A Novel Mathematical Model Describing Adaptive Cellular Drug Metabolism and Toxicity in the Chemoimmune System

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Abstract

Cells cope with the threat of xenobiotic stress by activating a complex molecular network that recognizes and eliminates chemically diverse toxic compounds. This "chemoimmune system" consists of cellular Phase I and Phase II metabolic enzymes, Phase 0 and Phase III ATP Binding Cassette (ABC) membrane transporters, and nuclear receptors regulating these components. In order to provide a systems biology characterization of the chemoimmune network, we designed a reaction kinetic model based on differential equations describing Phase 0–III participants and regulatory elements, and characterized cellular fitness to evaluate toxicity. In spite of the simplifications, the model recapitulates changes associated with acquired drug resistance and allows toxicity predictions under variable protein expression and xenobiotic exposure conditions. Our simulations suggest that multidrug ABC transporters at Phase 0 significantly facilitate the defense function of successive network members by lowering intracellular drug concentrations. The model was extended with a novel toxicity framework which opened the possibility of performing in silico cytotoxicity assays. The alterations of the in silico cytotoxicity curves show good agreement with in vitro cell killing experiments. The behavior of the simplified kinetic model suggests that it can serve as a basis for more complex models to efficiently predict xenobiotic and drug metabolism for human medical applications.

Introduction

Living organisms are permanently exposed to potentially toxic xenobiotic compounds including environmental toxins and also drugs administered for therapeutic purposes. Although tissue barriers, such as the skin, the intestinal epithelia or the blood brain barrier limit the entry of xenobiotics into the body or a specific organ, individual cells have to cope with significant xenobiotic stress. The majority of the xenobiotics are detoxified through the canonical Phase I, II, and III pathways [1–5]. Phase I pathways include oxidative, reductive and hydrolysis
role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors declare that co-author Gergely Szakacs is a PLOS ONE Editorial Board member. This does not alter the authors’ adherence to PLOS ONE Editorial policies and criteria.

reactions. The most prominent Phase I enzymes belong to the cytochrome P450 (CYP) family. CYPs (e.g. CYP3A4, CYP3A5, CYP2D6, CYP1A1, CYP1B1 and CYP2E1) recognize a wide range of chemicals as substrates, usually converting them into a more water soluble form [6]. The oxidized intermediates are further metabolized by the action of Phase II enzymes (e.g. UDP-glucuronosyltransferases—UGTs, glutathione S-transferases—GSTs), which neutralize Phase I products by conjugating them with small molecules [7]. Finally, conjugates are removed from the cells to avoid untoward accumulation. Phase III elimination is mostly linked to the activity of ABC (ATP Binding Cassette) transporters, including MRP1/ABCC1 and MRP2/ABCC2 [8, 9]. Additional ABC transporters (such as MDR1/ABCB1 and BCRP/ABCG2) can recognize unmodified xenobiotics and extrude them from the cell (or already from the cell membrane) in the so-called Phase 0 reaction, thus reducing the load on the entire metabolic process [4, 10].

The expression of phase 0-III enzymes and transporters is orchestrated by several nuclear receptors and transcription factors (e.g. NR1I2/PXR—pregnane X receptor, NR1I3/CAR—constitutive androstane receptor, and AhR—aryl hydrocarbon receptor) that recognize xenobiotics and often also their metabolites as ligands [1]. These regulatory processes converge to select for the most efficient set of proteins to protect the cell from the given xenobiotic. Based on the similarities of the cellular detoxification processes and the immune system (e.g. regulator and effector elements, differentiation of metabolites from xenobiotics (“self” vs. “nonself”, etc.), the complex network underlying cellular detoxication has been referred to as the “chemoimmune system” [4].

Therapeutic compounds are subject to cellular metabolism that influences both the ADME-Tox (absorption, distribution, metabolism, excretion, and toxicity) properties of drugs and also the drug-drug interactions [4, 9]. In many cases, metabolites may be more influential than parent xenobiotics. For example, it is estimated that three quarters of the carcinogens are activated from parent procarcinogens by CYPs [11]. Conversely, bioactivation of prodrugs is desirable. Cyclophosphamide is activated by CYP to form its pharmacologically active phosphoramid mustard metabolite [12, 13]. Similarly, morphine-6-glucuronide produced by UGTs from morphine is a more potent analgesic than morphine itself [14]. Although the pharmacological relevance of the chemoimmune system is universally appreciated, little is known about the interplay of the individual enzymes or the kinetic parameters of the regulatory mechanisms. To fully appreciate the complexity of this elaborate system, dynamic interactions between the participating enzymes should be considered [15].

