



Addressing Toxicity Risk when Designing and Selecting Compounds in Early Drug Discovery

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Abstract

Toxicity accounts for approximately 30% of expensive, late stage failures in development. Therefore, identifying and prioritising chemistries with a lower risk of toxicity, as early as possible in the drug discovery process, would help to address the high attrition rate in pharmaceutical R&D. We will describe how expert knowledge-based prediction of toxicity can alert chemists if their proposed compounds are likely to have an increased risk of causing toxicity. However, an alert for potential toxicity should be given appropriate weight in the selection of compounds to balance potential opportunities against downstream toxicity risk. If a series achieves good outcomes for other requirements, it may be appropriate to progress selected compounds and generate experimental data to confirm or refute a prediction of potential toxicity. We will discuss how multiparameter optimisation approaches can be used to balance the potential for toxicity with other properties required in a high quality candidate drug, such as potency and appropriate absorption, distribution, metabolism and elimination (ADME). Furthermore, it may be possible to modify a compound to reduce its likelihood of toxicity and we will describe how information on the region of a compound that triggers a toxicity alert can be interactively visualised to guide this redesign.

Introduction

Toxicity of drugs and clinical candidates remains a significant issue for pharmaceutical industry, leading to increased attrition and cost, late stage failures and market withdrawals. Recent data from CMR International [1] indicate that 22% of drug candidates entering clinical development in the period 2006 to 2010, failed due to non-clinical toxicology or clinical safety issues. In pre-clinical development, toxicity and safety issues accounted for 54% of failures (18% of all pre-clinical candidates). These expensive late-stage failures account for a large proportion of the cost of pharmaceutical R&D, recently estimated to be \$1.8B per marketed drug [2].

For many marketed drugs, toxicity remains an issue, causing adverse drug reactions (ADRs) and leading to black box warnings, restrictions on use and even withdrawals. These dramatically reduce or even eliminate the return on R&D and marketing investments and harm the reputation of pharmaceutical companies and the industry as a whole. A study by Lasser *et al.* [3] indicates that, of 548 new chemical entities approved by the U.S. Food and Drug Administration between 1975 and 2000, 10.2% acquired one or more black box warnings and 2.9% were withdrawn. Recent, high profile examples of market withdrawals include Cerivastatin (2001), Valdecoxib (2005, U.S.) and Rosiglitazone (2010, Europe). Of particular concern are idiosyncratic ADRs which, due to their rare occurrence, are unlikely to be detected during clinical trials.

From the sobering statistics above, it is clear that addressing failures due to toxicity would have a dramatic effect on the productivity of pharmaceutical R&D and the quality of the resulting drugs. Some toxicity is driven by the biological mechanism of the intended pharmacological action of a compound, particularly in the case of compounds intended for new, unvalidated targets. However, a significant proportion of observed toxicities are caused by unintended effects unrelated to the primary biological target. In the latter cases, it should be possible to reduce risk by focussing on structural motifs that are less likely to cause toxicity due to known mechanisms. Alternatively, if a risk of toxicity being observed in the clinic can be identified early in the process, in vitro or in vivo experiments can be prioritised to assess this risk before additional, downstream investments are made.

In the mid-1990s, a similar observation was made of a high rate of failure due to poor compound pharmacokinetics (PK) in clinical trials [4]. This led to the introduction of *in vitro* assays for high-throughput measurement of absorption, distribution, metabolism and elimination (ADME) properties in early drug discovery [5] and development of computational, or *in silico*, methods for the estimation of these properties [6] [7]. The result has been a reduction in the proportion of clinical failures due to PK issues from an estimated 39% in 1991 to approximately 10% in 2000 [8]. Unfortunately, in the same period, the overall failure rate was unchanged and the proportion of clinical failures attributed to toxicity or safety issues increased from approximately 14% to 30%. This, in turn has motivated a recent trend to develop and introduce *in vitro* assays earlier in the drug discovery process, to identify potentially toxic compounds and halt their progression. Similarly, *in silico* methods for the prediction of toxicity can help to guide the design and selection of compounds with reduced risk of toxicity.

