs were chosen because of previous evidence suggesting significant metabolic perturbation by these drugs. Measured metabolites were chosen to have broad coverage of pathways and target highly perturbed pathways as predicted by MetChange. Drug concentrations were chosen based on previous *in vitro* studies using these drugs.

First, treatment with the antipsychotic haloperidol revealed decreased uptake of vitamin B6, consistent with the calculated decrease in the B6 processing pathway (Fig. 4a). Second, treatment with metformin, a 5′ AMP-activated protein kinase (AMPK) activator, showed significant perturbation of tricarboxylic acid cycle and fatty acid oxidation metabolites. The observed change was consistent with the large calculated perturbation but was in the opposite direction of the transcriptional change, indicating substantial non-transcriptional control of the metabolite levels (Fig. 4b). Additional calculated expression are imperfectly correlated and enzyme functional state may change due to perturbation as well. However, change in metabolic gene expression may still indicate a pathogenic metabolic functional change.

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Figure 2 | Description of the MetChange algorithm. (1) Using a metabolic network reconstruction, sink (demand) reactions are added for each metabolite. Demand reactions are irreversible with the stoichiometry: metabolite $\rightarrow$ $\varnothing$. Each demand reaction is maximized in turn to obtain maximal production values for each metabolite using a linear programming problem (LP Problem 1). (2) Reaction presence/absence *P*-values are generated from gene expression data and mapped onto the metabolic network. A second linear programming problem is then solved (LP Problem 2) for each metabolite. LP Problem 2 identifies the flux solution that PubMed drug-metabolite associations are recalled in a statistically significant manner (non-parametric perturbation *P*-value *P* < 10<sup>−3</sup> for 1,000 perturbations of MetChange scores). Second, it was found that known metabolic drug targets as found in the DrugBank database28,29 are significantly closer in reaction proximity to known highly perturbed metabolites than less perturbed metabolites for these drugs (median Wilcoxon rank sum *P* < 1.65 × 10<sup>−10</sup> for the highest scoring bin). Third, MetChange scores were found to be able to predict drug–protein literature co-associations within the PubMed database in a statistically significant manner (non-parametric perturbation *P* < 0.01).

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metformin-induced changes supported by previous results include: (i) a downregulation of folate metabolism consistent with reported folate deficiency in metformin-treated patients\(^{30}\), (ii) upregulated oxidative stress response consistent with reported lower reactive oxygen species levels\(^{31}\) and (iii) increased polyamine synthesis and recycling pathways that may result from shared use of organic cation transport (OCT) proteins between metformin\(^{32}\) and polyamines\(^{33}\). Third, treatment with genistein, an isoflavone with hypolipidemic effects\(^{34}\), experimentally showed a preferential reduction of the fraction of mono-unsaturated fatty acids synthesized from glucose that was consistent with predictions (Fig. 4c).

**Identification of in vitro disease-linked metabolic pathways.** Using MetChange scores generated from the CMap data set, we then identified the drug-induced metabolic gene expression changes that are most conserved among drugs with particular clinically described side effects. We used a machine learning approach to select discriminating metabolite production pathway perturbations from the set of MetChange-calculated pathway changes based upon side effect frequency data reported in the Side Effect Resource (SIDER) database\(^{35}\) (Figs 1b and 5). A total of 357 side effects across 1,417 gene expression profiles from CMap were analysed, based on the criterion that at least 30 expression sets per side effect were available. The cutoff in number of expressions sets was chosen rationally as a balance between the scale of the study and the robustness of the side effect signature obtained. Overall, 2,422 disease-linked drug-changed (termed DISLoDGED) metabolic pathways were identified with this analysis. These pathways are MetChange calculated metabolite production pathways that are over-represented in perturbations by drugs with particular side effects. DISLoDGED pathways are potential side effect to pathway relationships, hypothesizing that drug-induced perturbations away from metabolic homeostasis are involved in side effect pathogenesis. The calculated DISLoDGED pathways thus comprise a large set of omics-driven hypotheses of side effect pathogenic mechanisms (Supplementary Data 2). These associations were initially supported by automated co-association searches of PubMed abstracts to examine disease-linked nutrient deficiencies across all 357 side effects (Table 1). Downregulated DISLoDGED pathways were found to be marginally significantly predictive of deficiencies associated with corresponding specific diseases (permutation \(P = 0.055\)), whereas upregulated pathways were not predictive (permutation \(P = 0.39\)). We then sought to determine whether...
B6 levels were measured by an enzymatic assay in media supernatant and lysates of MCF-7 cells, showing decreased utilization of the essential nutrient. (In the left panel, metformin gene expression perturbations are shown on the left, and resulting MetChange scores for related metabolites are shown on the right. The right panel shows response of tricarboxylic acid metabolites and acylcarnitines to metformin treatment. Metabolites show a large perturbation in the opposite direction of the observed transcriptional changes, validating the presence of a perturbation but indicating non-transcriptional control of metabolite levels in central metabolism as well. (In the upper panel, genistein is predicted to preferentially decrease production of mono-unsaturated fatty acids. The lower panel shows measurement of biosynthesis fraction of different fatty acid groups and precursors from glucose following genistein treatment measured by NMR spectroscopy, which validate modelling predictions. MUFA: Mono-unsaturated fatty acids. SFA: Saturated fatty acids. PG: Phosphoglycerate. Error bars represent standard deviations in all cases. Each experiment was performed in biological triplicate.

Comparison of DISLoDGED pathways with side effect pathogenesis. We first examined whether DISLoDGED metabolic pathways contain genes with variants that alter clinical side effect susceptibility, according to the data in MDDB (Table 1). Causal gene variants are typically considered to affect either drug pharmacokinetics, which involves drug exposure, or drug pharmacodynamics, which involves the interaction of the drug with the body. The majority of identified gene variants affecting drug side effect incidence are involved in drug pharmacokinetics, because these genes historically have been simpler to identify as drug metabolism genes are largely known. However, we focused upon genes affecting pharmacodynamics, as these genes are more directly indicative of the mechanisms underlying pathogenesis. In MDDB, we identified nine side effect susceptibility gene variants that overlap directly with metabolism but are not involved in drug pharmacokinetics (Supplementary Note 1). We found that for each of the nine side effect susceptibility genes, at least one overlapping DISLoDGED pathway for this side effect had been identified through our analysis (Table 2). This overlap between DISLoDGED pathways and non-pharmacokinetic side effect susceptibility altering genes was highly significant (joint hyper-geometric tests, \( P \)-value \( 2.2 \times 10^{-11} \), see Supplementary Note 2 for calculation).