Mathematical models are promising possibilities for the investigation of such elaborate systems. On the other hand, xenobiotics interact with many regulators, enzymes and transporters, but the kinetic parameters describing these reactions are often unknown. Thus the formulation of widely applicable, general models is a serious challenge. Recent attempts to model the detoxifying cascade followed a single, well characterized xenobiotic or focused only on limited aspects of cellular detoxication. Stamatelos et al. created models describing the metabolism of arsenic compounds. While their toxicokinetic model [16] describes only Phase I-III reactions, the combined toxicokinetic-toxicodynamic model [17] includes also a simple transcriptional-translational feedback loop in the form of the Keap1–Nrf2 pathway playing a role in oxidative stress response regulation [18]. The electrophilic stress response model of Zhang and Andersen confines itself only to Phase II-III reactions, but contains more details about the Keap1–Nrf2 pathway [19]. Up to now, probably the model of Zhang et al. covers most aspects of cellular detoxication [20]. Beside Phase I-III reactions and the Keap1–Nrf2 pathway (the latter is described by not only feedback but also by feedforward loops) it contains regulatory circuits based also on the AhR nuclear receptor. However, since this model is geared towards mechanisms underlying overcompensation by the homeostatic control systems, especially hormetic
response, and since the applied simplifications, it is suboptimal for more general modeling of
the chemoimmune system.

To be able to study the role of each metabolite, enzyme and reaction involved in cellular de-
toxication in the context of this complex network, we generalized and extended the scheme
proposed by Zhang et al. to create a more comprehensive model. We introduced ABC0 as a
transporter modeling Phase 0 efflux transporters operating at pharmacological barriers. Fur-
thermore, by making a clear distinction between xenobiotic metabolites and reactive species
produced by the endogenous metabolic processes of the cell, we were able to analyze their rela-
tive importance. Using time course simulations we demonstrated our model’s ability to recapit-
ulate elementary properties of the chemoimmune system. In addition, we introduced a novel
framework to describe the effects of toxic species on cellular fitness. This opened the possibility
for the calculation of in silico cytotoxicity curves resembling experimentally obtained results.
Our results of in silico toxicity analysis are in agreement with RNA silencing experiments de-
scribed in the literature [21, 22].

Methods

Generation of the model

The SBML (Level 2 Version 4) model format [23] was chosen for the implementation of the
model. The model was created by using the graphical user interface of COPASI (version 4.8),
the biochemical network simulator package [24] and custom Python scripts employing
libSBML programming library (version 5.8.0) [25]. Time course simulations were carried out
in COPASI, through its Python bindings. For numerical integration the deterministic
(LSODA) solver was applied along with its default settings. Typically, the duration of the simu-
lations was 60 hours with an interval size of 60 seconds. Initial conditions (here: concen-
trations) for time course simulations were steady state conditions corresponding to the given
parameter set, and to zero extracellular xenobiotic concentration ([Xe]). The initial conditions
were calculated by running a preliminary, 30 day long time course simulation, sufficient for the
system to reach steady state. Details of the model are provided as Supporting information.

Biological and chemical details of the model

The chemoimmune network model is based on the scheme proposed by Zhang et al. [20],
which describes the metabolism of xenobiotic X including the following reactions:

Phase I CYP oxidizes X to X’, Phase II GST conjugates GSH to X’ forming X”’, which in turn
is excreted from the cell by a membrane transporter (MRP). Transcription and translation of
the participating enzymes are regulated by the AhR nuclear receptor and the Nrf2 transcription
factor. The model contains three compartments: the extracellular space, the cytoplasm and the
nucleus. In order to create a model tailored to the study of the chemoimmune system, this ini-
tial scheme was significantly modified (S1 Fig.). In brief, the following changes were
introduced:

(1) ABC0, a Phase 0 transporter which is capable to eliminate the cytoplasmic form of X
(Xc) from the cell was introduced. (2) The transcriptional and translational regulation of ABC0
was added to the scheme. (3) Some parameters were altered to obtain biologically more realistic
concentration ranges. For example the transcription rate constant of ABCIII (a general Phase
III ABC transporter; formerly MRP) was reduced to decrease its level, while the transportation
rate constant of X”c was increased to keep its transportation rate roughly unaltered. For mathem-
tical details of the model see S1 Model and S1–S4 Tables. (4) Reactive species originating
from background reactions (X’bc) and its conjugated form (X”bc) were introduced (see Results
for rationale). X’bc and X”bc were treated as respective xenobiotic metabolites X’c and X”c.
and $X'_{bc}$ are substrates of GST, while $X''_c$ and $X''_{bc}$ are substrates of ABCIII. To avoid extremely complex enzyme kinetics functions describing the metabolism of these two pairs of alternative substrates, instead of using conventional competitive inhibition formulae the following simplification was applied. In equations describing conjugation by GST and transport by ABCIII, the total enzyme amount was divided according to the concentration ratios of the alternative substrates, and for each substrate this reduced amount of enzyme was used in the calculations. (5) $X''_e$ (conjugated and extracellular product) and $X''_{be}$ (conjugated and extracellular product from the background reactions) were introduced to be able to measure the amount of metabolites eliminated by ABCIII. (6) A new aspect, a flexible description of cellular fitness was implemented using SBML parameters and events (S1 Text, S5–S6 Tables). (7) The ABCIII dimerization step was eliminated, since members of the ABCC family, participating in this Phase are full transporters. (8) MRP and AhR were renamed to ABCIII and NR, respectively to emphasize that they represent a general Phase III ABC transporter (e.g. MRP1, MRP2) and a general xenobiotic nuclear receptor, respectively. Other components related to the regulation by NR were also renamed.

**Calculation of cellular fitness**

In order to simulate the impact of toxic species on cellular fitness, the model was extended with the variable Fitness (Fig. 5). To describe the toxicity of a drug or metabolite, its Critical concentration was defined as the threshold concentration, which must be exceeded to cause cellular damage when the given compound is assumed to form the sole toxic species in the cell. In addition to the inherent toxicity, the concentration and exposure time of toxic species also have to be considered for the assessment of cellular damage. To take the respective concentrations into account, the concentration of each toxic compound is divided by its Critical concentration and a nonlinear function of the sum of these fractions is defined as the Chemical load. Chemical load expresses chemical stress coming from all toxic species as a single value at any given time point.

The ability of the cells to recover from moderate damage caused by chemical stress is modeled by introducing a constant Regeneration capacity. According to the model, the cell can be in one of three states, defined by relations between Chemical load, Regeneration capacity, and Fitness. When Chemical load is lower than Regeneration capacity, repair mechanisms can compensate chemical stress, the cell may recover from previous damage. If Fitness is maximal, no regeneration occurs, since the cell is already in perfect shape. When Fitness is below its maximum, the cell undergoes regeneration to increase Fitness. When the Chemical load exceeds Regeneration capacity, repair mechanisms are insufficient to compensate chemical stress, the cell undergoes damage, Fitness is decreasing. The rate of regeneration and damage is the function of Chemical load. Thus, the exposure time is also taken into account in damage calculation. Regeneration and damage are described by the variables Regeneration and Damage, respectively.

Since cellular fitness is always the overall result of damage and regeneration, Fitness is calculated by adding the (scaled) positive values of Regeneration and the (scaled) nonpositive values of Damage to the maximal value of Fitness. At the start of the simulation experiment, Fitness has its maximal value (1), which represents the state of perfect health. Minimal Fitness (0) represents a state when the cell dies due to severe damage. Three characteristically different Fitness profiles are shown in S2 Fig. A more technical description of Fitness calculation is provided as Supporting information (S1 Text, S5–S6 Tables).

**Results**

We have generated a new reaction kinetic model of the cellular chemoimmune network, based on the model proposed by Zhang et al. [20]. The Zhang-model was created to investigate
hormesis, which was demonstrated through the function of participants of Phase I-III metabolism. Since our aim was to study the network as a whole system under a wider range of drug concentrations and various toxicity conditions, we have implemented several changes and additions.

A simplified view of the chemoimmune model is shown in Fig. 1 (a more detailed description is provided in S1 Fig.). The model describes the interaction of a general xenobiotic compound (X) with a cell. The extracellular concentration of the xenobiotic ([Xₐ]) is set to a constant value, based on the assumption that the extracellular space contains many orders of magnitude more X than the cell (at the end of the 60 hour long simulations in Fig. 2 the ratio of the cumulated drug intake and the total amount of drug in the extracellular space is in the range of 10⁻⁷), thus the amount of xenobiotic metabolized by the cell is negligible compared to the total available amount. The xenobiotic molecule enters the cell by diffusion through the plasma membrane. To model Phase 0 extrusion at “the gates” by plasma membrane ABC transporters such as ABCB1/MDR1, ABCC1/MRP1 or ABCG2/BCRP, we introduced ABC₀, which eliminates Xₐ, that is the unmetabolized, cytoplasmic form of the xenobiotic, thus limiting intracellular drug concentrations. Xₐ may also be converted to an oxidized metabolite (X'ₐ) by CYP, or may bound to a nuclear receptor (forming XNRₐ), or may be distributed to the nucleus (Xₐₙ) by diffusion. X'ₐ is conjugated by GST to form a more water-soluble metabolite (X''ₐ), which is in turn eliminated by ABC₃ (modeling Phase III efflux transporters) from the cell.