This article will focus on 'knowledge-based' methods for prediction of toxicity (also described as 'rule-based'), that produce a semi-quantitative estimate of toxicity risk, based on experimental precedence for similar compounds. Other approaches, broadly described as 'statistical' methods, rely on fitting a mathematical model of compound characteristics to empirical data [9]. The output of such statistical methods are classifications of compounds as toxic or otherwise or predictions of a numerical measure of toxicity, e.g. LD₅₀. The principals that we discuss herein for the application of *in silico* methods to address toxicity risk in early drug discovery can apply equally to both approaches.

In the following sections we will describe the principles of knowledge-based prediction of toxicity and the challenges posed by application in early drug discovery. We will discuss how these methods may be applied to the selection of compounds, giving appropriate weight to predictions of toxicity against other important factors, and provide feedback on strategies for redesign to reduce toxicity risk. Finally, we will present two applications of knowledge-based toxicity predictions — one of recently approved drugs and the other in the context of a hypothetical hit-to-lead project — before drawing some conclusions.

Knowledge-based prediction of compound toxicity

Expert knowledge-based predictive systems for small molecules are designed to emulate the decision-making process of an expert by applying a form of artificial intelligence whereby a knowledge base of facts is used to make a prediction by inferring relationships between facts through a process known as reasoning. This allows for the introduction of associated data such as reactivity or knowledge of the mechanism of action, and can cope with uncertainty and conflicting data which is common in the field of toxicity prediction. In contrast, purely statistical approaches derive probabilities of toxicity by taking a dataset of compounds, identifying descriptors that show a correlation to activity and use this to predict the toxicity of novel compounds. Statistical systems have the advantage of being fast to implement and can more efficiently cope with large datasets when the endpoint is relatively simple. Expert systems are particularly well suited to making predictions for toxicities derived through multiple mechanisms for which only incomplete datasets are available. Expert systems can often provide more interpretable predictions with detailed supporting documentation.

The Derek prediction engine (www.lhasalimited.org) applied in the examples below, provides a prediction (active/inactive) for each toxicity endpoint. If no evidence of toxicity has been found, then 'No report' (nothing to report) is returned. A prediction of activity is typically associated with a structural alert, identifying the motif triggering the positive prediction, along with an associated likelihood. The likelihood qualifies this prediction; some of the likelihoods relating to positive predictions are shown in Table 1. In practise, it has been demonstrated that likelihood can be taken as a level of confidence since it correlates well with the accuracy of a prediction [10].

Table 1 Examples of the reasoning levels within Derek and their definitions

certain	the proposition (prediction) is known to be true
probable	there is at least one strong argument for the proposition and none against it
plausible	the weight of evidence supports the proposition
equivocal	There is an equal weight of evidence for and against the proposition

Expert systems are frequently applied in the later stages of drug development, where it may be necessary to produce an assessment of risk suitable for regulatory acceptance, or to design *in vivo* studies that should be undertaken to support a submission. In such cases, features including mechanistic interpretation, expert commentary, documentation, validation statistics and supporting data are particularly valuable. At this stage of the process, relatively few compounds are assessed for toxicity and the endpoints can be relatively complex meaning that training sets for *in silico* models tend to be sparse and do not always sufficiently capture the different mechanistic pathways at work. To overcome this, collaborative data sharing, through organisations such as Lhasa Limited, enables participating companies to gain knowledge of toxicities from proprietary data, without revealing confidential information such as biological targets or chemical structures.

In contrast, these methods have been less commonly applied in early drug discovery, where the numbers of compounds considered are much larger and the scientists using the predictions are less likely to be expert toxicologists. This makes detailed examination of each prediction, using detailed supporting information, impractical. In this scenario, toxicity predictions must appropriately integrated into decision-making processes, to provide intuitive guidance on reducing toxicity risk and facilitate collaboration with expert toxicologists where expert guidance is required.

Guiding Compound Design and Selection

Balancing Toxicity with Other Factors

A high quality drug must simultaneously satisfy many property requirements. Primary among these is achieving sufficient potency against the intended therapeutic target(s); however, to be both safe and efficacious a successful compound must also have appropriate ADME properties and, of course, avoid causing toxic effects at a therapeutic dose. Therefore, identifying high quality lead and candidate compounds is a delicate balancing act, often described as 'multi-parameter optimisation' (MPO) [11].