To support the relevance of this overlap, we assessed the nine overlapping DISLoDGED pathways in the context of additional known factors related to side effect pathogenesis (Table 2). We found that, in each case, the overlapping DISLoDGED pathway had established ties to the clinical pathogenesis of the side effect
Figure 5 | Description of the genetic algorithm used to identify metabolic signatures of side effects (DISLoDGED pathways). (1) Inputs to the algorithm are a set of response variables for each gene expression set (either MetChange scores or gene expression changes), a binary presence/absence vector for whether each sample was treated with a drug that has the side effect, and the desired maximum number of predictor variables desired. (2) At initiation, the genetic algorithm generates a ‘population’ of random guesses at the predictor variables, termed ‘individuals’, and assigns them either a value of 0, 1, or 1. For each individual, all gene expression samples are scored as the response variables (MetChange or gene expression changes) multiplied by the candidate signature. (3) Each gene expression sample is then ranked and a receiver operator characteristic (ROC) curve is generated and area under the curve (AUC) is calculated using the input presence/absence vector for the side effect or indication. The sample AUCs are the maximization objective of the genetic algorithm. (4) The genetic algorithm sub-routines are then used to generate a new population, biasing towards higher AUCs. Best solutions are maintained without modification, and lower scoring individuals are combined (‘crossed over’) and modified (‘mutated’) to search the solution space in a heuristic manner. The termination criteria are typically a number of generations without improvement; however, we applied a simple maximum time limit. (5) The signature yielding the highest prediction AUC is considered the best predictor set. In the example case, the resultant AUC is 1.0, a perfect predictor for the sample set. (6) To assess over-fitting and hence the predictive potential of the metabolic signature, tenfold cross-validation is performed by generating ten partitions of 90% of the data to train signatures and predict the remaining ten partitions of 10% of the data. To find signatures that have constant predictive power, the cross-validation signatures were summed, and high scoring metabolites were considered the conserved metabolic response signature (DISLoDGED pathway) for the side effect or indication. FPR, false positive rate; TPR, true positive rate; TP, true positive; TN, true negative; FP, false positive; FN, false negative.
Table 1 | Summary of large-scale validation of drug-metabolic perturbation associations predicted by MetChange calculations.

| Property                          | Database     | # Drugs | Properties analysed | Statistical significance     |
|----------------------------------|--------------|---------|---------------------|------------------------------|
| Drug target distance             | DrugBank     | 134     | 536 Target reactions | P = 1.65 × 10⁻¹⁰ (Wilcoxon rank sum test) |
| Drug-protein association         | PubMed       | 645     | 501 Signalling proteins | P < 0.01 (Permutation test) |
| Drug-metabolite association      | PubMed       | 645     | 375 Metabolites      | P < 0.001 (Permutation test) |
| Drug side effect                 | SIDER, PubMed | 334     | 357 Side effects total | P = 5.6 × 10⁻² (Permutation test) |

For co-term analysis

Abbreviation: SIDER, Side Effect Resource.

Table 2 | Existing studies of pharmacodynamic-altering side effect susceptibility genes and overlapping DISLoDGED pathways*.

| 1 Side effect | 2 DISLoDGED Pathway (# metabolites in pathway) | 3 PD gene | 4 Nutrient overlap | 5 Pathophysiology | 6 Metabolic perturbation | 7 In vivo drug perturbation | 8 Supplement |
|--------------|----------------------------------------------|-----------|-------------------|-------------------|-------------------------|--------------------------|--------------|
| Weight gain  | Melatonin                                     | MC4R      | MC4R levels correlated with melatonin | Melatonin and MC4R energy regulators | Decreased melatonin tied to metabolic syndrome | Antipsychotics suppress melatonin synthesis | Melatonin |
| Parkinsonism | Poly-unsaturated fatty acids                  | ZNF202    | ZNF202 lipid regulator | PUFAs linked to Lewy bodies | Altered lipid oxidation | NA | RBC PUFAs altered |
| Tardive      | Serotonin                                     | 5-HT      | Direct ligand Dopaminergic neuron interaction | Dopamine primary mediator | NA | 5-Hydroxyindoleacetate perturbation | Essential fatty acids |
| Dyskinesia   | Ascorbate                                     | D3 receptors | Dopamine synthesis cofactor | NA | Metabolite levels perturbed | Vitamin C, vitamin E |
| Myotoxicity  | CoQ10                                        | COQ2      | CoQ10 synthesis | Mitochondrial function | Q10 deficiency associated w/ myopathy | Decreased under statin treatment | CoQ10 (conflicting) |
| Cardiac      | Urea cycle                                    | NOS1AP    | Nitrogen regulation | NO affects cardiac function | Altered NO linked to arrhythmia | Various affects NO production | L-arginine |
| arrhythmia   | Pentose phosphate pathway                     | GPDL1, ZFH3 | Oxidative stress response | Oxidative stress affects cardiac function | H₂O₂ induces arrhythmia | Clomipramine induces via oxidative stress | Various antioxidants |
| Hearing loss | Ascorbate                                     | COMT      | Dopamine synthesis cofactor | Dopamine innervation in inner ear | Dopa synth. inhibition tied to hearing loss | Cisplatin inhibits dopamine | Vitamin C, dopamine |
| Hearing loss | Ascorbate, lipoate                            | Glutathione S-transferases | Oxidative stress response | ROS-induced cochlear cell death | SOD deficiency tied to hearing loss | Cisplatin induces via oxidative stress | Vitamin C, lipoate |

Abbreviations: NA, not applicable; PUFA, poly-unsaturated fatty acids; RBC, red blood cell; ROS, reactive oxygen species; SOD, superoxide dismutase; 5-HT, serotonin.

*Existing studies of pharmacodynamic-altering side effect susceptibility genes and overlapping DISLoDGED pathways. The two cases of tardive dyskinesia haveNA in the ‘Metabolic perturbation in disease’ row as tardive dyskinesia is not typically described as a disease outside of its occurrence as an adverse drug response, therefore there is no drug-independent case for comparison. See the Supplementary Note 1 for full discussion and references related to each case.

In the seven cases where the side effect pathology had been reported independently of the drug, alterations in the DISLoDGED pathway have been associated with the disease (Table 2, column 6). Critically, in each of the nine cases, we found that drugs causing the side effect had been reported to induce a perturbation in the DISLoDGED pathway. In vivo, demonstrating that the in vitro-derived DISLoDGED pathways are similarly perturbed in vivo (Table 2, column 7). Furthermore, in seven of the nine cases, nutrient supplementation targeted at the DISLoDGED pathways have been shown to be effective in treating the drug side effect, whereas the remaining two cases are inconclusive due to reports of both positive and negative results (Table 2, column 8).

For example, among the 38 drugs within the CMap database reported in SIDER as causing increased risk of cardiac arrhythmias, we identified five DISLoDGED pathways (Supplementary Note 1). Three of these DISLoDGED pathways, which were a downregulation of oxidative pentose phosphate pathway and two related upregulations in nitrogen metabolism, overlap with genetic polymorphisms known to cause increased susceptibility to arrhythmias (Table 2). These pathways are known to have physiological ties to the pathogenesis of arrhythmias and have been shown to be perturbed in vivo by drugs causing arrhythmias (Table 2). Furthermore, nutrient supplementation targeted at these pathways has been found therapeutic in drug-induced arrhythmias (Table 2). The remaining examples are presented in Supplementary Note 1.