It should be noted that in the model of Zhang et al. X'ₐ represented not only the oxidized xenobiotic under investigation, but also reactive species (e.g. free radicals), produced by the basal metabolism of the cell. Since we focus on the effect of the xenobiotic and its metabolites, we introduced X'ₐₙ and X''ₐₙ to separately track background reactive species and their conjugated
forms, respectively. This distinction allows the separate monitoring of the amount of xenobiotic metabolites and those produced by background processes. The intracellular forms of the xenobiotic \((X_c, X_n, X'_c, X''_c)\) and the endogenous metabolites \((X'_bc, X''_bc)\) have distinct toxicity parameters, reflecting the respective toxicity of each metabolite.

Transcriptional and translational modulation of the effector transporters and metabolic enzymes in the cell is significantly affected by detailed regulatory circuits. In order to describe these processes the model contains a nuclear receptor, NR and the Nrf2 (NFE2L2) transcription factor [26]. NR provides a model for promiscuous nuclear receptors recognizing a wide range of drugs and xenobiotics. Since these regulatory elements interact with \(X_c, X'_c\) and \(X''_c\), the system contains multiple feedback and feedforward loops. The complete wiring diagram of the model is shown in S1 Fig.

Biochemical reactions of the modeled species are mathematically described by applying the laws of biochemical reaction kinetics. The resulting set of ordinary differential equations, their parameters and initial conditions together form the mathematical model. Since these differential equations are mostly nonlinear equations and cannot be solved analytically, numeric integration is required to solve them.

The kinetic model recapitulates basic properties of the chemoimmune system

We performed simulations to investigate the basic behavior of the modeled network. First, the effect of extracellular xenobiotic concentrations on the expression levels of the network elements and the cellular concentrations of the produced metabolites were studied. Second, levels of various components were monitored as a function of the cellular uptake properties of the

![Image](https://example.com/image1.png)

**Fig 2. Effect of extracellular drug concentration on the level of network components.** Six time course simulations were run from steady state as described in Methods. At \(t_0 = 0 \text{ h}\) the addition of drug was simulated by setting the extracellular drug concentration \([X_e]\) to a constant positive value between \(1 \text{ nM}\) and \(10 \mu\text{M}\). a-c Concentration profile of the cytoplasmic form of the drug \([X_c]\), its CYP-oxidized metabolite \([X'_c]\) and GST-conjugated form \([X''_c]\). d \(n(X'_c)/n(X_{in})\) ratio, where \(n(X'_c)\) is the amount of the ABC01-excreted extracellular form of \(X'_c\) and \(n(X_{in})\) is the amount of drug intake. (Details of drug intake calculation are provided in S1 Text.) e-h Concentration profile of four key enzymes (ABC0, CYP, GST and ABCIII).

doi:10.1371/journal.pone.0115533.g002
xenobiotic (by changing the passive diffusion rate). Third, the impact of changes in the affinities of a metabolic enzyme (GST) and a transporter (ABC₀) to their substrates were followed.

As expected, increasing the extracellular xenobiotic concentration ([Xₑ]) resulted in an increased cytoplasmic concentration of the xenobiotic ([Xₘ]) and its oxidized ([X’ₑ]) and conjugated ([X”ₑ]) forms in the time course simulations (Fig. 2A-C). Following extracellular xenobiotic administration at 0 h, [Xₑ] usually reaches a quasi-steady state level in seconds. Due to ABC₀ activity, this level is always lower than the corresponding extracellular xenobiotic concentration. Xₑ triggers a regulatory response, which results in increasing ABC₀ levels and monotonically decreasing [Xₑ] on an hourly scale (Fig. 2A).

At relatively low xenobiotic doses, levels of X’ₑ and X”ₑ show a transient increase (at ca. 0–10 hours), before starting to decrease at 5 to 10 hours (Fig. 2B, C). This behavior is the result of multiple effects. First, Xₑ and X’ₑ upregulate the expression of transporters and metabolic enzymes (Fig. 2E-H). Initially, the CYP level grows faster than the GST level (compare Fig. 2F and G). As a result, the production rate of X’ₑ increases faster than its consumption rate, which results in the increase of [X’ₑ]. Faster consumption of Xₑ by CYP at the same time leads to the faster decrease of [Xₑ]. Decreasing Xₑ concentrations slow down the production of X’ₑ by decreasing the reaction rate of the CYP-catalyzed oxidation and by transcriptional and translational downregulation of CYP.