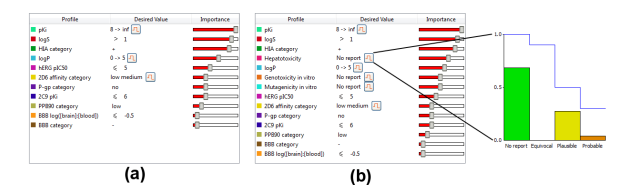


Figure 1 Example scoring profiles defining the ideal criteria for a range of experimental and predicted properties and the importance of each individual criterion to the overall objective of the project. (a) shows an example of a profile includes experimental potency against the target and predicted ADME properties. (b) illustrates a profile combining these properties with knowledge-based predictions of toxicity endpoints. Also shown in (b) is an expansion of the criterion for hepatotoxicity, demonstrating how the impacts of different predicted likelihoods for this toxicity on the chance of a compound's success can be reflected by a 'desirability function' shown in blue. The histogram shows the distribution of the different predictions in the current data set.

Predictions of toxicity risk must be balanced against other properties and given appropriate weight in the selection and design of compounds in early drug discovery. As discussed above, knowledge-based methods for toxicity prediction indicate if a compound has an increased likelihood of toxicity, but a toxicity alert is not a guarantee that a compound will be toxic. Therefore, one would give priority to compounds with no indications of toxicity over those with an alert, all other factors being equal; however, an alert may not be a sufficient reason to 'kill' a compound that meets many other requirements. The cost of incorrectly rejecting a good compound based only upon an uncertain prediction may be high, particularly in the absence of alternative options or if methods to mitigate the risk (such as a change to the dosing regimen) have not been considered.

Methods for MPO, such as Probabilistic Scoring [12], allow a project team to define a profile of property criteria that they require in an ideal compound. Furthermore, as illustrated in Figure 1, each property criterion can be assigned an importance to reflect the impact of a property outcome on a compound's chance of success. The results of predictions or experimental property measurements for each compound are then assessed against the profile to generate a score representing the compound's likelihood of success, i.e. the probability of achieving an ideal property profile. This allows compounds with the best chance of downstream success to be effectively prioritised. Furthermore, the uncertainty in each property value, due to experimental variability or statistical errors in predictions, can be explicitly taken into account to estimate the uncertainty in the overall scores. This, in turn, makes it clear when compounds can be confidently distinguished, based on the available data, and avoids inappropriate rejection of compounds based on an uncertain prediction or measurement.

Guiding Compound Re-design

An advantage of a knowledge-based approach to toxicity prediction is that the structural feature of a compound that is associated with an increased toxicity risk is identified. Highlighting this alert on the structure of the compound provides valuable information for medicinal chemists considering optimisation strategies. Coupled with predictive models of other properties and an MPO method in an interactive environment, this information can be used to guide the design of an alternative compound to reduce the risk of toxicity without having a negative effect on other required properties. Figure 2 shows an example of such an 'interactive designer'.

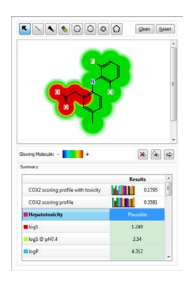


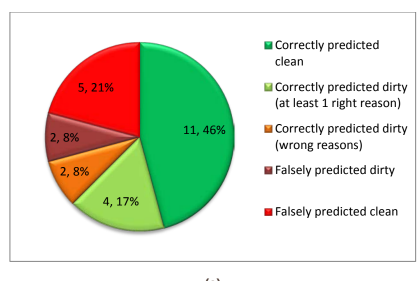
Figure 2 An example of an interactive designer in which the structural alert giving rise to the prediction of hepatotoxicity risk for Lumiracoxib is highlighted in red. Such an environment enables exploration of strategies to reduce toxicity risk while providing instant feedback on the predicted impact of structural changes on multiple, relevant properties.