Comparison of DISLoDGED pathways with general disease pathways. We further expanded the scope of validation beyond the nine available cases of non-pharmacokinetic genetic variants directly affecting side effect incidence. The MDDB database was used to determine whether the identified DISLoDGED pathways...
are conserved within in vivo data related to non-drug-induced pathologies as well, where a significantly larger body of literature exists than for side effect pathogenesis. This analysis hypothesizes that non-drug-induced pathogenesis and corresponding side effect pathogenesis share a common basis. Using the data collected, we assessed the calculated DISLoDGED pathways in terms of the five causal relationships in MDDB. Downregulated and upregulated DISLoDGED pathways were assessed independently to examine directional causal relationships. Approximately one-sixth of calculated DISLoDGED pathways were investigated for validation, and compared with an equal number of randomized predictions as a control. The next sections describe each causal link examined.

**Gene variants affecting disease incidence.** We first sought to determine whether DISLoDGED pathways contain known disease-linked gene variants. To do this, we compared the metabolic subsystems into which disease-linked gene variants and DISLoDGED pathways occur. We analysed 239 metabolic disease-linked genes from MDDB and found an enrichment of disease-linked genetic variants among transporters (one-tailed hypergeometric \( P = 0.0048 \)) and inositol metabolism (one-tailed hypergeometric \( P = 0.002 \)), as well as a depletion of variants in central carbon metabolism (one-tailed hypergeometric \( P = 0.035 \)). We found that DISLoDGED pathways showed similar results, including an enrichment of downregulated DISLoDGED pathways in inositol metabolism \( (P = 0.018) \) mirroring the enrichment found in disease-linked genetic variants, as well as enrichment of DISLoDGED pathways in non-central metabolism (Supplementary Table 1). These results indicate certain metabolic pathways may be inherently less robust to pathological perturbation.

**DISLoDGED pathway associations with disease physiology.** Next, we examined whether DISLoDGED pathways are known to be essential to system physiology in a manner that could result in disease when perturbed. To compare DISLoDGED pathways to disease pathophysiology, side effects were grouped based on affected physiological systems, such as renal diseases or autoimmune complications. We first grouped DISLoDGED pathways by nearest nutrient for better coverage in the literature (Fig. 6a). DISLoDGED nutrient pathways preferentially altered by drugs causing side effects in specific physiological systems were then calculated. In 17 of the 18 cases of enrichments of downregulated DISLoDGED nutrient pathways within particular physiological systems, the downregulated pathways had directionally consistent links to the disease pathophysiology (Fig. 6b and Supplementary Data 2). For example, inositol metabolism downregulation was enriched among drugs with side effects affecting the kidney, including kidney failure. Supporting this relationship, the kidney is a primary site of inositol synthesis, and inhibition of inositol transport has been reported to cause renal failure.

Similarly, pathways that were upregulated by drugs affecting particular physiological systems also showed directionally consistent links to pathophysiology. In 9 out of 17 DISLoDGED nutrient pathways that were upregulated in particular physiological systems, inhibitors targeted at the upregulated pathway were established therapeutics within the disease class (Fig. 6c). For example, drugs causing cancer- or autoimmune-related side effects were enriched in upregulation of folate metabolism, and anti-folates are commonly used in treatment of diseases in both classes. Supporting the implications of this upregulation in side effect pathogenesis, studies have shown that folate supplementation is tied to increased incidence of both cancer and childhood asthma.

**Altered activity of DISLoDGED pathways in disease.** We then sought to determine whether DISLoDGED pathways are significantly over- or under-active in corresponding clinical disease and disease models, based on several metrics in the MDDB. To perform this analysis, DISLoDGED metabolic pathways were associated with dietary nutrients nearest in the metabolic network. In an initial analysis of 323 DISLoDGED nutrient pathways, downregulated DISLoDGED pathways were found to be significantly enriched in disease-associated nutrient deficiencies (59% enrichment, binomial \( P = 0.0017 \)), whereas upregulated DISLoDGED pathways were depleted in disease-associated nutrient deficiencies (47% depletion, binomial \( P = 0.036 \); Fig. 7a).

These results were confirmed in an independent set of 453 investigated DISLoDGED pathways added to MDDB. Downregulated DISLoDGED pathways were more likely to have a causal downregulation associated with the corresponding pathology (17% enrichment of downregulation, binomial one-tailed \( P = 0.045 \); Fig. 7b), whereas upregulated DISLoDGED pathways were significantly predictive of consistently over-active pathways tied to corresponding pathologies (67% enrichment of over-activity, binomial one-tailed \( P = 0.003 \); Fig. 7c).

**Effect of targeted inhibition of DISLoDGED metabolic pathways.** Next, we analysed in vivo data on the effect of non-drug chemical inhibitors targeted at calculated DISLoDGED pathways. A causal relationship would be indicated by (i) inhibitors targeted at downregulated DISLoDGED pathways reproducing the clinical side effect and (ii) inhibitors targeted at upregulated DISLoDGED pathways treating the clinical disease. We found that metabolic inhibitors targeting downregulated DISLoDGED pathways specifically were significantly more likely to cause corresponding side effects (28% enrichment, binomial one-tailed \( P = 0.04 \); Fig. 7d). Also, we found that both downregulated and upregulated DISLoDGED pathways were more likely to be known targets for effective metabolic inhibition to treat corresponding diseases, indicative of imperfect directionality of predictions (64% enrichment, binomial one-tailed \( P = 0.0015 \) and 29% enrichment, binomial one-tailed \( P = 0.09 \); respectively; Fig. 7e).

**Effect of supplementation targeted at DISLoDGED pathways.** Finally, we compared calculated DISLoDGED pathways grouped by nearest nutrient with clinical therapeutic nutrient supplementation data (Fig. 6a). We sought to determine whether downregulated DISLoDGED pathways might be targets for nutrient supplementation as a disease therapy. We observed that predicted downregulated DISLoDGED pathways were preferentially targets of nutrient supplementation to alleviate corresponding pathologies (24% enrichment, binomial one-tailed \( P = 0.065 \); Fig. 7e), whereas overexpressed pathways were preferentially depleted as effective nutrient supplements (19% depletion, binomial \( P = 0.14 \)), although the relationships did not reach statistical significance for given sample sizes available.

**Discussion** The systems biology-based workflow developed in this study predicts side effect-linked dysregulated metabolic pathways (termed DISLoDGED pathways) from drug treatment of human cells in culture. We contextualize numerous independent data types that together support a key role of drug-induced non-pharmacokinetic metabolic dysregulation in side effect pathogenesis. Through the construction of a comprehensive resource on metabolic involvement in disease, we corroborate the predictions made using five independent clinical, genetic and biochemical
The class are coloured according to the legend. Many of the enriched interactions are consistent with known effects of nutrient/pathway perturbation on the number of side effects with a downregulation in the production pathway for a metabolite. (Figure, the number of side effects with an upregulation in the production pathway for a metabolite is shown in yellow boxes, whereas blue boxes show inhibition of metabolic pathway beneficial.)