Interestingly, considering a wide range of extracellular xenobiotic concentrations, two characteristically different X’ₑ and X”ₑ concentration profiles can be distinguished (Fig. 2B, C). At lower xenobiotic doses, the transient concentration increase is moderate, as described above. When applying higher xenobiotic concentrations, X”ₑ levels reach a maximum, due to the saturation of GST (Fig. 2C). Since saturation of CYP happens only at higher substrate levels compared to GST, the oxidized xenobiotic accumulates, resulting very high X’ₑ concentrations (Fig. 2B).

Cellular accumulation of metabolites—an important factor in drug action—can also be inferred by calculating the ratio of the amount of the ABCIII-excreted extracellular metabolite (n(X”ₑ)) and the amount of cumulated xenobiotic intake (n(Xₑ)) (Fig. 2D). At lower doses, the n(X”ₑ)/n(Xₑ) ratio is close to 100%, suggesting that most of the xenobiotic molecules are eliminated from the cell. Conversely, at higher xenobiotic concentrations this ratio drops significantly during the first hours of the simulation, which indicates severe metabolite accumulation.

It is important to note that the concentration of Xₑ is mostly in the nanomolar, while concentrations of X’ₑ and X”ₑ are in the micromolar range (Fig. 2A-C). This difference originates from the high capacity of CYP, producing X’ₑ. Parameters of conjugation are based on experimental evidence, while parameters of oxidation were set by assuming CYP is not easily saturated. Since this enzyme catalyzes a practically irreversible reaction, and X’ₑ has no other way to be converted back to Xₑ, in our model the consumed Xₑ can only be supplemented from other processes, which are the diffusion from the extracellular space (infinite source) and the nucleus, and dissociation from various complexes containing the unmetabolized form of X. Taking together, the model suggests that CYP activity can easily lead to X’ₑ concentrations orders of magnitude higher than the concentration of Xₑ.

Role of individual components in detoxication

The impact of individual components was studied by selectively altering the respective kinetic parameters. Increasing the passive diffusion rate of the xenobiotic through the membranes (which can reflect e.g. increased hydrophobicity of the xenobiotic) results in elevated cytoplasmic xenobiotic concentrations ([Xₘ]) and—to a smaller extent—in elevated metabolite (X’ₑ and X”ₑ) concentrations (Fig. 3A-D). When the diffusion rate is high, xenobiotic extrusion by
ABC0 cannot significantly limit xenobiotic entry (Fig. 3A). Therefore, even though the concentration of Xc is much lower than the concentration of Xc', higher diffusion rates (or the presence of uptake transporters) potentially have severe adverse effects in the cell. Thus, similar to the case of high extracellular xenobiotic concentration (Fig. 2A, B), at high diffusion rates the saturation of GST leads to extremely high Xc' concentrations (Fig. 3A, B). These results highlight the relevance of xenobiotic diffusion rates in pharmacology, and also suggest that the incorporation of uptake transporters into the model could open further perspectives in understanding the kinetics of cellular xenobiotic metabolism [27]

We simulated the consequence of increased ABC0-mediated xenobiotic efflux by decreasing the apparent Michaelis constant (Km; Fig. 3E-H). In case of moderate substrate concentrations, lower Km values result in increased efflux, which leads to lower [Xc], and consequently also to reduced metabolite (Xc' and Xc") concentrations. As expected, the increased diffusion rate can be compensated by an increase in the affinity of ABC0 (S3 Fig.). Similarly, by lowering the Km of GST, Xc' concentrations decrease (Fig. 4B), while Xc" levels increase due to the higher production rate (Fig. 4C).

Interestingly, our model predicts that reactive species produced by normal cell metabolism (Xbc) may play an important role in preconditioning the chemoimmune system. Decreased affinity of GST to Xbc (increased Km) results in elevated baseline concentrations of Xbc, which in turn leads to increased baseline ABC0 levels (since Xbc upregulates ABC0 through the Nrf2 pathway), even in the absence of extracellular xenobiotics. (Compare basal ABC0 levels at t = 0 h on Fig. 4H which are steady state values calculated before simulating the addition of the xenobiotic to the system.) The baseline ABC0 level has a twofold significance. As expected, ABC0 function results in lower Xc levels (Fig. 4A, E). In addition, when higher xenobiotic doses are
applied, increased baseline ABC0 activity protects the cells against initial $[X_c']$ peaks (Fig. 4F). Thus, the presence of reactive species (e.g. free radicals) increases the initial detoxication performance of the cell. This feature of the system can be viewed also as a hallmark of hormesis: moderate stress may be beneficial for the cell to react faster during the need of chemodefense [28].