Analysis of Recent Drug Approvals

In order to assess the potential value of applying knowledge-based toxicity prediction, all small molecule drugs approved in 2012 by the FDA Center for Drug Evaluation and Research were analysed against available endpoints in the Derek Nexus module for StarDrop (www.optibrium.com). The structures of these compounds were obtained from PubChem or NCI and regulatory label information was obtained from the FDA and/or EMA to identify clinically observed toxicities together with relevant black-box warnings. Three endpoints, skin sensitisation and irritation of the eye or skin were subsequently removed from the analysis since only one compound was topically administered hence predictions for these adverse events could not be validated. It should however be noted that the single topically administered compound, Ingenol Mebutate, was correctly predicted as a skin sensitiser. The predictions covered a range of important endpointsincluding hepatotoxicity, hERG channel inhibition, developmental toxicity, teratogenicity, chromosomal damage (in vitro and in vivo), mutagenicity in vitro and carcinogenicity. All of the alerts returned were at the plausible level (meaning that the weight of evidence is for activity to be observed). This full dataset is available as supplementary material. Of the 24 compounds, 11 were correctly predicted clean, 6 were correctly predicted with toxicities, 5 were falsely predicted clean and 2 were falsely predicted to have toxicities that were not observed. This is summarised in Figure 3(a). For the dataset of 24 compounds across 8 endpoints a total of 16 predictions of toxicity were made, showing a sensitivity of 55% and a specificity of 85%. The breakdown of those alerts is shown in Figure 3(b).

Looking at the performance of individual alerts, three complex high-level endpoints were responsible for the majority of false predictions. The first, hepatotoxicity is a challenging endpoint to predict since there are a large number of causes of liver toxicity, and reasons why it may not be observed. One compound, Aclidinium, was predicted to show hepatotoxicity but this is administered as a small inhaled dose making hepatotoxicity an unlikely event. Two compounds (Bosutinib and Bedaquiline) were not predicted to show the observed hepatotoxicity but these were dosed at very high levels (>400mg daily), and in the latter case, toxicity was only shown when co-administered with other drugs which inhibit CYP3A4 (the major clearance route for Bedaquiline). Large doses of hepatically cleared compounds increases the risk of liver toxicity through the saturation of processes or the build-up of metabolites. Developmental toxicity currently has only a limited number of alerts in the Derek engine and in this dataset only 1 out of the 7 observed instances was correctly predicted. Three of the 'false negative' compounds were kinase inhibitors which may indicate a lack of historical data from which to build good models since kinase inhibitors represent a relatively new class of There is however, growing evidence of a relationship between kinase inhibition and chromosomal/developmental toxicity [13] which should support further development of this alert. A third endpoint, teratogenicity, is incompletely understood, complex and driven by a complex array of pathways – for example, Lomitapide exhibited teratogenicity in rats and ferrets but not in rabbits Teriflunomide is believed teratogenic through its primary mechanism of action - inhibition of dihydroorotate dehydrogenase - an essential enzyme for nucleotide synthesis.

 $^{{}^{1}}http://www.fda.gov/downloads/Drugs/DevelopmentApprovalProcess/HowDrugsareDevelopedandApproved/DrugandBiologicApprovalReports/UCM342733.pdf$



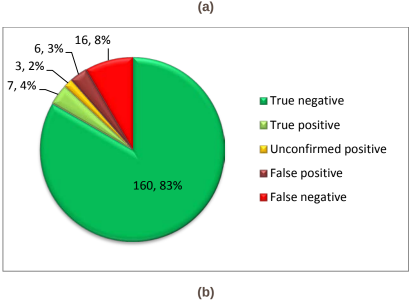


Figure 3 Results of predictions from the Derek Nexus module for StarDrop on the 24 compounds approved by the US Food and Drug Administration in 2012. (a) shows the analysis on a per-compound basis and (b) shows the analysis on a per-endpoint basis.

This analysis shows that knowledge-based toxicity predictions can be an effective tool to identify potential toxicities before a compound reaches the clinic. By flagging potential toxicities early in the drug development process, hazards can be assessed through early screening before significant investments have been made and by applying MPO analyses, these risks can be balanced against the potential benefits a drug may provide. This is highlighted in the case of Carfilomib which despite giving 6 alerts, 4 of which were confirmed (2 were not assessed) has been accepted as a treatment for cancer when other treatments are unsuccessful.

Application in Early Drug Discovery

To illustrate one workflow for the practical application of these methods in the context of a hit-to-lead project, we have used a public domain data set, derived from the ChEMBL database (https://www.ebi.ac.uk/chembl/). This data set contains 152 compounds from multiple chemical series for which the inhibition of the Cyclooxygenase 2 (COX2) enzyme has been determined experimentally, including the drugs Celecoxib and Lumiracoxib. This is typical of a data set containing primary screening data in a hit-to-lead project targeting a fast-follower for an existing drug.