The work presented relied upon the development of a systems biology workflow to identify side effect-linked features and causal protein-binding events and genetic susceptibility factors.

The present work we believe is the first to validate omics-driven predictions of post-binding mechanisms underlying side pathogenesis using clinical data at a large scale, which was empowered by the generation of a new database through manual curation of the literature.

The workflow presented is highly dependent upon large amounts of gene expression data obtained under treatment by diverse drugs to filter out perturbations due to factors other than common side effects. Normalization of data across multiple studies and platforms is a substantial issue. Thus, data generated within a single study is ideal, but such large studies are rare. For this reason, the potential to extend the workflow to new data types, for example, in vivo gene expression data, may be limited. Owing to the use of in vitro data, corroboration against in vivo evidence is limited.

Figure 6 | Maps of interactions between DISLoDGED pathways grouped by nutrient domain and corresponding side effects grouped by disease class. Only nutrients and disease classes with at least one marginally enriched nutrient-class interaction (hypergeometric P<0.1) are shown. (a) Using the metabolic network reconstruction Recon 1, side effect-specific metabolic perturbations (DISLoDGED pathways) are grouped into nutrient domains to enable comparison with existing disease-related genetic, clinical and pre-clinical data to assess the potential causality of observed perturbations. In this figure, the number of side effects with an upregulation in the production pathway for a metabolite is shown in yellow boxes, whereas blue boxes show the number of side effects with a downregulation in the production pathway for a metabolite. (b) Downregulated DISLoDGED pathways. Nutrient-disease class interactions indicating an enrichment of downregulations in drugs causing side effects within the class are coloured according to the legend. (c) Upregulated DISLoDGED pathways. Nutrient-disease class interactions indicating an enrichment of upregulations in drugs causing side effects within the class are coloured according to the legend. Many of the enriched interactions are consistent with known effects of nutrient/pathway perturbation on the corresponding disease classes and physiological systems.

bodies of the literature. These results provide understanding of the mechanisms underlying side effect pathogenesis, which have remained largely opaque despite recent progress identifying causal protein-binding events and genetic susceptibility factors.

The work presented relied upon the development of a systems biology workflow to identify side effect-linked features and validate the relevance of these features using historical in vivo data. The use of systems biology approaches to study drug side effects has become an active field in recent years. Previous studies have predicted drug–protein-binding events responsible for side effects, effective combinations of drugs to minimize side effects, and drug mechanisms of action underlying side effect pathogenesis. Still, uncovering how causal binding events result in disease is an essential and largely unanswered question, as identified pathogenic mechanisms can potentially be used to design therapies to circumvent drug side effects.
data appears critical. Further deployment of the presented workflow may hinge upon expansion of curated disease data necessary to corroborate the clinical relevance of *in vitro* drug treatment features. Furthermore, the definition of pathways used to integrate disparate data types has yet to be concretely established and may be an area for further workflow optimization.

Owing to the ubiquity of gene set enrichment analysis (GSEA)\(^5\) in pathway-based analysis of gene expression, a discussion on the differences between the MetChange algorithm and GSEA is warranted. Both methods attempt to aggregate ‘signal’ in gene expression data along pathway definitions to increase the interpretability of the data and decrease the effect of noise. GSEA uses a variety of pathways, including manually curated metabolic pathways, and results are typically *P*-values of a Kolmogorov–Smirnov test for the likelihood that the cumulative distribution of expressions of genes in each pathway have not changed between conditions. MetChange defines a different production pathway for each metabolite in the network, based on calculation of functional states of the metabolic network, and scores for each metabolite how gene expression along this production pathway has changed between conditions. As a result, MetChange has the potential to give finer resolution results, as its results are defined at the level of individual metabolite scores rather than pathway scores. However, the overall performance of the methods is difficult compared due to the fundamental difference in resolution of outputs.

In the comparison between MetChange and comparable methods (Fig. 3), it is apparent that, on the data set used, output correlations of none of the methods with measured metabolite perturbations in yeast were particularly high. However, given that this analysis is possibly using these methods out of their intended use cases, no presumptions should be made regarding the general usefulness of any of these methods. Rather, this may highlight the difficulty in establishing standards by which to compare pathway analysis algorithms with statistical rigor. In this study, we performed the analysis primarily to provide some context for the relative performance of the MetChange algorithm at one specific task relevant to predicting metabolic changes occurring within the cell. We note that a recent method to predict metabolism changes from gene expression has reported statistically significant correlations on three other yeast data sets\(^5\), including values that exceed those reported in Fig. 3.

There are at least two obvious potential improvements that could be made to the workflow used in this work. First, as the cell lines within the CMap are within the NCI60 cell line panel, and expression, growth and most recently exometabolomic\(^12\) data have been measured for these lines, cell-specific metabolic models could be used in place of a global model. Second, the human metabolic network reconstruction has been updated with the publication of Recon 2 (ref. 55), and thus improvements might be made from the increased scope of the model. We evaluated the potential for improvements using these changes by constructing cell-specific models for the MCF-7, HL-60 and PC-3 cell lines based on Recon 2 and running the MetChange algorithm on randomized simulated expression data drawn from an empirical distribution of MAS5.0 normalized *P*-values from the CMap data used in this study. Briefly, between cell-specific Recon 2 models, error was between 10 and 25% across replicates when given the same metabolite uptake constraints. The largest difference observed was by enforcing measured metabolite uptake constraints. Differences in MetChange scores between models constrained and unconstrained by metabolite uptakes were around 70%, indicating that significantly constraining the flux state of the model alters MetChange scores more so than topological differences. Thus, further improvements in prediction accuracy might be seen by accounting for the baseline metabolic differences between cell lines.