Model of cellular fitness to study toxicity

In addition to describing changes in the concentration of intracellular xenobiotic forms, we wished to model the cytotoxic effect of the compounds. To this end we extended the model with a versatile description of the impact of toxic species on cellular fitness (Fig. 5), described by the variable $Fitness$ (blue curve). This is a general description which does not contain any assumption about the mechanism of toxicity, and should be considered as an approximation rather than a detailed representation of toxic effects.

Typical toxicity studies use only single parameters (e.g. $EC_{50}$, $LD_{50}$), and are not informative regarding the toxicity of the various intracellular drug forms. In our model, we have assigned a toxicity parameter to each intracellular form of the xenobiotic, and indirectly calculated an overall $EC_{50}$-like value, allowing the simulation of toxic effects of drugs with different ‘toxicity profiles’. See Methods for implementation details of $Fitness$ calculation.

In silico cytotoxicity curves

In order to test one of the possible practical applications of the kinetic model, we aimed to calculate drug cytotoxicity curves. The analogous diagrams in the model are $Minimal Fitness vs$
Curves, generated by running multiple time course simulations, by using increasing extracellular xenobiotic concentrations and measuring the minimal Fitness values in 48 hours (the length of a typical in vivo cytotoxicity test) (Fig. 6). Minimal Fitness is used instead of the Fitness value belonging to the endpoint of the simulation, as the final value reflects the actual state of the cell and does not include information on decreased metabolism (e.g. cell division), which is measured in in vivo experiments. Therefore, similar to an experimental cytotoxicity curve, an in silico cytotoxicity curve also allows the deduction of the EC50 value: it can be defined as the external xenobiotic concentration which causes 50% decrease of Fitness in the time course of 48 hours.

An important difference between experimentally determined and in silico cytotoxicity curves is that the former is calculated by investigating a cell population, and the latter is calculated by deterministic modeling of a hypothetical single cell. In the theoretical case, minimal Fitness values between 1 and 0 can be interpreted like partial growth inhibition, and zero Fitness as cell death.

First, we investigated the impact of the toxicity of the unmetabolized form ([Xe,crit]) on the in silico cytotoxicity curve (Fig. 6A). Assuming that X'_c is more toxic than X''_c (this is the typical case) the critical concentrations of X'_c and X''_c were set to constant values of 1 μM and 100 μM, respectively. The critical concentration of X_c was varied between 0.05 nM and 5 μM. When [X_c,crit] = 0.05 nM, and X_c is more toxic than X'_c, the corresponding curve was shifted to lower concentrations (EC50 = 4.38 nM) from the curve with [X_c,crit] = 0.5 nM (EC50 = 25.72 nM). Contrary, when [X_c,crit] = 5 μM, and X_c is less toxic than X'_c, the corresponding curve was shifted to higher concentrations (EC50 = 47.97 nM). The curves with [X_c,crit] = 5 nM and 5 μM are close to each other, indicating a maximal EC50 threshold close to [X_c] = 53 nM. The
**Fig 6.** *In silico* cytotoxicity curves reveal impact of drug’s toxicity profile and protein level on survival.

Time course simulations were run for up to 48 hours from steady states belonging to different parameter sets and extracellular drug concentrations ([Xe], set at t0 = 0 h) as described in Methods. The minimal Fitness values reached in simulations were plotted against [Xe] (colored circles; connected by interpolation curves—see S1 Text). Critical concentrations of Xc and X'c are constant on all panels with values 1 μM and 100 μM, respectively. Critical concentration of Xc ([Xc,crit]) is indicated on the panels. 

**a** Impact of the toxicity of the unmetabolized form of the drug ([Xc,crit]) on *in silico* cytotoxicity. *In silico* cytotoxicity curves were plotted for [Xc,crit] values from 0.05 nM to 5 μM. The EC50 value is indicated for [Xc,crit] = 0.5 nM. 

**b** Impact of transporter and oxidase levels on *in silico* cytotoxicity when Xc is more toxic than X'c ([Xc,crit] = 0.05 nM).