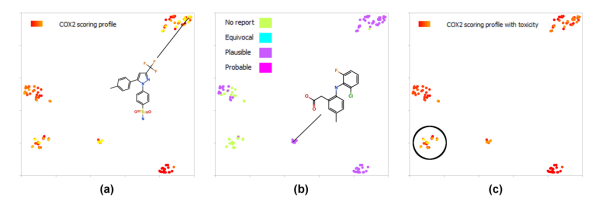


Figure 4 These 'chemical space' plots illustrate how predictions of toxicity risk can be combined with other experimental and predicted data to guide the selection of lead series in early drug discovery. Each point in a chemical space represents a single compound and the proximity of points indicates their structural similarity (2D path-based similarity calculated by a Tanimoto index [14]). (a) shows the compounds in a library of compounds with COX2 inhibition data containing 5 clusters of similar compounds, coloured by compound score from red (low) to yellow (high). The score was calculated using the profile shown in Figure 1(a), taking into account only potency and ADME properties. From this it can be seen that multiple clusters contain compounds with high-scoring compounds. For reference, the point corresponding to Celecoxib is identified. (b) shows the points coloured by predicted likelihood of hepatotoxicity, from which it can be seen that many regions of chemistry are predicted to have increased risk of exhibiting hepatotoxicity. The point corresponding to Lumiracoxib, a known hepatotoxin, is highlighted in this plot. In (c), this information is combined with the data for compound potency, predicted ADME properties and predictions for mutagenicity and genotoxicity using the scoring profile shown in Figure 1(b). The colours indicate low scoring compounds in red and high scoring compounds in yellow and the cluster containing the majority of high scoring compounds is circled.

Figure 4(a) shows the 'chemical space' of this library, in which the colour of a point represents the score of each compound against the scoring profile shown in Figure 1(a), including the experimentally measured target inhibition and a range of predicted ADME properties, but not considering predicted toxicity. This illustrates the distribution of the compound scores across the chemical diversity of the library and indicates that there are three clusters of similar compounds that are likely to yield compounds with a good balance of potency and ADME properties. These high-scoring compounds include the drugs Celecoxib and Lumiracoxib.

The toxicity risks of these compounds were then predicted using the Derek Nexus module for StarDrop for endpoints including mutagenicity, hepatotoxicity and genotoxicity. Figure 4(b) shows the prediction of hepatotoxicity mapped onto the chemical space of the COX2 library, which clearly shows that several of the clusters have plausible evidence of hepatotoxicity and should be considered with care. Among those compounds with evidence of hepatotoxicity is Lumiracoxib, which was withdrawn from the market in several countries, mostly due to hepatotoxicity concerns, and has never been approved for use in the United States.

The toxicity predictions can be combined with the *in vitro* and *in silico* data for other properties in an overall scoring profile, shown in Figure 1(b), giving appropriate weight to the predictions of toxicity against the other factors. The resulting scores are plotted in the chemical space shown in Figure 4(c), in which one cluster clearly stands out as having several compounds with the highest likelihood of yielding a high quality lead series with good ADME properties and reduced toxicity risk.

It is noteworthy that Celecoxib (the gold-standard COX2 inhibitor) is also identified as having plausible evidence of toxicity, illustrating the potential to balance the potential for toxicity against the benefits. One advantage of using a weighted scoring profile and taking into account the uncertainty in the underlying data, instead of a hard filter, is that the chemistry including Celecoxib would not be rejected outright. The score for Celecoxib (0.15 \pm 0.08) is not statistically significantly different from the top-scoring compound (0.45 \pm 0.30). This indicates that a rigorous strategy should select a small number of compounds from this series in order to experimentally confirm the required properties before making a final choice of lead series.

Finally, considering the structure of Lumiracoxib in Figure 2, a single functionality is highlighted as the cause of the structural alert for increased risk of hepatotoxicity, in common with all other members of this series. This suggests that approaches for reducing the this risk, while retaining potency and other desirable properties, can be investigated at an early stage before rejecting this class of compounds.