One interesting outcome of the work was that, although previously identified genetic susceptibility factors in the literature are dominated by genes involved in pharmacokinetics, we observe overlap of drug-induced metabolic changes with nine genes that affect pharmacodynamics. This may suggest that such non-pharmacokinetic genes may play a larger role in side effect pathogenesis than currently appreciated. We did, however, see some alteration of pharmacokinetic genes in a few cases as well (Supplementary Note 3). In addition, DISLoDGED pathways overlapped with clinical disease-linked pathways for both drug-treated and drug-independent studies, suggesting a common basis in pathogenesis. Furthermore, our results suggest that targeted nutrient supplementation may be a relatively simple and inexpensive path to broadly reduce side effect incidence. The impact of drugs on patient metabolic status is thought to be an underappreciated but important aspect of drug response\(^5\), and this work further suggests this interaction is worthy of significant investigation. Patient attrition may be reduced through effective

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**Figure 7 | Multiple lines of clinical evidence in MDDB implicate a causal role for drug-induced metabolic transcription changes in side effect pathogenesis.** (a) Downregulated DISLoDGED pathways were found to preferentially alleviate the corresponding pathology. (c) Upregulated DISLoDGED pathways are found to be enriched in consistently over-active pathways associated with corresponding pathologies. (d) Inhibitors targeted at DISLoDGED metabolic pathways are more likely to be effective treatments in corresponding pathologies. (e) Metabolic inhibitors targeted at downregulated DISLoDGED pathways are significantly more likely to cause corresponding side effects. (f) Nutrient supplementation targeted at downregulated DISLoDGED pathways was found to preferentially alleviate the corresponding pathology. *P*-values are defined for one-tailed binomial tests based on the level of the treated sample relative to the control (random). Sample sizes were chosen *a priori* based on feasibility of data collection, without an expectation for a particular effect size. Symbols: * indicates significant or $z < 0.05$, + indicates marginally significant or $z < 0.1$. 

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nutrient supplement to drug pairing during the development process. A natural question that may arise is whether certain pathways or side effect disease classes are more successfully predicted by the method used in this work than others. A sensitivity analysis shows that the method appears to be fairly robust in being able to predict DISLoDGED pathways in various areas of the metabolic network and across disease classes. Restricting to at least five pairs investigated for both the down-regulated DISLoDGED pathways and random pairs, deficiencies of nine nutrients (polyunsaturated fatty acids, coenzyme Q10, melatonin, niacin, riboflavin, steroids, thiamine, vitamin A and vitamin D) had disease associations better predicted by down-regulated DISLoDGED pathways compared with random, whereas one (choline) did not. Similarly, in side effects with at least two pairs investigated in both random and real, seven side effect/nutrient deficiency relationships were better predicted by down-regulated DISLoDGED pathways (dyspepsia, epilepsy, hyperlipidemia, interstitial nephritis, tardive dyskinesia, testicular atrophy, and thrombocytopenia), whereas only two were better predicted by random nutrients (eczema and tendosynovitis). These types of sensitivity analyses suggest that predictive capability for particular nutrients and side effects are vulnerable to error because of small sample size but corroborate overall results that dysregulated DISLoDGED pathways are predictive of metabolic pathways associated with corresponding pathologies.

Although challenges remain, the ability to observe perturbations important to in vivo side effect susceptibility within in vitro data suggests that early-stage drug screening to identify and manage side effect risk may become possible, analogous to other ‘disease in a dish’ efforts5,6. Notably, high use drugs such as statins and the most commonly used cancer drugs (e.g., taxanes) were predicted to have side effects with at least two pairs investigated in both random and real, seven side effect/nutrient deficiency relationships were better predicted by down-regulated DISLoDGED pathways (dyspepsia, epilepsy, hyperlipidemia, interstitial nephritis, tardive dyskinesia, testicular atrophy, and thrombocytopenia), whereas only two were better predicted by random nutrients (eczema and tendosynovitis). These types of sensitivity analyses suggest that predictive capability for particular nutrients and side effects are vulnerable to error because of small sample size but corroborate overall results that dysregulated DISLoDGED pathways are predictive of metabolic pathways associated with corresponding pathologies. Thus, although we do not discount that other methods may exhibit better performance by certain standards, we chose our approach to meet the specific requirements of our problem, as typical in the field. An overview of the methods is below and a MATLAB implementation is provided (see Supplementary Software).

**CMap data processing and integration.** Gene expression data for the AffyMetrix HT Human Genome U133A platform were obtained from the CMap database22 for MCF-7, PC-3 and HL-60 cell lines. Data were MAS5.0-normalized with the BioConductor package54. Human Entrez Gene identifiers associated with probes were used to map detection P-values to their corresponding reactions based on the Boolean gene–protein–reaction associations as was previously described59. Reactions that were not associated with a gene were assigned a P-value of 0.

**Metabolite exchange was set to an exchange value of −1 for DMEM (MCF-7) or −0.1 for PC-3 and HL-60 media metabolites. In cases where metabolite production was possible with open constraints, the metabolite was removed from further analysis. For the cases of in vitro experimental validation under genistein treatment, for which data was generated for the fraction of metabolites generated from glucose, the MetChange algorithm was run using glucose as the sole carbon source to enable direct comparison with the data.**

**The MetChange algorithm.** The Metabolite-Centered Hotspots of Altered Network Gene Expression (MetChange) algorithm was used to analyse differential quantitative gene expression profiles in the context of a genome-scale metabolic network57. This algorithm is built upon the Gene Inactivity Moderated by Metabolism and Expression (GIMME) algorithm previously developed to build context-specific metabolic networks based on gene expression data58,59.

The MetChange algorithm defines a consistency metric of an expression profile with optimally producing each metabolite in the metabolic network. For each metabolite, a sink reaction is created and flux through the sink reaction is maximized using flux balance analysis:

$$v_{max} = \max (v_x \cdot v - S \cdot v = 0, \ a \leq v \leq b)$$  \hspace{1cm} (1)

where $S$ is the stoichiometric matrix, $v$ is the reaction flux vector, $x$ and $b$ are vectors for the lower and upper bounds of the reactions and $e_x$ corresponds to the imposed reaction objective for each $i$th sink reaction. This results in a set of optimal production fluxes for each network metabolite.

Each reaction is then weighted by a detection P-value from mapped expression data to solve a second optimization problem. To obtain metabolite production ‘consistency’ scores, $x_{r}$ for each $r$th metabolite, scores were generated by setting the lower bound of the metabolite sink reaction to its maximal production flux, $v_{max}$, and minimizing the inner product of the reaction detection P-values:

$$x_i = \min (v_x \cdot v - S \cdot v = 0, \ a \leq v \leq b, x_i = v_{max})$$  \hspace{1cm} (2)

where mapped detection P-values $p$ were stored in a weighting vector $e_x$ and each vector component $e_{x_i} = p_i$ mapped to each $i$th reaction. This optimization is the GIMME algorithm-like step of the MetChange algorithm. Each $r$th component in $e_x$ serve as a cue for whether a reaction is detected or absent (that is, a lower P-value indicates a reaction is more likely to be present). Hence, this second optimization determines: (i) a flux distribution which maximizes the production of a metabolite weighted (ii) generates a weighted flux distribution score defining consistency of the production of that metabolite with expression data. These scores cannot be directly compared across metabolites because of the fact that each metabolite has a different network flux state at maximal production. Thus, we compared metabolite production consistency scores between treated and control samples with a standardized score:

$$Z_{max} = \frac{X_{max} - X_{min}}{X_{97.5} - X_{2.5}}$$  \hspace{1cm} (3)

A MATLAB implementation of the MetChange algorithm is provided in the Supplementary Software File.