**c** Impact of transporter and oxidase levels on *in silico* cytotoxicity when Xc is less toxic than X'c ([Xc,crit] = 5 μM). In panels **b** and **c** *in silico* cytotoxicity curves were calculated after setting the transcription rates of ABC0 or CYP 0.2 or 5 times of its original value (S3 Table) to model lower or higher expression levels.

doi:10.1371/journal.pone.0115533.g006
existence of the threshold can be explained as follows: when \([X_c]\) is above this threshold, \([X'_c]\) exceeds \([X'_c,\text{crit}]\) during a long period of time, resulting in cell death.

Next, we investigated the influence of altered levels of ABC0 and CYP in two cases of different toxicity conditions. In the first case the drug is more toxic than its oxidized metabolite. This is a typical scenario, e.g. most docetaxel metabolites are more than 500 times less toxic than the drug itself [29]. In the opposite case the drug is less toxic than its oxidized metabolite. Among the important examples are antitumor prodrugs like cyclophosphamide [12]. As expected, when the ABC0 substrate \(X_c\) is more toxic than \(X'_c\), changes in ABC0 levels lead to much more pronounced effects than changes in CYP expression levels (Fig. 6B). Decrease of ABC0 level results in lower EC50 values, while an increase of ABC0 level results in higher EC50 values. However, when \(X_c\) is less toxic than \(X'_c\) (Fig. 6C), the effect of CYP modulation is more significant. Modulation of ABC0 is still important, since it indirectly influences the \(X'_c\) level. In this case both decrease of ABC0 and increase of CYP levels result in lower EC50 values, while increase of ABC0 and decrease of CYP level results in higher EC50 values. These results are in agreement with RNA silencing experiments, where cytotoxicity to doxorubicin increased after silencing of \(ABCG2\) [22], whereas cytotoxicity to cyclophosphamide decreased after the silencing of \(CYP3A4\) [21].

Discussion

In order to analyze the cellular chemoimmune network from a systems biology perspective, we have developed a detailed reaction kinetic model. Our model is the first to describe the metabolism of a single, general xenobiotic or drug covering all aspects of cellular detoxication from Phase 0 to Phase III and reflecting xenobiotic dependent transcriptional regulation. In addition to effectors (transporters and metabolic enzymes converting chemicals) the kinetic model also contains regulatory elements, including a general xenobiotic sensing nuclear receptor, the Keap1–Nrf2 oxidative stress response pathway, and sophisticated transcriptional and translational regulatory circuits. These pathways form multiple feedback and feedforward loops, through which the xenobiotic and its oxidized metabolite regulate their own transport and metabolism. Of course, any of these regulatory mechanisms are independently switchable by setting the appropriate reaction kinetic parameters. This way modeling the metabolism of xenobiotics e.g. without Phase 0 excretion is also possible.

The chemoimmune model recapitulates basic properties of the chemoimmune system. Increasing extracellular xenobiotic concentrations result in increased cytoplasmic concentrations of the xenobiotic metabolites, which in turn lead to the upregulation of transporters and metabolic enzymes through transcriptional and translational feedback and feedforward loops (Fig. 2). The cytoplasmic concentration of the unmetabolized xenobiotic is jointly affected by its diffusion rate, Phase 0 transporter activity and consumption by metabolic processes (Fig. 3). We find that increased affinity (decreased \(K_m\)) of a Phase 0 xenobiotic transporter for the xenobiotic can counteract increased xenobiotic influx (increased rate of diffusion) (S3 Fig.).

In various time course simulations the concentrations of the xenobiotic, its metabolites, metabolic enzymes and efflux transporters were followed and relevant scenarios corresponding to particular drug effects were studied. Simulations clearly indicate that the metabolites can reach much higher concentrations than the external xenobiotic concentration especially when an enzyme, which forms a bottleneck in the metabolic pathway is saturated, thus leading to an extremely elevated concentration of its substrate (Fig. 2). An example of this feature in our model is GST, which conjugates GSH to the oxidized form of the xenobiotic. Since the oxidized form neither can be reverted back to reduced form, nor can be removed by other pathways, saturation of GST leads to an extreme increase in the level of the oxidized form (Fig. 2B, D).
real cell, the oxidized form may be cleared by pathways other than GSH conjugation. On the contrary, the unmetabolized form of the xenobiotic cannot be concentrated in the cell, since even in the case of limited oxidase (CYP) action, it still can be cleared at Phase 0 due to ABC transporter action. The model suggests that the role of Phase 0 exporters extends beyond limiting the cellular entry of toxic molecules to prevent the buildup of extreme metabolite concentrations and the saturation of metabolic enzymes.