Conclusions

A key strategy to reduce the long timelines and spiralling cost of pharmaceutical R&D is to target safe and efficacious compounds as early as possible in the drug discovery process. Taking all available information into account, from predictive and experimental sources, as early as possible, increases the likelihood of delivering a high quality lead and, ultimately, a development candidate with an improved chance of success in the clinic. Furthermore, a lead series with a good balance of properties is less likely to require many, long and costly lead optimisation cycles; an important factor identified to reduce the overall cost per marketed drug [2].

Knowledge-based prediction of toxicity has an important role to play in this process, guiding the selection and optimisation of compounds when *in vitro* and *in vivo* toxicity data is often not available, due to the high cost and long timescales of experimental measurements. However, as with any predictive method, the uncertainties in the predicted outcomes should be taken into account and appropriate weight should be given to these results, relative to other property requirements for a high quality compound for a drug discovery project's objective. In this article, we have illustrated how toxicity predictions can be incorporated into an MPO approach to quickly identify compounds with an appropriate balance of properties and guide the optimisation of compounds with potential liabilities.

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Supplementary Information

The detailed results of the application of Derek Nexus for StarDrop to the 24 drugs approved by the US FDA in 2012 are provided in the supplementary information to this article.

References

- 1 Leeson, P. J. Am. Med. Assoc. 220. In *SCINOVO, Unlocking the Value of Drug Candidates* (Stevenage, UK 2013).
- 2 Paul, SM, Mytelka, DS, Dunwiddie, DT, Persinger, CC, Munos, BH, Lindborg, SR, and Schacht, AL. (2010) How to improve R&D productivity: the pharmaceutical industry's grand challenge. *Nat. Rev. Drug Discov.*, 9, 203-14.
- 3 Lasser, K.E., Allen, P.D., Woolhandler, S.J., Himmelstein, D.U., Wolfe, S.M., and Bor, D.H. (2002) Timing of New Black Box Warnings and Withdrawals for Prescription Medications. *JAMA*, 287, 2215-2220.
- 4 Kennedy, T. (1997) Managing the drug discovery/development interface. Drug Discov. Today, 2, 436–444.
- 5 Tarbit, M.H. and Berman, J. (1998) High-throughput approaches for evaluating absorption, distribution, metabolism and excretion properties of lead compounds. *Curr. Opin. Chem. Biol.*, 2, 411-416.
- 6 Van de Waterbeemd, H and Gifford, E. (2003) ADMET in silico modelling: towards prediction paradise? *Nat. Rev. Drug Discovery*, 2, 192-204.
- 7 Ekins, S, Boulanger, B, Swaan, PW, and Hupcey, MA. (2001) Towards a new age of virtual ADME/TOX and multidimensional drug discovery. *J. Comp. Aided Mol. Design*, 16, 381-401.
- 8 Kola, I. and Landis, J. (2004) Can the pharmaceutical industry reduce attrition rates. *Nature Rev. Drug Discov.*, 3, 711-716.
- 9 Tropsha, A. Recent trends in statistical QSAR modeling of environmental chemical toxicity. In Luch, A., ed., *Molecular, Clinical and Environmental Toxicology. Volume 3: Environmental Toxicology.* Springer, Basel, 2012.

- 10 Judson, P.N., Stalford, S.A., and Vessey, J. (2013) Assessing confidence in predictions made by knowledge-based systems. *Tox. Res.*, 2, 70-79.
- 11 Segall, M.D. (2012) Multi-Parameter Optimization: Identifying high quality compounds with a balance of properties. *Curr. Pharm. Des.*, 18, 1292-1310.
- 12 Segall, MD, Beresford, AP, Gola, JMR, Hawksley, D, and Tarbit, MH. (2006) Focus on Success: Using in silico optimisation to achieve an optimal balance of properties. *Expert Opin. Drug Metab. Toxicol.*, 2, 325-337.
- 13 Olaharski, A.J., Gonzaludo, N., Bitter, H., Goldstein, D., Kirchner, S., H., Uppal, and Kolaja, K. (2010) Identification of a kinase profile that predicts chromosome damage induced by small molecule kinase inhibitors. *Toxicol. Sci.*, 118, 266-275.
- 14 Rogers, D.J. and Tanimoto, T.T. (1960) A computer program for classifying plants. Science, 132, 1115-1118.