**Analysis of expression data for yeast under carbon and nitrogen starvation.** Metabolite concentration25 and gene expression26 data for S. cerevisiae were mapped to a genome-scale yeast metabolic reconstruction, iMM904 (ref. 26). For all methods, scores were compared to log-2 metabolite concentration changes relative to the initial time point for carbon or nitrogen data. Spearman correlations were calculated to avoid biases due to variable score distributions among the different methods. K-means clustering analysis was performed using 100 replicates. In all cases, the same clusters were obtained in repeated runs. For the MetChange algorithm, reported expression values were used to generate reaction presence/absence P-values. Expression values were ranked, and based upon the apparent distribution, a value corresponding to the bottom second percentile of expression was used to define the noise level. Expression values below this threshold were assigned a P-value of 1 as ‘not present.’ The P-values below the threshold were then inverted across the threshold to generate a symmetric distribution. The mean and standard deviation of this approximated noise threshold were then inverted across the threshold to generate a symmetric distribution. The mean and standard deviation of this approximated noise level. Expression values below this threshold were assigned a P-value of 1 as ‘not present.’ The P-values below the threshold were then inverted across the threshold to generate a symmetric distribution.

**Methods**

**Overview of computational approach.** As described schematically in Fig. 1, with more detailed methods diagrams in Figs 2 and 5, we employed a combination of constraint-based modelling and machine learning to look for metabolic gene expression perturbations that are conserved in adverse drug reactions. The first step is the analysis of drug-specific metabolic perturbations using the constraint-based MetChange algorithm described below and comparison of these perturbations to drug-specific response properties. We then combined MetChange scores based on side effects and used an area under the Receiver Operating Characteristic curve (AUC)-maximizing classification genetic algorithm described below to determine small subsets of metabolic changes highly conserved in certain side effects. We note in general that the fields of constraint-based modelling and machine learning have developed a wide variety of methods that perform similar tasks. We compare our method with several other constraint-based methods and with gene expression data alone, as described in the Results. We also qualitatively contrast our method with GSEA in the Discussion. In general, performance of machine learning methods largely depends on the particular problem. We note that in cases of rare events, such that only a small fraction of samples have a property, AUC (or rank) maximizing algorithms have been shown to perform particularly well. In addition, we choose to place a hard constraint on the number of variables rather than using a traditional SVM with an L2-norm approach, for example. This was done to maximize interpretability of the resulting signatures, which was critical for later comparison of DISLoDGED pathways with pathology deficiency relationships and disease-linked and side effect-linked genetic susceptibilities.

**Supplementary Software.** Thus, although we do not discount that other methods may exhibit better performance by certain standards, we chose our approach to meet the specific requirements of our problem, as typical in the field. An overview of the methods is below and a MATLAB implementation is provided (see Supplementary Software).
the same reaction, the minimum value was used. The MetChange algorithm was then applied using these mapped reaction presence/absence \( P \)-values. As the data were longitudinal in time and multiple controls were not available, log-2 scores were calculated with respect to the initial time point, rather than taking standard scores.

Reporter metabolite analysis was then implemented. \( P \)-values for the significance in expression change were used as inputs to the reporter metabolite analysis. Each expression value was first standardized across conditions and then \( P \)-values were calculated using a standard \( Z \)-test for each gene across condition. 10,000 permutations of the data were used to generate the background levels of perturbation.

In addition, the E-Flux method was implemented, with metabolite sinks set as separate objective reactions similar to the MetChange algorithm for comparison. Optimal flux through each metabolite sink reaction was then calculated using flux balance analysis for each gene expression data set, and log-2 scores compared with the initial time point for each carbon and nitrogen data set were calculated.

For a direct comparison with gene expression levels cues, we implemented the following approach. First, standardized scores of expression level changes were calculated with respect to all gene expression data sets for the particular perturbation (carbon or nitrogen starvation). The scores were mapped to their respective reactions according to the gene–reaction association in the metabolic network. We then assessed whether absolute changes in expression levels are predictive of the magnitude of metabolite change by adding the standardized reaction scores, weighted by the absolute stoichiometric coefficients. We then assessed whether higher gene expression for a reaction was indicative of a higher reaction scores, weighted by the absolute stoichiometric coefficients for each metabolite.

Analysis of metabolic response phenotypes across drugs. The Recon 1 metabolic network was first converted into an irreversible network, such that each reaction proceeds only forward, and reactions that can proceed in multiple directions are split into two forward-proceeding reactions. The MetChange algorithm was run using gene expression presence/absence MASS.0 \( P \)-values from the Cmap database build 02 (ref. 22) when multiple controls were present, a standard score was generated. When a treated sample was from a batch with a single control, the mean and standard deviation of all control samples was used instead.

Cell line standard scores were then generated in the following manner. First, for each cell line, the median scores of all samples for each drug were found and used as the cell line-specific score. Then, to simplify the comparison and model the general metabolite response, cytosolic metabolite scores were taken when available. If no cytosolic metabolite existed, the median of scores across all compartments was taken as the metabolite score. Finally, a standard score across all drugs was calculated for each cell line. Consensus drug perturbations across cell lines were calculated by averaging cell-specific MetChange scores and standardizing across all drugs.

PubMed querying of drug associations. To identify drug-metabolite associations in the literature in an automated manner, PubMed/MEDLINE records were downloaded from the National Library of Medicine and abstracts were parsed to raw text. Chemical entities were tagged using Reflect61. A training set of 44 abstracts describing true drug-metabolite correlations (positive set) and 150 abstracts with other chemical entities (negative set) were used to train a Bayesian network to recognize biological association and causative relationship between an administered drug and the presence of a certain metabolite. A second Bayesian network was trained to recognize sentences within the abstracts that refer to metabolites from those that refer to other chemical entities. For each drug-metabolite co-occurrence the two Bayesian networks were used to assign a posterior for abstract occurrence and a posterior for sentence occurrence and the joint probability of the two posterior was used as the drug-metabolite score. The evaluation of the output is provided in Supplementary Data 1. The text-mining output was evaluated using a random sample of 100 pairs. The abstracts that describe each sampled pair were manually checked and identified as true or false positives. The score cutoff was set to 0.3, which provides an 83% true positive rate and 90% accuracy.

Text querying on side effect–metabolite, protein–metabolite and drug–protein literature co-associations was carried out using the Entrez Programming Utilities (NCBI) using a simple ‘AND’ specification. Side effects were obtained from the SIDER database35. Signalling protein lists were obtained from various publically hosted protein literature associations for the drug corresponding to the sample, and the protein–metabolite and drug–protein literature co-associations. These proteins were used as guesses for true drug–protein literature associations for the drug corresponding to the sample, and the protein–metabolite and drug–protein literature co-associations was then determined using the protein–metabolite literature co-associations. These frequencies were then mapped to all expression samples from CMap corresponding to the appropriate drugs. A minimum of 30 expression sets for a side effect were required for inclusion in the analysis. A total of 357 side effects were analysed for 850 expression sets corresponding to 334 drugs.