Interestingly, saturation of metabolic enzymes is also prevented by reactive species produced by endogenous metabolic processes of the cell. Our simulations suggest that the increased levels of reactive species produced by normal cell metabolism prevent extreme metabolite peaks (Fig. 4F) by elevating basal enzyme expression levels (Fig. 4H). Thus, a low level exposure to toxic molecules can prepare the system for a massive attack of xenobiotics. This characteristic of the chemoimmune system resembles the phenomenon of active immunization in the context of the immune system, making the analogy between the immune and chemoimmune systems even more fascinating.

By means of the newly developed toxicity framework, the model became suitable to predict toxicity parameters of \textit{in silico} cytotoxicity assays (Fig. 6). The toxicity module—despite being simplified—is able to qualitatively reproduce the results of cell killing experiments [21, 22]. An important limitation of our model in this regard is that in order to provide predictive systemic effects, the toxicological properties of the modeled drug metabolites have to be independently estimated. Still, from relevant experiments these values can be reinserted into the model to study the effects of variable cellular parameters.

An important medical aspect is the involvement of the chemoimmune network and more specifically multidrug ABC transporters in cancer drug resistance, a major cause of the failure of chemotherapy in malignant diseases. Potent inhibitors against ABCB1 have been developed to overcome cancer multidrug resistance (MDR), but the therapeutic applications were unsuccessful in major clinical trials [30]. These results underline the importance of ABC transporter analysis in the context of the whole chemoimmune network [31, 32]. By providing a system level description of this network, the chemoimmune model provides powerful means for the comprehension of side-effects caused by ABC transporter inhibition, and may serve as a valuable aid in the development of novel therapeutic strategies.

In summary, the chemoimmune model describes cellular xenobiotic metabolism well at the level of the applied simplification. In its present form, it is a capable tool for the investigation of dynamic interactions between components of the chemoimmune system and for the assessment of toxic effects on cellular viability. Also, it can serve as a basis of forthcoming, more advanced models. Introduction of xenobiotic uptake transporters (e.g. members of the SLC family) with saturable properties as compared to passive xenobiotic diffusion, implementation of further xenobiotic sensing nuclear receptors or further metabolic enzymes and transporters may lead to more realistic models suitable not only for qualitative, but also for quantitative predictions.

**Supporting Information**

**S1 Fig. Detailed representation of the model of the chemodefense network.** Wiring diagram of the chemodefense network represented as an SBGN [33] diagram. (PDF)

**S2 Fig. Characteristic ‘Fitness profiles’ for three different external drug concentrations.** Three time course simulations were ran from steady state as described in Methods after setting different extracellular drug concentrations ($[X_e]$) at $t_0 = 0$ h. Critical concentration of $X_c$, $X'_c$ and $X''_c$ was set to 5 nM, 5 μM and 5 mM, respectively (S3 Table). Concentration of $X'_c$ (solid
curves), and Fitness (dashed curves) were plotted. Simulating drug administration by setting relatively low external drug concentration (green curves) results only in transient Fitness decrease, since decreasing concentration of toxic compounds allows regeneration. Moderate drug concentration (blue curves) leads to monotonic Fitness decrease to an intermediate level till the end of the simulation experiment. Considering that typical cytotoxicity tests are run up to 24 or 48 hours, the observed behavior can be interpreted as the sign of cytostasis or partial growth inhibition. When applying relatively high drug concentration (red curves), Fitness decreases to zero before the end of the experiment, which signs the death of the cell. When the cell dies, simulation is interrupted.

S3 Fig. Simultaneous alteration of diffusion rate and ABC0 affinity to drug can compensate each other’s effect. Five time course simulations were run from steady states belonging to different parameter sets as described in Methods. The Michaelis constant of ABC0 and the diffusion rate constants were altered simultaneously. The extracellular drug concentration ([Xe]) was set to 75 nM at t0 = 0 h. Similarity of concentration profiles indicate that opposing effects affecting xenobiotic transport through membranes can compensate each other. Parameter values are expressed as multiples of their default value (S3 Table) a-c Concentration profile of the cytoplasmic form of the drug ([Xc]), its CYP-oxidized metabolite ([X’c]) and the GST-conjugated form of the latter ([X”c]). d Concentration profile of ABC0.
Supplementary methods. Details of cellular fitness, in silico cytotoxicity curve, and drug intake calculation.

Acknowledgments
We appreciate the helpful discussions with Anna Lovrics (Institute of Enzymology, RCNS, Hungarian Academy of Sciences).

Author Contributions
Conceived and designed the experiments: AT AB GS BS TH. Analyzed the data: AT GS BS TH. Conducted simulations: AT.

References