A genetic algorithm was then implemented, termed SiderFinder (Fig. 5). The matrix of MetChange scores for the 850 expression sets was input as well as the corresponding side effect frequencies for a particular side effect. A maximum number of predictor metabolites was set to 20 metabolites. A set of 125 candidate sidedness scores were generated, assigning values of –1, 0 or 1 to each metabolite to indicate negative, no or positive prediction of a high MetChange metabolite score on occurrence of side effect. Each expression set was then scored for the side effect for each individual by multiplying the predictor set for the individual by the SideFnder scores for the expression set. These side effect scores were ranked and a pseudo-ROC curve was generated by comparison of scores with the side effect frequencies for the current side effect. At each threshold, expression sets with a side effect score greater than the threshold were called as having the side effect. To weigh more heavily affecting samples with higher side effect frequency, the base 10 logarithm of each side effect frequency was taken and adjusted such that the lowest frequency had a value of 1, each order of magnitude greater in frequency is a unit greater, and all zero frequency side effects remain zero. An ROC curve was then calculated with true positive hits being assigned the value of the adjusted side effect frequency, with no effect to false positive, true negative and false negative values. The number of true values was then taken to be the sum of the adjusted frequencies, and the ROC area under the ROC curve (AUC) is calculated. The AUC of this curve was then used as the objective function to maximize in the genetic algorithm. Tenfold cross-validation was performed, and perturbations that appeared cumulatively in at least four of the ten sets we selected. Using the genetic algorithm directly on gene expression data achieved similar classification performance (results not shown), but metabolite scores were chosen as the variables due to their previously established performance in predicting drug-specific metabolic perturbations.

Genetic algorithm creation, mutation and crossover parameters were used as implemented in the OptGene function of the COBRA Toolbox 2.0 (ref. 62). The genetic algorithm was solved using the Global Optimization Toolbox in MATLAB (MathWorks). A MATLAB implementation is provided.

Co-association analysis of drug side effect signatures. Statistical analysis of literature co-associations of side effects with side effect metabolite signatures was performed with (i) a non-parametric permutation test on the drug side effect
metabolite signatures against 1,000 permutations of the signatures for the AUC of predicting presence/absence of literature side effect/metabolite co-association in PubMed abstracts and (ii) a hypergeometric test for the enrichment of literature association among side effect/metabolite pairs in predicted signatures.

Construction of MDDB. To enable statistical analysis of predicted DISLoDGED pathways, disease–nutrient pairs were randomly selected from observed down-regulations, observed up-regulations and random associations chosen by resampling the former two lists through bootstrapping. These disease–nutrient pairs were then evaluated using established relationships using manual literature searches while blind to the origin of the pair to prevent investigator bias. The list of collected metabolism–disease relationships is not yet comprehensive because of the scope of the effort, but instead relationships were investigated in a targeted manner.

Data collection for the database covered literature up to and including May 2013.

Pathway analysis. We curated a database of 1,394 distinct GWAS publications34 and extracted 239 distinct disease-associated metabolic genes that overlapped with the 1,496 genes in Recon 1. We then assigned pathways to each disease-associated metabolic gene as well as to each DISLoDGED pathway calculated in our analysis (Supplementary Data 1). Disease-associated genes were assigned pathways based on the previously assigned pathway of corresponding reaction assigned in Recon 1. DISLoDGED pathways (which are metabolite centred) were associated with disease-linked pathways by determining the most frequent pathway among all reactions in which a metabolite takes part. To assess enrichment, hypergeometric tests were performed to determine whether the observed coverage of metabolites or genes was enriched or depleted in particular pathways, controlling for multiple tests were performed to determine whether the observed coverage of metabolites or DISLoDGED pathways (which are metabolite centred) were associated with the previously assigned pathway of corresponding reaction assigned in Recon 1. Only a single disease within the class was discarded, as were in conflicting cases (in which a nutrient was downregulated in certain diseases in the class and upregulated in others).

Side effect pathophysiology classifications. To assess whether common metabolic perturbations were observed among related diseases, we manually grouped side effects by pathophysiological disease class. Enrichment was assessed with the hypergeometric test, at a significance level of \( p < 0.1 \). Enrichment where there was only a single disease within the class was discarded, as were in conflicting cases (in which a nutrient was downregulated in certain diseases in the class and upregulated in others).

Side effect-linked gene polymorphism search. We attempted to identify all cases of genetic basis for side effect incidence reported in the literature, including GWAS and targeted genetic studies. To be eligible for comparison, we required that the side effect be a near or exact match and the metabolic pathway of the susceptibility gene be within the scope of our model. For example, immune-related genes were excluded due to nonspecific metabolic association, whereas G protein–coupled receptors known to be regulated by particular metabolic pathways were included. We generally excluded genes related to drug pharmacokinetics, including drug metabolism and transport, as the effects of polymorphisms on susceptibility are generally nonspecific to the pathology. We also required that the pathology be manifested in nucleated cells (that is, excluding red blood cell pathologies), as gene expression changes are assumed to be irrelevant to pathologies of nucleated cells. Based on these criteria, well over 20 studies were excluded, whereas 9 genetic susceptibilities spanning 6 side effects were valid for comparison with predictions. We also mention two cases in which pharmacokinetic genes do overlap with conserved side effect–linked gene perturbations, suggesting possible interactions between pharmacokinetic and gene expression effects of drug perturbation. Lists of included and excluded studies are found in Supplementary Notes 3 and 4, respectively.

Nutrient deficiency literature search. To populate MDDB, we searched the literature for associations between the pathology of the side effect and deficiency relationships between the closest nutrient to the metabolic perturbation and the occurrence of the pathology of the side effect. In an automated search, PubMed abstracts were searched for a number of side effects using a list of deficiency synonyms (Supplementary Data 2). Statistical significance of enrichment of co-association PubMed abstract hits (presence/absence) among downregulated and upregulated nutrient pathways was assessed through a dual permutation analysis. For each perturbed side effect–nutrient pair, 1,000 permuted pairs were generated by first randomly selecting the presence/absence result for a random nutrient with the same side effect, and then randomly selecting the presence/absence result for a random side effect with the same nutrient, then averaging the result. Then, the number of permutations with total presence calls greater than the true observation was counted and divided by the total permutations, as is typical in permutation tests.

In the manual alteration search, a number of possible nutrient–disease relationships were identified, such as an inhibitor of the metabolic pathway causing the side effect, an inhibitor of the metabolic pathway curing the side effect and so on (Supplementary Data 2). Then, search terms were generated using synonyms, and PubMed abstracts were searched. Results were then filtered such that each nutrient–side effect pair was assigned as ‘upregulated activity associated with the disease’, ‘downregulated activity associated with the disease’, ‘no associated with the disease’ or ‘conflicting associated with the disease’. Inconsistent results were assigned as conflicting and were excluded from further analysis. Significance of enrichments of particular relationships among upregulated and downregulated pathways were then evaluated with the hypergeometric test.

In the manual deficiency search, deficiency relationships were defined such that the DISLoDGED nutrient pathway could meet any of three criteria to be considered a ‘hit’. First, the deficiency of the nutrient could be known to be associated with the occurrence of the side effect pathology. Second, supplementation with the nutrient is known to alleviate the side effect pathology. Third, a physiological dysregulation of the pathway is known to be associated with the side effect pathology. PubMed and Google Scholar were both used for this study. Patents were not accepted as valid references, unless associated with a peer-reviewed publication.

To determine whether DISLoDGED pathways are significantly predictive of side effect/nutrient deficiency relationships, we generated a list of ‘random’ side effect/nutrient pairs for comparison with DISLoDGED pathway–disease pairs through the random generation of the distribution and thus were unaware of the treatments or treatment distributions. Statistical tests were only performed on the blinded literature results. The initial blinded list was then expanded to examine additional DISLoDGED pathways and curated to ensure consistent criteria between investigators. Although the curated list is unblinded, we have a greater confidence in this list for research purposes due to the consistent criteria and expanded list of investigated relationships.

Overview of experimental validation of MetChange results. Drug–induced metabolic perturbations calculated using the MetChange algorithm were validated using in vitro experiments in the MCF-7 cell line for three drugs: metformin, genistein and haloperidol.

Sample preparation. Human MCF-7 breast carcinoma cells (American Type Culture Collection, HTB-22) were maintained in supplemented DMEM media (CellGro Mediatech, 10013CV) with 10% (v/v) fetal bovine serum (HyClone, SH3093003), 1% (v/v) antibiotic/antimycotic solution (Omega Scientific, PS-20), 2 µg/mL non-essential amino acids (HyClone, SH3028001), 1% (v/v) MEM vitam- ins (CellGro Mediatech, 25020CI), 1 mM L-glutamine (CellGro Mediatech, 25005CD). 4 × 10^6 cells were seeded in supplemented MEM medium (CellGro Mediatech, 15010CV) into 150 mm dishes. For labelling with [U-13C] glucose (Sigma-Aldrich, 389374), the medium was replaced with supplemented MEM with glucose total of which 50% was [U-13C] glucose. Approximately 2.6 × 10^6 cells were harvested by incubation with trypsin for 5 min at 37°C (Gibco, 25200-056) and adjusted for total protein amount (Thermo Scientific Pierce, 23227). Intracellular polar metabolites were extracted by rapid quenching with 50% methanol at −40°C; total lipids were extracted using chloroform (Sigma-Aldrich, 950330). The cell extracts were dried by vacuum evaporation. Lipid extracts were derivatized to form the corresponding oximes and trimethylsilyl derivatives. Acyl-carnitines were derivatized to their corresponding methyl esters. Polar metabolites were dissolved in 99.9% H2O with 0.75% 3-(trimethylsilyl)propionic-2,2,3,3-d4 acid (Sigma-Aldrich, 293040). Lipophilic metabolites were dissolved in 2H-chloroform with 1% tetramethylsilane (Sigma-Aldrich, 151831).

Drug treatment. Cells were exposed for a 24 h period to varying drug concentrations (as indicated in their respective citations below) while control cells were exposed to the corresponding DMSO (CAS 67-68-5, Sigma-Aldrich, D8418) dilution: metformin (glucophage; CAS 1115-70-4, 15169101, MP Biomedicals, stock 3 M in PBS), genistein (CAS 446-72-9, 10005167100 Cayman Chemicals, stock 100 mM in DMSO), haloperidol (haldol; CAS 52-86-8, 15369690 MP Bio- chemicals, stock 10 µM in DMSO). For acyl-carnitine measurements, the medium was supplemented with 1 mM L-carnitine (CAS 6645-46-1, Sigma-Aldrich, 0283). Each experiment was performed in three biological repeats.

NMR spectroscopy. NMR experiments were performed on a 500-MHz Bruker Avance spectrometer with a 5-mm TXI z-gradient probe (Bruker-BioSpin, Karls-ruhe, Germany) at 298 K. 13C enrichments were determined by one-dimensional 1H NMR difference spectroscopy (13C-coupled spectrum minus the 13C-decou- pled spectrum). Two-dimensional indirect-detected 13C, 1H-j resolved HSQC spectra were recorded over a time of 1.5 d per spectrum with 5,120 indirect 13C data points, at 80 p.p.m. 13C sweep width, 40 p.p.m. carrier position, 4,096 direct 1H increments, 3 s recycling delay and 8 scans. Before Fourier
transformation, the data were multiplied with a squared sine-bell window function, phase corrected and zero-filled to 8,192 data points, for indirect 13C sampling.

**Mass spectrometry.** Quantification of organic acids was accomplished by gas chromatography-mass spectrometry (ThermoFisher Trace GC Ultra/DSQ II single quadrupole mass spectrometer). Calibration standards were prepared in water and spiked with stable isotope-labelled internal standards. Detecting the derivatized organic acids was achieved by single ion monitoring of each derivative after gas chromatography separation. Acyl-carnitines were quantified by LC/MS/MS (Waters Ultra Performance LC/Triple quadrupole mass spectrometer) using fluo-injection analysis with electrospray ionization. Calibration standards of the acyl-carnitines were prepared in bovine serum spiked with stable isotope-labelled internal standards. Parent ion scanning was used to detect parent, acyl-carnitine molecular ions that produced a characteristic acyl-carnitine fragment ion, m/z = 99, formed by collision-induced dissociation.

**Enzymatic assays.** Pyridoxal 5'-phosphate or vitamin B6 was determined in an enzymatic assay (A/C Diagnostics, ACB6001), which we adapted to quantify tissue culture supernatant or intracellular Vitamin B6 content by using PBST buffer with 0.1% Tween-20 (CAS 9005-64-5, Sigma-Aldrich, P9416).

**Additional methods for analysis of metformin response.** Acyl-carnitine species from 2 to 16 carbons and organic acids of the tricarboxylic acid cycle were analysed by liquid chromatography mass spectrometry and show dose-dependent associations upon metformin treatment in MCF-7 cells. Acyl-carnitine (C2) values were determined in nmol mg⁻¹ protein (scaled by factor 1/1,000) and were compared with acyl-carnitine (C12-C16) levels in pmol mg⁻¹ protein. Organic acids were determined in nmol mg⁻¹ protein.

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D.C.Z., A.B., M.J.H., M.L.M. and B.O.P. conceived and designed the computational study. D.C.Z., A.B., K.J., M.J.H. and M.L.M. generated and analysed the computational data. F.V.F., J.W.S., M.L.M. and B.O.P. conceived and designed experimental study. F.V.F. generated and analysed the experimental data. D.C.Z., F.V.F., A.B., K.J., J.W.S., M.J.H., M.L.M. and B.O.P. wrote the paper. All authors have read and approved the final manuscript.

Additional information

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